

**Air quality in a BaU
scenario**

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Effects of business-as-usual anthropogenic emissions on air quality

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Abstract

The atmospheric chemistry general circulation model EMAC has been used to estimate the impact of anthropogenic emission changes on global and regional air quality in recent and future years (2005, 2010, 2025 and 2050). The emission scenario assumes that population and economic growth largely determine energy and food consumption and consequent pollution sources with the current technologies (“business as usual”). This scenario is chosen to show the effects of not implementing legislation to prevent additional climate change and growing air pollution, other than what is in place for the base year 2005, representing a pessimistic (but feasible) future.

By comparing with recent observations, it is shown that the model reproduces the main features of regional air pollution distributions though with some imprecisions inherent to the coarse horizontal resolution (~ 100 km) and simplified bottom-up emission input.

To identify possible future hot spots of poor air quality, a multi pollutant index (MPI), suited for global model output, has been applied. It appears that East and South Asia and the Middle East represent such hotspots due to very high pollutant concentrations, although a general increase of MPIs is observed in all populated regions in the Northern Hemisphere. In East Asia a range of pollutant gases and fine particulate matter ($PM_{2.5}$) is projected to reach very high levels from 2005 onward, while in South Asia air pollution, including ozone, will grow rapidly towards the middle of the century. Around the Arabian Gulf, where natural $PM_{2.5}$ concentrations are already high (desert dust), ozone levels are expected to increase strongly.

The per capita MPI (PCMPI), which combines demographic and pollutants concentrations projections, shows that a rapidly increasing number of people worldwide will experience reduced air quality during the first half of the 21st century. Following the business as usual scenario, it is projected that air quality for the global average citizen in 2050 would be almost comparable to that for the average citizen in the East Asia in the year 2005, which underscores the need to pursue emission reductions.

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1 Introduction

Clean air is prerequisite to human health and well-being. In spite of the introduction of less polluting technologies in industry, energy production and transport during the past decades, air pollution remains a major health risk (HTAP, 2010). Many epidemiological studies have demonstrated the negative impacts of air pollutants on human health, in particular in the urban environment (e.g. Mokdad et al., 2004; Pope et al., 2002). Air quality is a major issue in the worldwide increasing size and number of megacities, i.e. population centres with more than 10 million inhabitants. Megacities are strong localized pollution sources though their effects can extend over large distances of more than 1000 km downwind (Lawrence et al., 2007). In many OECD (Organisation for Economic Co-operation and Development) countries air pollution is monitored routinely to comply with air quality legislation (EPA, 2005; UNECE, 1999; European Commission, 2008; Tørseth et al., 2012) and emission abatement strategies (Schöpp et al., 2010, and reference therein). Despite abatement strategies in some regions, air pollution may be expected to increase globally with industrial activity, with serious implications on a range of scales. In this work, the global EMAC (ECHAM5/MESSy, Atmospheric Chemistry) model is used to analyse the influence of anthropogenic emissions on air quality worldwide. Following Fenger (1999) the analysis includes the five major air pollutants which can significantly impact human health, i.e. Particulate Matter (particles with a diameter less than $2.5\ \mu\text{m}$, $\text{PM}_{2.5}$), nitrogen dioxide (NO_2), sulphur dioxide (SO_2), ozone (O_3) and carbon monoxide (CO). Our analysis of the influence of anthropogenic emissions is performed for selected time periods, i.e. preindustrial conditions, recent conditions (the year 2005) and projected future conditions (the years 2010, 2025 and 2050) using the recently developed Emission Database for Global Atmospheric Research, EDGAR-CIRCE (Climate change and Impact Research: the Mediterranean Environment).

Several emissions scenarios are present in the literature. The Special Report on Emissions Scenarios (SRES) scenarios families (Nakicenovic et al., 2000) developed by the Intergovernmental Panel on Climate Change (IPCC) are widely used in the

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literature for assessing the possible effects of future emissions on air quality on a global scale (e.g. Hogrefe et al., 2004; Jacobson and Streets, 2009). These scenarios presented a range of optimistic-pessimistic scenarios for gases influencing climate change, but did not consider further air pollution emissions controls, except for SO₂.

5 Recently the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP, www.giss.nasa.gov/projects/accmip/) was initiated, which focuses on emissions scenarios based on the Representative Concentration Pathways (RCP) (Lamarque et al., 2011; Meinshausen et al., 2011; van Vuuren et al., 2011b,a, and references therein). These scenarios, however, implicitly assume air pollution controls and are hence missing a “worst case” scenario.

10 Some studies performed similar analysis to this work (e.g. Szopa et al., 2006; Horowitz, 2006; Dentener et al., 2005, 2006; Stevenson et al., 2006). These simulations did consider optimistic, realistic and pessimistic cases. However, the resolution used in these studies was generally much lower than that used in this work and the projected simulation were performed only for the year 2030. The results presented here are expected to be more representative for densely populated regions. Finally, to our knowledge, no studies so far focused specifically on a combination of pollutants which effect human health, but rather on single pollutants (e.g. Knowlton et al., 2004).

15 In this work only the effects of anthropogenic emissions are investigated, thus ignoring possible effects of climate change on the atmospheric composition. Only long term air quality levels are analyzed (annual averages), as the short term concentrations are strongly influenced by the local meteorology, transport and deposition processes which, in turn, are influenced by climatic changes, not included in this work. We present a Business as Usual (BaU) scenario (which represents a possible future with continuation of the past emission trends and without additional legislation to abate increasing air pollution and climate change) to illustrate the effects on the atmospheric composition that can be avoided by implementing additional emission control measures. Although a range of emission scenarios is available (see Sect. 2) we focus on the pessimistic BaU scenario, firstly because the other scenarios are more similar to the current situation

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and secondly because a BaU scenario could be not too far from reality (Garnaut et al., 2008).

The work has two objectives. The first is to compare the influence of anthropogenic emissions on air quality in different regions according to a multi-pollutant index; the second is to demonstrate the impact of no-further legislation in the deterioration of air quality worldwide. In Sect. 2 the model is presented and the scenario simulations are briefly described. In Sect. 3 the model results are analysed, firstly by comparing the results for the year 2005 with observational datasets and then by calculating the changes of pollutant concentrations from the different simulations (projected future). In Sect. 4 the multi-pollutant index is defined to evaluate air quality levels and its time evolution is estimated on regional and global scales. Section 5 presents the conclusions.

2 Model description and setup

EMAC is a combination of ECHAM5 (Roeckner et al., 2006) (version 5.3.01) and the Modular Earth Submodel System (MESSy, version 1.9; Jöckel et al., 2005). An extensive evaluation of the model can be found in Jöckel et al. (2006) and Pozzer et al. (2007). In this study, the general circulation model (GCM) ECHAM5 has been used with a T106L31 resolution, corresponding to an horizontal resolution of $\approx 1.1 \times 1.1^\circ$ of the quadratic Gaussian grid, and with a vertical resolution of 31 level, up to 10 hPa in the lower stratosphere. The model set-up used in this work was presented by Pozzer et al. (2012), and here only differences and the central features are described. Additionally in Pozzer et al. (2012) an evaluation of the model results was performed, showing that the model reproduces the main spatial and temporal atmospheric distributions of sulfate and ammonium aerosols while nitrate aerosol shows some discrepancies compared to observations. The latter is partly attributed to the loss of nitrate from filters at relatively high temperatures, so that the observations are not representative for summer conditions.

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The atmospheric chemistry is simulated with the MECCA (Module Efficiently Calculating the Chemistry of the Atmosphere) submodel by Sander et al. (2005), while the aerosol microphysics and gas-aerosol partitioning are calculated by the Global Modal-aerosol eXtension (GMXe) aerosol module (Pringle et al., 2010a,b). For descriptions of the emission and deposition routines we refer to Kerkweg et al. (2006a,b), Pozzer et al. (2006) and Tost et al. (2007a,b).

To evaluate the effects of the emissions in the different air quality projections, the chemistry and the dynamics have been decoupled, hence there is no connection between changing greenhouse gas concentrations and the dynamics of the atmosphere, i.e. the GCM is used as a chemistry-transport model. More specifically, two sets of greenhouse gases (GHG) are included in the EMAC model: one used by the radiation code which is estimated from a 2000–2009 average and is the same in all simulations performed, and one used in the chemistry calculations, described below. The boundary conditions of the general circulation model ECHAM5, the Sea Surface Temperature (SST) and the ice coverage, were pre-calculated from a 10 yr climatology (2000–2009) from the AMIP II data (Taylor et al., 2000; Hurrell et al., 2008). Accordingly, all simulations follow the same (i.e. binary identical) dynamics and meteorology, which can be considered as a climatology of the first decade of the 21st century. The same dynamics between the experiments is essential for a comparison of the different simulations, as only the emissions/chemistry allowed to cause differences, as the meteorology is identical. In all model runs performed two years are simulated from which only the latter year output is analysed as the first one is considered as spin-up time.

In total five simulation experiments were performed. In each simulation only the anthropogenic emissions were changed, while all other emissions and the model set-up were kept the same. The emissions refer to:

- natural conditions (i.e. no anthropogenic emissions), termed the standard case natural (*SC_natural*).

- anthropogenic emissions for the year 2005, termed standard case 2005 (*SC_2005*).
- anthropogenic emissions for the year 2010, termed standard case 2010 (*SC_2010*).
- anthropogenic emissions for the year 2025, termed standard case 2025 (*SC_2025*).
- anthropogenic emissions for the year 2050, termed standard case 2050 (*SC_2050*).

More detailed information about the emissions is given in Sect. 2.

To take into account the effects of the GHG on atmospheric chemistry, mostly through changes of the hydroxyl radical by methane (CH_4) concentration changes, the GHG have been scaled to the observed or projected values in each simulation. The GHG used in the chemistry calculation are not used in the dynamics/radiation calculations. The projected GHG concentrations have been adopted from the scenario A1B of the IPCC recommendation (see IPCC, 2001, appendix II), and it includes CO_2 , CH_4 and N_2O , while CFCs and HFCs are from the WMO98 Scenario A1(baseline) following the Montreal Amendments. GHG estimates for the *SC_natural* have been obtained for CO_2 from Etheridge et al. (1998), for NO_2 from Machida et al. (1995), for CH_4 from Etheridge et al. (2002) and for CFCs from Walker et al. (2000).

Anthropogenic emission scenario

The global anthropogenic emission inventory (EDGAR-CIRCE) used in this study has been prepared in the framework of the CIRCE Project (Climate Change and Impact Research: the Mediterranean Environment) by the EDGAR group (EDGAR, Emission Database for Global Atmospheric Research) of the EC-Joint Research Center Ispra (Italy), Air and Climate Unit (Doering et al., 2009a,b, 2010). The European project

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CIRCE focuses on the EU neighbouring countries with emphasis on the Mediterranean area, using a scenario developed with a European Commission perspective. This dataset includes emissions of the nitrogen oxides family (NO_x), carbon monoxide (CO), Non Methane Volatile Organic Compounds (NMVOCs), ammonia (NH_3), sulphur dioxide (SO_2), Black Carbon (BC) and Particulate Organic Matter (POM) from fossil fuel and biofuel combustion, industrial processes, the production and usage of solvents, agriculture and waste treatment.

The emission totals of all world countries in the base year 2005 were projected up to 2050. Comparable to the study of Shindell et al. (2012) and in contrast to the RCP scenarios, the CIRCE projections differentiate the climate change (CC) measures from the air pollution control (AP) measures because the implementation of CC measures require a much larger and longer process of structural changes than the AP measures. Future emissions scenarios were developed to identify the regions that will become most affected and might be most concerned about co-benefits from emission control measures. For this paper we used a reference scenario based on energy and fuel projections of the POLES model (Prospective Outlook for the Long term Energy System), and agriculture, land use and waste projections of the IMAGE model (Integrated Model to Assess the Global Environment), incorporating only the presently agreed policies affecting emissions.

With this BaU scenario that leaves out potential additional future emission control measures, we focus on the worst case. It explores the potential consequences of absence of further climate and air pollution policies beyond what is in place since the year 2005. Other scenarios were produced under the EDGAR-CIRCE project, namely the Global climate policy only (CC), the European air quality policy (BAP) and the Integrated air quality and climate policy (CAP), which explore different combination of air quality and climate change reduction policies which may be applied in the future. The future emissions of pollutants and GHG equivalent for the CC, BAP and CAP scenarios lead to a stabilisation or an improvement of the current air quality (Doering et al., 2010). For example, the projected CO and BC emissions in the CC scenario is ~ 559

and $\sim 5.4 \text{ Tg yr}^{-1}$ respectively for the year 2050, which is -4.7% and -8.4% of the emissions estimated for the year 2005 while the BAP and CAP scenarios assume even larger decreases. As the scenarios present a large range of uncertainties without a clear indication of the most probable future situation, we focus on the scenario where no further effective implementation of air pollution legislation or structural measures in energy production systems will be applied (i.e. BaU). This rather pessimistic scenario is chosen to show the impact of refraining from further measures against air pollution and climate, indicating current legislation is associated with serious challenges even in EU and the US where air quality control has a tradition of many decades.

Beyond 2005 no change in technologies for combustion (power industry, transport), manufacturing processes (steel, cement, chemicals), waste treatment, crop production and manure management are assumed. Either the activity data or implied emission factors are scaled up. The energy-related activities are driven by population and economic growth but following the oil, gas and coal markets modeled in POLES for the different world regions. The BaU scenario of POLES assumes a continuation of the trend from 1990-2005: the energy consumption per GDP will decline further but will be more than offset by economic growth and is projected to be 79 % above its 2005 levels by 2050. The fossil fuels dominate for 75 % the fuel mix until 2030 and then they are assumed to decrease with 10 % assuming that the renewable and nuclear power will grow. The assumptions are well in line with those incorporated in world energy models from the International Energy Agency (IEA) World Energy Outlook 2006 as shown by Schade and Wiesenthal (2007). The agriculture-related activities are mainly driven by population and corresponding food demand, but following the global agricultural trade with regional food prices and with livestock-crop-land use interactions as calculated by IMAGE. As described by van Vuuren et al. (2009), the IMAGE baseline scenario projects the global agricultural production to grow by more than 50 % through increased productivity and an extension of the agricultural area with about 10 %. Both models, POLES and IMAGE used the medium-fertility population projection variant of the 2004 revision of world population prospects of United Nations Department of Economic and Social

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Affairs/Population Division (UNDP) (2005), expecting 9 billion people by 2050, ~45 % more than in 2005.

The total amounts of trace gases and aerosol emissions and the sectoral breakdown is shown in the Supplement.

5 The EDGAR-CIRCE emission dataset is gridded on a 0.1×0.1 degree latitude \times longitude grid using point sources locations for power industry, fuel production and process industry, line sources for the transport sectors and diffusive distributions using the Food and Agriculture Organization Geonetwork (www.fao.org/geonetwork) global land cover map and the Center for International Earth Science Information Network (CIESIN), rural and urban population as proxy data. A monthly distribution for agricultural sources based on LOTOS (Long Term Ozone Simulation; Builtjes, 1992; Veldt, 1992) and GENEMIS (Generation and Evaluation of Emission Data; Friedrich and Reis, 2004) is imposed. The emissions are distributed vertically in the model as described in Pozzer et al. (2009).

15 Natural/biogenic emissions have been calculated as in Pozzer et al. (2012) while volcanic emissions (particularly important for SO_2) are based on the AEROCOM dataset (Dentener et al., 2006). The natural/biogenic emissions are the same in all simulations.

The biomass burning contribution was added estimating the climatological emissions of the years 1997–2009 from the Global Fire Emissions Database (GFED version 3, van der Werf et al., 2010). The data has a $0.5 \times 0.5^\circ$ spatial resolution and a monthly temporal resolution. The same biomass burning emissions are used in all simulations performed.

3 Results

25 Though the EMAC model accounts for high time resolution variability (10 min time step) and applies monthly emissions, we largely focus on annual averages in the analysis of the results. The model has been used in a free running mode, i.e. without nudging to actual meteorological conditions. This prohibits comparing the results to instantaneous

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observations. The model is expected to represent the atmospheric dynamics on a climatological, i.e. decadal, time scale while single episodes do not temporally coincide with those in reality. Furthermore, many observational datasets are readily available as annual averages, without detailed information about sampling frequencies (Sect. 3.1).

5 Finally, we focus on surface values of the pollutant species because it enables comparison with station observations which are generally in locations where pollutant concentrations are considered to be most relevant for human health.

3.1 Year 2005 evaluation

The EMAC model has been extensively evaluated previously in different publications, both for gas phase trace gases (e.g. Jöckel et al., 2006; Pozzer et al., 2007), aerosol compounds (e.g. Pringle et al., 2010a; Pozzer et al., 2012) and for deposition (Tost et al., 2007a). In these efforts large numbers of observations have been used to test the model performance, e.g. in reproducing O₃, NO_x, CO, Volatile Organic Compounds (VOC) and semi-volatile inorganic aerosol species and gases such as HNO₃ and NH₃.
15 Nevertheless, additional comparison of model output to observations at the surface is useful, e.g. for SO₂, O₃ and NO₂, which play an important role in air quality.

Three different datasets were used to analyse the model results:

- EPA (the United States Environmental Protection Agency) network: ambient concentrations of pollutants in outdoor air are measured at more than 4000 monitoring stations operated mainly by state environmental agencies. The EPA provides an annual summary for each monitoring station (Edgerton et al., 1990), of which the data can be obtained from the AirData web site (<http://www.epa.gov/airdata>).
- EEA (The European Environment Agency): this agency of the European Union offers a range of information on the regional environment. The public air quality database system of the EEA is the AirBase (<http://www.eea.europa.eu/themes/air/airbase>). It contains air quality monitoring information of more than 30 participating countries throughout Europe. AirBase is managed by the European Topic

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Centre on Air pollution and Climate change Mitigation (ETC/ACM) on behalf of the EEA. In this work only annual statistics have been used (also available in the Airbase database).

- EANET (The Acid Deposition Monitoring Network in East Asia) network: this initiative supports regional cooperation among the participating countries (i.e. China, Indonesia, Japan, Malaysia, Mongolia, Philippines, Republic of Korea, Russia, Thailand, Viet Nam, Cambodia, Lao People's Democratic Republic and Myanmar). Regular monitoring started in January 2001. In this work, data from 28 stations of the EANET network were used. The data were downloaded from <http://www.eanet.cc/product/index.html>, (Asia Center for Air Pollution Research (ACAP), 2011).

Simulation *SC_2005* was used for comparison with the observations as the emissions for the year 2010 are projection based. Thus we compared the model results for the year 2005 with observations taken for the EEA, EPA and EANET dataset. It must be stressed that the EEA and EPA database were preferred to other database because they cover all the pollutants of interest. Additionally, the large numbers of stations allow a sound statistical comparison with the model using annually averaged values.

As shown in Table 1, and Figs. 1 and 2, the model reproduces the observed distribution well for O_3 and $PM_{2.5}$. Ozone is accurately simulated, especially compared to the EPA dataset. In this case the model average is close to the observations, and almost all modelled values lie well within a factor of two of the observations. In general more than ~80 % of the modelled values are within a factor of two, considering all datasets.

The dry $PM_{2.5}$ mass simulated by the model agrees well with the observations. Although the simulated values are generally low biased by 30–40 % compared to the EPA and EEA dataset, they are mostly within a factor of 2 of the observations (~68 % and ~81 % of the values, respectively). As shown by Tsyro (2005, and references therein), this underestimation is a known issue present in many air quality models. The water content of the aerosol at 50 % relative humidity (required when using the gravimetric

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measurement method) could explain the model underestimation, accounting for 20–35 % of the total aerosol mass, depending on the composition. Additionally, the coarse resolution of the model, which cannot reproduce local effects on fine aerosols, could also contribute to the underestimation. Nevertheless we obtain (see Fig. 2) very similar patterns (although low in absolute values) compared to those derived by van Donkelaar et al. (2010), which estimated the PM_{2.5} mass at 50 % relative humidity from satellite observations.

In the case of SO₂, the modelled mixing ratios are associated with larger discrepancies. For the EPA network, the model reproduces the observed values within a factor of two in more than 71 % of the locations, though with some small underestimations. For Europe and East Asia, the observations are more scattered, i.e. with larger variance (not shown). In Europe the model tends to overestimate the mean observed values by almost a factor of two. The SO₂ emissions of the EDGAR-CIRCE dataset might overestimate the sulfur content of the European coal and are also higher than the officially reported EMEP (European Monitoring and Evaluation Programme Vestreng et al., 2007, 2009) emissions (Pozzer et al., 2012). In Asia, the modelled averages agree well with the observations, though this is merely due to balancing of extremes: in fact, only ~32 % of the modelled values lies within a factor of two to the observations. As shown by Zhao et al. (2012), the EDGAR-CIRCE emissions present generally higher SO₂ emissions compared to other datasets, with strong differences on regional or smaller scales. Finally, the low number of observations for the EANET network could influence the robustness of the results, which could be statistically weak as based only on 35 points/stations.

Despite the reasonable agreement between model results and observations for O₃, PM_{2.5} and SO₂, for CO and NO₂ significant discrepancies with the observational datasets are apparent. The model tends to underestimate the observed values by up to a factor of two to three for NO₂ and CO, respectively. These results for CO with the EMAC model seems to be in contrast with previous findings (Jöckel et al., 2006; Pozzer et al., 2007), which showed much better agreement with the observations.

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To further investigate these discrepancies, an additional observational dataset has been used: the CO observations from the NOAA ESRL GMD (National Oceanic and Atmospheric Administration, Earth System Research Laboratory, Global Monitoring Division, Boulder, CO, USA) cooperative air sampling network (Novelli et al., 1998). The same dataset was used in the work of Jöckel et al. (2006) and Pozzer et al. (2007), and these observations were collected for non-polluted conditions avoiding local contamination (Haas-Laursen and Hartley, 1997). Compared to these observational data, the simulated annual CO differs by maximum 40 %; and for ~90 % of the locations the differences are within 30 % (not shown). Finally, considering only locations of the NOAA ESRL GMD dataset outside Antarctica, all modelled calculated annual means are within 20 % of the observations. These results are very similar to what obtained by Pozzer et al. (2007). It can be hence concluded that, if the same observational dataset is used (NOAA ESRL GMD), the agreement between model results and observations is comparable to what obtained by Pozzer et al. (2007). This agreement is much better than what suggested by the comparison of model results with the EPA and EEA observations. Apparently, the total CO source in the model is quite realistic and the concentration of OH, the main sink of CO, is also well reproduced globally, as described in Sect. 3.3, and it is very similar to what was obtained by (Jöckel et al., 2006).

To further investigate if the global CO distribution is well reproduced, an additional comparison for CO near the surface was performed using satellite data. Satellite observations of CO are provided by the MOPITT (Measurements Of the Pollution In The Troposphere) instrument (Drummond and Mand, 1996), which flies on the Terra platform. The MOPITT dataset has been extensively evaluated and used in many studies (e.g. Emmons et al., 2004, 2009; Pfister et al., 2005; Liu et al., 2011). Here we used the level 3 (with monthly temporal resolution) version 4 (V92.0.1) of the MOPITT product, with a spatial resolution of $1 \times 1^\circ$. It should be mentioned that the MOPITT product is not only derived from direct observations but also contains modelled information (see Deeter et al., 2003, 2004, for information regarding the MOPITT CO

retrieval algorithm). Nevertheless, its use can provide additional insight into the EMAC representation of CO, having a similar spatial resolution. In Fig. 3, the annual mean modelled and remotely sensed near-surface CO are compared. The appropriate averaging kernels of the satellite retrieval algorithm have been applied to the model output to enable a meaningful comparison. Generally the model results are within 30% of the satellite observations. Some small overestimations are present in the tropics and over highly elevated topography, due to a model overestimation of CO mixing ratios in the free troposphere. Relatively small model underestimations occur in the extratropics. In particular over the USA the model underestimates the satellite observed CO by 15%, well below that suggested by the comparison of the model results with the EPA data. Such small differences also occur over Europe (generally below ~10%), East Asia (below ~6%), while larger differences occur over the Sahara and the Middle East (~15%). Hence the model tends to underestimate the CO concentrations, although much less (below ~15%) than suggested by the comparison with the EPA and EEA observations. The general underestimation also appears by the total CO emissions (~1052 Tg yr⁻¹), which is ~20% lower than that suggested by Hooghiemstra et al. (2011), based on inverse modeling.

Overall the CO source strength is reasonably represented: when compared to MO-PITT, which has a similar resolution as the simulations used in this work, the EMAC model is able to reproduce the observations consistently or with a minor underestimation. Hence this analysis suggests that the CO observations in the EPA, EEA and EANET networks are strongly influenced by local pollution, which is difficult to accurately simulate by the EMAC model at T106 resolution. The observational networks used here often include stations in locations with heavy traffic or industry. For example ~25% (160 Tg yr⁻²) of the total CO emissions are due to fossil fuel combustion by traffic. Similar findings were obtained by Qian et al. (2010), who showed that CO presents a large sub-grid variability and it is strongly influenced by local source.

NO₂ is also generally underestimated by the model (note however the low number of stations in the EANET network) while there are no indications that the emissions are

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equally incorrect. We attribute the discrepancy to underrated direct emissions of NO_2 (all NO_x is emitted as NO) and possibly a too slow conversion of NO into NO_2 by peroxy radicals or other oxidants in polluted air. Furthermore, a recent study by Miyazaki et al. (2012), based on model inversion of satellite data, suggests that anthropogenic NO_x emissions are generally underestimated in inventories. The model probably underestimates the photochemistry of reactive hydrocarbons near pollutant sources, e.g. because it does not account for aromatics. Nevertheless, what is most relevant for air quality is the concentration of total oxidant ($\text{NO}_2 + \text{O}_3$), which is well represented. In Table 1, the comparison with the total oxidant O_x ($\text{NO}_2 + \text{O}_3$) is shown, though only for the EPA and EEA networks, because EANET does not include a sufficient number of stations to derive meaningful statistics. The model results show very good agreement with both the EPA and EANET measurements, as more than 99 % and 56 % of the modelled values are within a factor of two and 0.2, respectively, of the observations, implying that the total oxidant is well represented in the model.

3.2 Future projections

3.2.1 SO_2

In Fig. 4 the annual mean of near-surface sulphur dioxide is shown for the different simulations. As expected, the SO_2 mixing ratios in background locations from simulation *SC_natural* are below 1 nmol mol^{-1} , since SO_2 is only emitted by relatively small natural sources (e.g. volcanoes are typically on land and release 29 Tg yr^{-1}) and SO_2 in the marine atmosphere is formed by the oxidation of dimethyl sulphide (DMS). The annual mean near-surface mixing ratios from other simulations are remarkably different, as the anthropogenic emissions increase dramatically in simulations *SC_2005*, *SC_2010*, *SC_2025* and *SC_2050*. As shown in Fig. 4, including the anthropogenic emissions drastically increases the near-surface mixing ratios with time. This is also illustrated by Table 2 in which the atmospheric burdens of selected tracers are listed. While SO_2 sources in simulation *SC_natural* lead to a burden of 0.3 Tg , in simulation *SC_2005* the

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burden strongly increases by a factor of 3 (0.9 Tg). The projected increase for the year 2050 (from simulation *SC_2050*) is around 1.8 Tg, which is six times the burden in the atmosphere estimated in the simulation *SC_natural*. A doubling of the burden of SO₂ is projected between the years 2005 and 2050, consistent with the increase of the anthropogenic emissions. It is clear from Fig. 4 that China has become the major emitter of SO₂ since 2005, and it is expected to remain so up to 2050, also shown in previous studies (Pozzer et al., 2012; Lu et al., 2010, 2011). In 2050 the densely populated northern part of India will also be exposed to very high SO₂ mixing ratios. Interestingly, the model results suggest that toward 2050 India may have SO₂ mixing ratios comparable to those in eastern China since 2005, while in simulation *SC_2025*, apart from some local hot spots, the large scale SO₂ mixing ratios are still an order of magnitude lower than in eastern China. Analogously, increasing emissions in some countries in eastern Europe and southern Russia are projected to lead to high levels of SO₂, and comparably also some locations in central Asia. The SO₂ is not uniformly distributed on large scales, due to its short lifetime (few days). Hence, local mixing ratios can vary tremendously. Generally, SO₂ reaches high levels in urban locations in Europe, eastern USA, South America and South Africa in all simulations with anthropogenic emissions included.

It must be stressed that the increase of the burden does not directly translate in a direct increase of the maximum of SO₂ levels. The model calculates local annual mean maxima in hot spot locations of about 24, 86, 91, 89, and 88 nmol mol⁻¹ for *SC_natural*, *SC_2005*, *SC_2010*, *SC_2025* and *SC_2050*, respectively. The large increases of SO₂ mixing ratios are associated with the rapidly growing emissions in countries with emerging economies (e.g. India, China), while sources in other nations that strongly emitted in 2005 tend to not contribute to future increases of the total SO₂ burden (but rather decrease).

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3.2.2 O₃

Different from SO₂, O₃ is more homogeneously distributed in accord with its atmospheric lifetime of several weeks (see Fig. 5), with global annual mean hot spot maxima of 41, 78, 77, 84 and 95 nmol mol⁻¹ for simulations *SC_natural*, *SC_2005*, *SC_2010*, *SC_2025* and *SC_2050*, respectively. Similar to SO₂, northern India is progressively becoming strongly influenced by high O₃ mixing ratios, with a continual increase up to 2050. The already high O₃ mixing ratios over the Arabian Gulf in simulation *SC_2005* are projected to drastically increase further in future, with annual average mixing ratios higher than 90 nmol mol⁻¹ in the year 2050 with an increase of the annual average equal to 70 nmol mol⁻¹ with respect to simulation *SC_natural*. The high levels of ozone in this region are caused by a generally high background by atmospheric transport, strong insolation, a shallow boundary layer over the Gulf and large local precursor emissions (Lelieveld et al., 2009; Smoydzin et al., 2012). Unless the anticipated emission trend will be reversed, the Arabian Gulf is destined to become a hot spot of poor air quality due to the high ozone levels. Interestingly, China is not expected to be subjected to very high levels of ozone, but instead mean mixing ratios appear rather constant up to the year 2050, due to tritiation of O₃ (see Sect. 3.2.3), and lower concentrations of OH due to the large increase of CO. Also in most of Europe and eastern North America ozone mixing ratios are not expected to change drastically, although the projected local precursor emission decreases can be counteracted by a hemispheric increase of background values (Lelieveld and Dentener, 2000). Overall, the tropospheric burden of ozone is calculated to increase by 15 % between 2005 to 2050, from 387 to 442 Tg (Table 2).

3.2.3 NO₂

As shown in Fig. 6, the increasing ship traffic noticeably affects the NO₂ distributions in the different simulations. According to *SC_2005* and *SC_2050* air traffic emissions also increase substantially in this period, from ~0.9 to ~1.4 Tg(N) yr⁻¹, while the ship

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emissions are expected to increase from ~ 4 to ~ 11 Tg(N) yr⁻¹. Different from SO₂, in Europe and the eastern USA we find similar NO₂ mixing ratios as in India. China is expected to become the most prominent NO_x emitter and in future the near-surface mixing ratios of NO₂ are projected to be the highest in the world, which contributes to ozone formation downwind. For example, over the central Pacific Ocean the annual mean near-surface mixing ratio of O₃ is calculated to increase from less than 15 to nearly 50 nmol mol⁻¹ from *SC_natural* to *SC_2050*. The resultant ozone levels in China, close to the NO_x sources, may not increase very strongly after 2005 (see Sect. 3.2.2) since the growing NO sources also cause a local titration of O₃.

3.2.4 CO

The atmospheric carbon monoxide burden almost doubles from simulation *SC_natural* to simulation *SC_2005*, increasing from ~ 200 to 393 Tg. Results from simulation *SC_2050* indicate that the burden projected for the year 2050 will be 484 Tg, suggesting a $\sim 23\%$ increase between 2005 and 2050. As shown in Fig. 7, the increasing emissions from *SC_natural* and *SC_2005* influence the atmosphere globally, as the relatively long lifetime of CO (several months) allows the efficient transport on large scales. While the peak increases in CO concentrations in *SC_2050* relative to *SC_2005* are located in the poor air quality hot spots of southern and eastern Asia, large effects are also found over remote regions, with substantially increasing mixing ratios over the northern Indian Ocean, the central Pacific Ocean, the central Atlantic Ocean and Siberia. The typical large scale contrast that appears between source regions and the background atmosphere in *SC_natural* decreases in *SC_2005*. This is a result of the global anthropogenic CO emissions which are not in phase with the biogenic and biomass burning emissions (see also the seasonal variability of the burden in Table 2). Hence the anthropogenic emissions damp the annual cycle and the horizontal contrast in CO mixing ratios. In the year 2050 the anthropogenic emissions are projected to substantially

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exceed the natural ones, so that the amplitude of the annual cycle (see Table 2) forced by the anthropogenic emissions is comparable to that in *SC_natural*.

3.3 Hydroxyl radical (OH)

The hydroxyl radical is the main oxidant in the troposphere and its average concentration determines the “cleansing capacity” of the atmosphere. In Fig. 8, the annual average of the OH radical concentration at the surface is shown for the different simulations performed. The highest concentrations are found in the tropics, as expected (Prinn et al., 1992; Spivakovsky et al., 2000). In all simulations the highest values are found over the tropical oceans. In this environment primary OH production is facilitated by the strong insolation together with the high abundance of water vapor. Over the tropical forest the OH present a minimum due to the strong emission of non methane volatile organic compounds (NMVOC), which deplete the available OH. This is in contrast with the recent study of Taraborrelli et al. (2012), which shows that the OH recycling over tropical forest is very efficient, a process which has not been incorporated in this study.

In the simulations with anthropogenic emissions included, the highest annual average OH concentrations (up to 6×10^6 mcl cm⁻³) are found off the coast of South and East Asia, over the Gulf of Mexico and the Red Sea. The ship tracks are very pronounced with high OH levels, due to the localized NO_x emissions in the otherwise relatively unpolluted environment, which enhance the recycling of OH (Lawrence and Crutzen, 1999; Lelieveld et al., 2002). Despite the high OH in ship tracks, in more remote regions over the tropical oceans OH levels have reduced, mostly due to the increasing methane. This is confirmed by the global average OH concentrations for the different simulations, presented in Table 3. Global OH is projected to reduce according to future scenarios, although not very significantly (Krol et al., 1998; Prinn et al., 2001). In general, OH concentrations over the tropical oceans respond relatively sensitively to anthropogenic perturbations (Lelieveld et al., 2002). Although the marine locations of relatively high OH in simulation *SC_natural* are still present in the simulations for 2005 up to 2050, the land-ocean contrasts are reduced due to enhanced OH recycling by

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NO_x in continental locations. Global mean OH changes in future will be dependent on the trade-off between enhanced OH recycling by NO_x, increased O₃ formation, and OH reductions caused by growing concentrations of reactive carbon (CH₄, CO, NMVOC). The relatively low latitude location of India, combined with relatively high NO_x and O₃ levels, may cause this country to become a continental OH hot spot in future (see Fig. 8).

PM_{2.5}

PM_{2.5} is very important for air quality, and its influence on human health is generally recognized (Brunekreef and Holgate, 2002, and reference therein). Even low levels of exposure have effects on the lungs and the cardiovascular system (Dockery, 2001). It must be stressed that PM_{2.5} is not only formed by bulk aerosol species, but also by the formation of secondary aerosol (Orel and Seinfeld, 1977) especially where direct emissions of precursors are very high. As shown in the Supplement, the anthropogenic emitted BC is projected to increase of only ~25 %, while SO₂ and NO_x emissions are projected to increase both of a factor of two.

As shown in Fig. 9 the largest changes of PM_{2.5} between simulation *SC_natural* and *SC_2005* are located in Europe, North East USA and particularly in southern and East Asia. In future scenarios (*SC_2025* and *SC_2050*), the increase of PM_{2.5} is larger in East and South Asia, associated with the strong increase of aerosol and precursors emissions.

4 Multi pollutant index

4.1 General definition

As we are focusing on multiple gases/aerosols which are harmful to human health, it is useful to combine these into a single index of air quality. Different indexes are

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present in the literature, and a full review of air quality indexes can be found in Plaia and Ruggieri (2011). We selected the multipollutant index (MPI) as defined by Gurjar et al. (2008) because it was tested for the representation of long term air quality (Gurjar et al., 2008), and hence appropriate to be used with annual average data. It must be stressed that the MPI is a linear index of air quality related to health and does not represent the effects of air quality on human health, which is strongly non linear (determined by the exposure time), and dependent on the different pollutants.

Following the approach of Gurjar et al. (2008), a MPI can be defined as:

$$\text{MPI} = (1/n) \left[\sum (\text{AC}_i - \text{GC}_i) / \text{GC}_i \right]$$

where n is the total number of pollutants taken in consideration, AC_i is the atmospheric concentration of a pollutant in a certain location and GC_i is the guideline concentration of a pollutant recommended by a national or international agency such as the WHO (World Health Organization – Regional Office for Europe, 1987, 2000; World Health Organization, 2005). As shown by Gurjar et al. (2008), a clean environment (in the absence of air pollutants) has an MPI equal to -1 , while for a location with AC_i equal to the GC_i (i.e. a location with borderline air quality), the MPI would be around 0. No upper limit is given for the MPI, with relatively higher values corresponding to poorer air quality. We refer to Gurjar et al. (2008) for a detailed discussion on such index.

As mentioned in Sect. 3, in this work only the annual average values are taken into account. Different recommendations and air quality standards for annual averages have been published in the past decades (i.e. World Health Organization – Regional Office for Europe, 1987, 2000; World Health Organization, 2005), reflecting the evolving understanding about the impact of long term exposure to these compounds on human health (Krzyzanowski and Cohen, 2008; HTAP, 2010). For example, the WHO recently abandoned the guideline limit value for particulate matter, reflecting that there is no “safe” lower threshold for this pollutant. Notwithstanding this discussion and the specific interpretation of the WHO air quality standard on regional and national levels, we adopted the limits as defined by Gurjar et al. (2008) for SO_2 (17 nmol mol^{-1}) and NO_2

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(20 nmol mol⁻¹), for consistency with his work, while for PM_{2.5}, the lowest level suggested by the WHO (10 μm⁻³) is used. For O₃ and CO no annual recommendations are generally given. For these compounds we linearly scaled the recommendation from hourly to annual, as for NO₂ (i.e. one fifth from 1-h-mean to annual mean) of the WHO recommendation (World Health Organization, 2005) for CO (i.e. 7000 nmol mol⁻¹) and the EPA recommendation EPA (2005) for O₃ (i.e. 24 nmol mol⁻¹). As mentioned before, although these values cannot quantitatively reflect the health risk associated with the exposure to mixtures of pollutants (which depends strongly on the compounds selected), we believe it provides an objective metric to compare air pollution in different regions, based on our model output. Additionally, following the discussions in Sect. 3, we did not use NO₂ or O₃, but rather the combination of the two gas tracers (O_x), which is well represented in the model. The air quality limits thus used are 10 μm⁻³ for PM_{2.5}, 17, 44 and 7000 nmol mol⁻¹ for SO₂, O_x and CO, respectively. It must also be stressed that the MPI values are strongly influenced by the value used for CG_{*i*} (recommended concentrations). In this work, the PM_{2.5} has the strongest influence on the overall MPI value, while CO has the least influence, due to its annual value generally being much lower than 7000 nmol mol⁻¹. The MPI definition has hence to be used carefully with respect to the absolute values while the comparison of locations over time and space is more robust.

In Fig. 10, the MPI values for the preindustrial case and for the year 2005 are presented. Not surprisingly, locations that are strongly influenced by desert dust have a very high MPI (i.e. low air quality), while even the pristine marine environment has a reduced (though not poor) air quality due to sea spray. The desert dust areas include northern Africa, the Arabian Peninsula, the region east of the Arabian Gulf, northern China (influenced by the Gobi desert) and central Australia. The influence of anthropogenic emissions is apparent in simulation *SC_2005*, especially over China and northern India, where the strong emissions of pollutants bring the annual MPI close to 1.0.

In Figs. 11 and 12 the differences between the MPIs from the four recent and future simulations (*SC_2005*, *SC_2010*, *SC_2025*, *SC_2050*) and the MPI obtained for

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the “preindustrial case” (*SC_natural*) are presented. In the year 2005, five or six main regions of decreased air quality due to anthropogenic emissions emerge: North America (east and west coast), Europe and the Mediterranean Basin, the Middle East, India and eastern China. In recent years, i.e. between 2005 and 2010, air quality noticeably degraded in the latter regions, especially in China.

In the projections for the future periods several regions with very high MPIs become apparent, mainly by the expansion of the present ones. Anthropogenic emissions will continue to degrade air quality in eastern China in 2025 with an MPI increase in excess of a one full point in the northeast, while northern India is also emerging with high MPIs. The northeastern USA, Central East Europe and the Middle East also appear as high MPI regions, though with lower MPI values than the Asian regions. The increase of MPI due to anthropogenic emissions in the Middle East and North Africa has to be added to the already high MPI by natural causes present in these regions, leading to very high MPIs. The main cause of the increase of the MPI in the Middle East is the formation of photochemical smog with high ozone levels. In contrast, in India and China particulate matter and SO₂ play an important role.

The large scale influence of these emerging anthropogenic air-pollution hot spots is shown in Fig. 12, where an intercontinental “connections” are formed. The increased MPI regions (> 0.2 increase with respect to the natural conditions) extends across the entire Northern Hemisphere especially in the middle and subtropical latitudes, including the USA and the Mediterranean Basin. In India the MPI is additionally enhanced by growing O₃ and PM_{2.5} pollution, while in China the high levels of NO₂, SO₂ and PM_{2.5} contribute substantially.

4.2 Per Capita MPI

4.2.1 Definition

Although the MPI helps in defining the degradation of air quality in polluted regions, it does not account for the numbers of people subjected to these conditions. Typically,

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polluted regions are also densely populated, in particular in the urban environment. Following the approach introduced in the previous section, we defined the Per-Capita Multi Pollutant Index (PCMPI), thus extending the definition to include population density. We used the (normalised) human population estimate to define a weighting factor to the MPI in calculation of PCMPI:

$$\text{PCMPI} = \frac{1}{W} \sum_{i=1}^N w_i \text{MPI}_i$$

where w_i is the weighting factor (the population in a certain model grid), $W = \sum_{i=1}^N w_i$ is the normalisation factor (the total population in the area) and $i = 1 \dots N$ refers to the model grid box in the region of interest. According to this method, the index is not only proportional to the air quality level (with respect to the international recommendations) but also weighs the number of people that is subjected to it. One feature is that the PCMPI offers the possibility of assessing the air quality on a global scale, over large regions or of countries, as the index reflects the mean level air quality to which the average person is subjected. To calculate the weighting factors, we used the population estimates for the year 2005, 2010, 2025 and 2050 by the Center for International Earth Science Information Network (CIESIN). For the *SC_natural* simulation the weighting factors are based on the year 2005 population for reference. This population weighting is a standard approach, also used by Brauer et al. (2012) focusing on ozone and PM_{2.5}.

4.2.2 PCMPI by regions

In Table 4 the PCMPI index is presented globally and for seven regions of interest: North America, Europe, Middle East, East Asia, South Asia, Central Africa and South America (see Fig. 13 for the region boundaries).

The largest apparent changes in the PCMPI occur between simulation *SC_natural* and *SC_2005*, related to growing air pollution in the 19th and 20th centuries, which rather drastically impacted air quality for much of the world population.

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Although the PCMPI changes between *SC_2005* and *SC_2050*, may seem smaller than those up to 2005, they occur on a shorter time scale. For North and South America, Europe and Central Africa the estimated time normalized (i.e. decadal) air quality degradation is more-or-less gradual. For the economically emerging regions in South and East Asia and the Middle East the PCMPI increases in the 21st century are very drastic.

Globally, the PCMPI increased by ~80 % between *SC_natural* and *SC_2005*, and an additional factor of two rise is projected up to the year 2050. The increase of ~80 % is a lower limit, as we neglected the urbanisation from preindustrial to 2005, using the same population distribution in both cases. North America and Europe have developed (and will continue to develop) very similarly, presumably typical for countries that have rapidly industrialised in the 20th century. After a strong increase of the PCMPI since preindustrial conditions (by ~0.3–0.4), the projected changes up to the year 2050 will be much smaller (around 25 %) in these regions compared to the globally projected changes. Even though air quality will deteriorate further in North America and Europe, in 2050 it will be significantly better than that experienced by the global average citizen. Remarkably, the global PCMPI in 2050 is projected to be similar to that in the Middle East in the year 2005.

It should be emphasized that this result is relatively strongly influenced by the high population density in Asia (i.e. low air quality affecting a large number of people). In fact, East and South Asia, have both the highest MPI values (see Fig. 10) and the highest population densities. The PCMPI for East Asia has strongly increased from –0.71 to 0.30 between *SC_natural* and *SC_2005*, more strongly than the increases experienced by any other region. Interestingly, all regions except the Middle East have a similar PCMPI under non-anthropogenically polluted conditions (*SC_natural*), while in 2050 East and South Asia, closely followed by the Middle East, are projected to have much higher PCMPIs, up to 0.82 (Table 4). The projected changes in air quality in East and South Asia are so drastic (~0.5–0.7 difference between *SC_2050* and *SC_2005*) and the population densities so high that, the global PCMPI is strongly influenced.

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South Asia (i.e. India) is the region experiencing the strongest changes between 2005 and 2050 with an increase of 0.75, by far the largest of all regions. Impressively, this change is even larger than the MPI change between *SC_2005* and *SC_natural* (~ 0.6).

Finally, the Middle East is a hot spot of poor air quality, for which we calculated a MPI change between *SC_2005* and *SC_2050* equal to the MPI change between *SC_2005* and *SC_natural* (i.e. ~ 0.2). However, the high abundance of desert dust and the consequently low air quality since preindustrial times and the continuous industrialisation (mainly related to the petrol extraction and processing industry) will drastically deteriorate the atmospheric environmental conditions.

In Central Africa the MPI values are close to those in North America. This is due to the particular population distribution and the fact that some subregions are strongly influenced by desert dust. The relatively best air quality and the lowest PCMPIs are calculated for South America, although also in this region pollution conditions are gradually becoming worse. The projected PCMPI is estimated to remain lower compared to all other regions during the entire period 2005–2050.

4.2.3 PCMPI by countries

In Table 5 the PCMPI is presented for the 100 world countries with the highest PCMPI in 2005. The complete list is available in the Supplement. Countries with a high abundance of natural desert dust are ranked high in the list due to the strong influence of $PM_{2.5}$ in the PCMPI calculations. It is interesting to notice the changes over time of the PCMPI for different countries. As shown in Fig. 15, India is projected to have the most rapid degradation of air quality per capita, followed by Bangladesh and China. Figure 15 also shows many countries in the Middle East, which are shown to have poor air quality already in simulation *SC_2005*. Finally, although the PCMPI is projected to increase globally by ~ 0.31 , most countries will face a much lower increase. A very few densely populated countries such as China, India and Bangladesh, on the other hand, are expected to be affected by a much higher increase (almost twice as high).

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5 Conclusions

We used the global EMAC model to estimate the potential impact of anthropogenic emission changes on air quality in recent and future years (2005, 2010, 2025 and 2050), based on a “business as usual” scenario. No feedbacks between the changing atmospheric composition and dynamics were allowed in the GCM, so that the model simulated air quality is controlled by air pollution emissions rather than climate change, though the latter may be an additional important factor in the future atmospheric composition (Giorgi and Meleux, 2007; Jacob and Winner, 2009; Dawson et al., 2009).

We focussed on the five major pollutants that impact human health: PM_{2.5}, nitrogen dioxide (NO₂), sulphur dioxide (SO₂), ozone (O₃) and carbon monoxide (CO).

The model reproduces the main characteristics of recent air quality according to observations in North America, eastern Asia and Europe, although with some imprecisions inherent to the coarse resolution of the model. Evaluation of the model results indicates that near the surface the model underestimates CO and NO₂ mixing ratios. On the other hand, SO₂ is slightly overestimated over Europe and Asia and underestimated over the USA, though the annual means are within 40 % of the observations.

Both O₃ and PM_{2.5} are simulated in reasonably good agreement with observations. While NO₂ is underestimated, the calculated total oxidant O_x, (i.e. the sum of O₃ and NO₂) agrees well with observations with more than 56 % of the model results being within 20 % of the observations and 100 % within a factor of two.

The simulations for the year 2025 and 2050 suggest that East Asia will be exposed to high levels of a range of pollutants (SO₂, NO₂, PM_{2.5}), though ozone may not increase very strongly. In contrast, in northern India and the Arabian Gulf region ozone levels may grow rapidly. To quantify the possible future air quality hot spots, a multi pollutant index (MPI) was introduced. This index, applied to the model projections, point to three main hot spots of poorest air quality in the future, located in eastern China, northern India and the Middle East/North Africa.

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By extending the MPI definition with population data, a Per-Capita MPI (PCMPI) has been derived. The PCMPI calculations indicate that global air quality has decreased significantly since pre-industrial times, which is projected to continue in future. In 2050 most of the world population will be subjected to degraded air quality. Especially the densely populated parts of South and East Asia contribute strongly to the loss of air quality for the average global citizen. The air quality (PCMPI) in 2025 and 2050 is projected to be worst in the Middle East and North Africa, due to a combination of natural desert dust and anthropogenic ozone, followed by East and South Asia where anthropogenic pollution emissions will be particularly high.

It must be stressed that the results obtained in this work are based on a business as usual scenario. This rather pessimistic though feasible scenario represents a future where no additional climate change and air pollution reduction policy will be implemented using the year 2005 as starting point. The results of this work indicate that strong actions and further legislations are essential to avoid the drastic deterioration of air quality which can have severe effects on human health.

Supplementary material related to this article is available online at:

<http://www.atmos-chem-phys-discuss.net/12/8617/2012/acpd-12-8617-2012-supplement.pdf>.

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Table 1. Summary of the comparison of model data with observations. N.stat. is the number of locations/stations used in the comparison. OAM and MAM are the arithmetic mean of the observations and of the model, respectively. O_3 , SO_2 , NO_2 , CO and O_x are in $nmol\ mol^{-1}$, while $PM_{2.5}$ is in $\mu g\ m^{-3}$. The model has been sampled at the locations of the observations, and the calculations are based on annual averages. PF2 is the percentage of model results within a factor of two of the observations and PF0.2 is the percentage of model results within 20% of the observations. O_x represents $NO_2 + O_3$.

species	network	All stations					
		N.stat.	MAM	OAM	MAM/OAM	PF2	PF0.2
O_3	EPA	1253	41.6	52.6	0.8	99.9	44.4
O_3	EEA	2039	36.1	24.9	1.4	86.3	21.4
O_3	EANET	17	41.4	36.4	1.1	82.3	58.8
SO_2	EPA	527	2.7	3.8	0.7	71.9	24.1
SO_2	EEA	2285	4.3	2.5	1.7	40.1	11.9
SO_2	EANET	35	3.4	3.1	1.1	31.4	17.2
NO_2	EPA	407	6.5	11.9	0.5	53.8	17.2
NO_2	EEA	2793	6.7	15.0	0.4	43.8	11.0
NO_2	EANET	5	5.4	15.4	0.3	20.0	20.0
CO	EPA	452	170.5	545.0	0.3	25.7	0.2
CO	EEA	1280	184.6	563.4	0.3	36.8	8.4
O_x	EPA	364	49.2	62.4	0.8	100.0	56.4
O_x	EEA	1930	42.5	37.5	1.1	99.3	68.1
$PM_{2.5}$	EPA	1245	8.7	12.3	0.7	68.1	17.4
$PM_{2.5}$	EEA	275	11.1	17.9	0.6	81.1	34.3
$PM_{2.5}$	EANET	2	8.3	11.6	0.7	100.0	50.0

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Table 2. Global burdens of CO, SO₂, NO₂ and O₃ in Tg. The yearly standard deviations are listed within parentheses. O₃ represents the tropospheric ozone, only integrated from the surface to the tropopause.

species	<i>SC_natural</i>		<i>SC_2005</i>		<i>SC_2010</i>		<i>SC_2025</i>		<i>SC_2050</i>	
CO	200.6	(22.8)	393.3	(18.9)	400.4	(19.7)	446.4	(21.4)	484.7	(22.9)
SO ₂	0.3	(0.0)	0.9	(0.2)	1.0	(0.3)	1.3	(0.4)	1.8	(0.6)
NO ₂	1.9	(0.1)	2.1	(0.1)	2.3	(0.1)	2.3	(0.1)	2.4	(0.1)
O ₃	273.4	(12.6)	387.7	(18.3)	390.5	(18.2)	416.2	(19.6)	442.4	(20.8)

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Table 3. Global average tropospheric OH concentration in 10^6 cm^3 , estimated with different weightings (mass, volume, methyl chloroform reaction and methane reaction) factors for the different simulations presented in this work. S2000 is the same calculation from the work of Spivakovsky et al. (2000).

method	S2000	<i>SC_natural</i>	<i>SC_2005</i>	<i>SC_2010</i>	<i>SC_2025</i>	<i>SC_2050</i>
mass	1.14	1.16	1.07	1.07	1.04	1.04
volume	1.10	1.19	1.07	1.07	1.04	1.03
k_{MCF}	1.29	1.30	1.23	1.24	1.21	1.21
k_{CH_4}	1.32	1.28	1.21	1.22	1.19	1.18

Table 5. PCMPI values for different counties (ranked according to the PCMPI in *SC_2005*, top 100 countries).

country	<i>SC_natural</i>	<i>SC_2005</i>	<i>SC_2010</i>	<i>SC_2025</i>	<i>SC_2050</i>	country	<i>SC_natural</i>	<i>SC_2005</i>	<i>SC_2010</i>	<i>SC_2025</i>	<i>SC_2050</i>
Niger	1.38	1.51	1.52	1.55	1.57	Yemen	-0.41	-0.23	-0.22	-0.19	-0.12
Mauritania	1.21	1.30	1.29	1.30	1.33	Uzbekistan	-0.50	-0.26	-0.24	-0.21	-0.16
Chad	1.12	1.21	1.21	1.23	1.25	Eritrea	-0.40	-0.26	-0.25	-0.22	-0.17
Egypt	0.67	0.94	0.95	1.01	1.11	Netherlands	-0.83	-0.27	-0.27	-0.25	-0.21
Iraq	0.44	0.70	0.70	0.73	0.82	Ivory Coast	-0.35	-0.28	-0.28	-0.26	-0.24
Mali	0.54	0.62	0.61	0.63	0.65	Mongolia	-0.43	-0.30	-0.30	-0.27	-0.23
Libyan Arab Jamahiriya	0.42	0.61	0.62	0.63	0.67	Myanmar	-0.70	-0.34	-0.32	-0.24	-0.15
Kuwait	0.21	0.57	0.57	0.60	0.72	Greece	-0.66	-0.35	-0.34	-0.33	-0.27
Saudi Arabia	0.34	0.56	0.56	0.59	0.66	Bulgaria	-0.73	-0.35	-0.32	-0.33	-0.24
Nigeria	0.20	0.52	0.52	0.58	0.59	Dem. Rep. Congo	-0.45	-0.36	-0.36	-0.35	-0.34
Sudan	0.35	0.46	0.46	0.48	0.52	Chile	-0.79	-0.38	-0.35	-0.32	-0.29
United Arab Emirates	0.16	0.44	0.44	0.49	0.59	Tajikistan	-0.60	-0.38	-0.37	-0.34	-0.29
Bahrain	0.04	0.40	0.40	0.44	0.56	Cambodia	-0.68	-0.38	-0.37	-0.32	-0.30
Burkina Faso	0.30	0.40	0.41	0.43	0.45	Kyrgyz. Republic	-0.56	-0.38	-0.37	-0.34	-0.29
Senegal	0.29	0.39	0.39	0.42	0.45	Cyprus	-0.62	-0.38	-0.37	-0.32	-0.25
China	-0.69	0.37	0.47	0.71	0.92	Czech Republic	-0.83	-0.39	-0.38	-0.37	-0.34
Qatar	0.01	0.36	0.35	0.40	0.51	Italy	-0.74	-0.39	-0.38	-0.37	-0.33
Pakistan	-0.28	0.29	0.29	0.40	0.53	Gibraltar	-0.67	-0.40	-0.39	-0.36	-0.31
Gambia	0.16	0.25	0.25	0.27	0.30	Turkey	-0.67	-0.40	-0.38	-0.32	-0.24
Syrian Arab Republic	-0.05	0.19	0.21	0.27	0.37	Albania	-0.69	-0.40	-0.38	-0.38	-0.32
Jordan	-0.15	0.18	0.20	0.30	0.47	Poland	-0.84	-0.40	-0.38	-0.38	-0.36
Guinea-Bissau	0.08	0.16	0.17	0.18	0.21	Indonesia	-0.75	-0.40	-0.40	-0.32	-0.28
Cameroon	0.04	0.16	0.16	0.19	0.20	San Marino	-0.77	-0.40	-0.39	-0.38	-0.34
Palestinian Terr.	-0.20	0.13	0.16	0.27	0.46	Liberia	-0.48	-0.42	-0.41	-0.39	-0.38
Korea	-0.76	0.11	0.18	0.33	0.46	Taiwan	-0.80	-0.42	-0.37	-0.27	-0.17
Turkmenistan	-0.05	0.08	0.09	0.11	0.15	Azerbaijan	-0.61	-0.42	-0.42	-0.38	-0.32
Macao	-0.79	0.07	0.15	0.35	0.55	Slovenia	-0.79	-0.42	-0.40	-0.39	-0.35
Israel	-0.29	0.05	0.07	0.19	0.38	Macedonia	-0.72	-0.42	-0.40	-0.40	-0.34
Hong Kong	-0.79	0.02	0.10	0.28	0.45	Germany	-0.84	-0.42	-0.42	-0.40	-0.37
Algeria	-0.19	0.02	0.03	0.06	0.09	Thailand	-0.76	-0.42	-0.41	-0.32	-0.26
Guinea	-0.06	0.01	0.01	0.03	0.05	Japan	-0.79	-0.42	-0.41	-0.38	-0.35
Benin	-0.15	-0.01	-0.00	0.01	0.03	Slovakia	-0.81	-0.43	-0.40	-0.39	-0.36
Cape Verde	-0.13	-0.01	-0.00	0.01	0.04	Kazakhstan	-0.62	-0.43	-0.41	-0.38	-0.33
Togo	-0.13	-0.02	-0.02	0.00	0.02	Spain	-0.75	-0.43	-0.43	-0.41	-0.37
Oman	-0.23	-0.03	-0.03	0.00	0.07	Croatia	-0.77	-0.44	-0.42	-0.41	-0.36
Central African Republic	-0.11	-0.06	-0.06	-0.05	-0.03	Hungary	-0.81	-0.45	-0.42	-0.41	-0.37
Bangladesh	-0.75	-0.08	-0.05	0.14	0.64	Peop. Dem. Rep. Lao	-0.74	-0.45	-0.43	-0.36	-0.29
Iran	-0.30	-0.08	-0.07	-0.04	0.05	Austria	-0.82	-0.45	-0.44	-0.42	-0.39
India	-0.67	-0.08	-0.04	0.15	0.79	Serbia and Montenegro	-0.75	-0.45	-0.40	-0.40	-0.34
Lebanon	-0.39	-0.09	-0.08	-0.01	0.11	France	-0.82	-0.45	-0.45	-0.43	-0.39
Tunisia	-0.37	-0.10	-0.09	-0.06	-0.02	Bhutan	-0.76	-0.45	-0.44	-0.35	-0.11
Ghana	-0.22	-0.12	-0.12	-0.09	-0.07	Jersey	-0.78	-0.46	-0.45	-0.42	-0.37
Dem. Peop. Rep. of Korea	-0.74	-0.12	-0.07	0.06	0.20	Angola	-0.52	-0.46	-0.46	-0.44	-0.44
Sierra Leone	-0.24	-0.16	-0.16	-0.14	-0.12	Guernsey	-0.77	-0.46	-0.45	-0.42	-0.38
Viet Nam	-0.80	-0.17	-0.15	-0.00	0.08	Romania	-0.78	-0.46	-0.43	-0.43	-0.35
Belgium	-0.84	-0.19	-0.18	-0.16	-0.12	Uganda	-0.65	-0.46	-0.46	-0.43	-0.39
Nepal	-0.70	-0.20	-0.18	-0.05	0.27	Bosnia-Herzegovina	-0.75	-0.46	-0.44	-0.43	-0.38
Malta	-0.49	-0.22	-0.21	-0.19	-0.14	Andorra	-0.79	-0.46	-0.46	-0.44	-0.40
Afghanistan	-0.41	-0.22	-0.22	-0.18	-0.15	Luxembourg	-0.85	-0.47	-0.47	-0.45	-0.42
Marocco	-0.43	-0.22	-0.22	-0.18	-0.12	Armenia	-0.67	-0.47	-0.47	-0.44	-0.38

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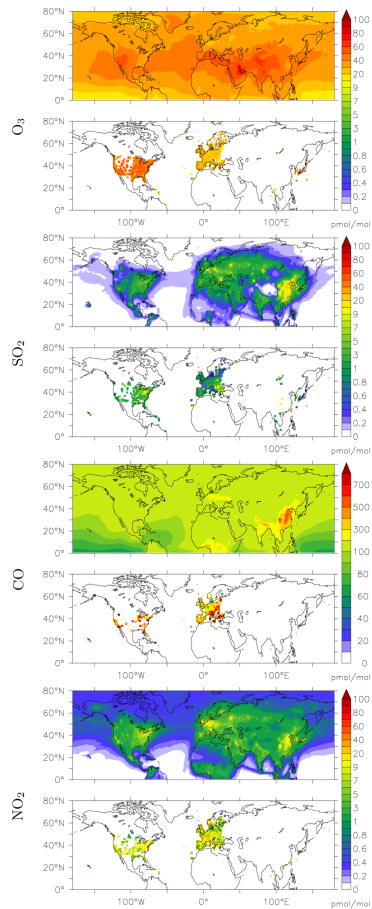


Fig. 1. UPPER PANELS: simulated annual mean mixing ratios in pmol mol^{-1} for the year 2005. LOWER PANELS: annual mean in pmol mol^{-1} from the EPA, EEA and EANET network observations for the year 2005. The trace gases (from top to bottom) are NO_2 , SO_2 , CO and O_3 .

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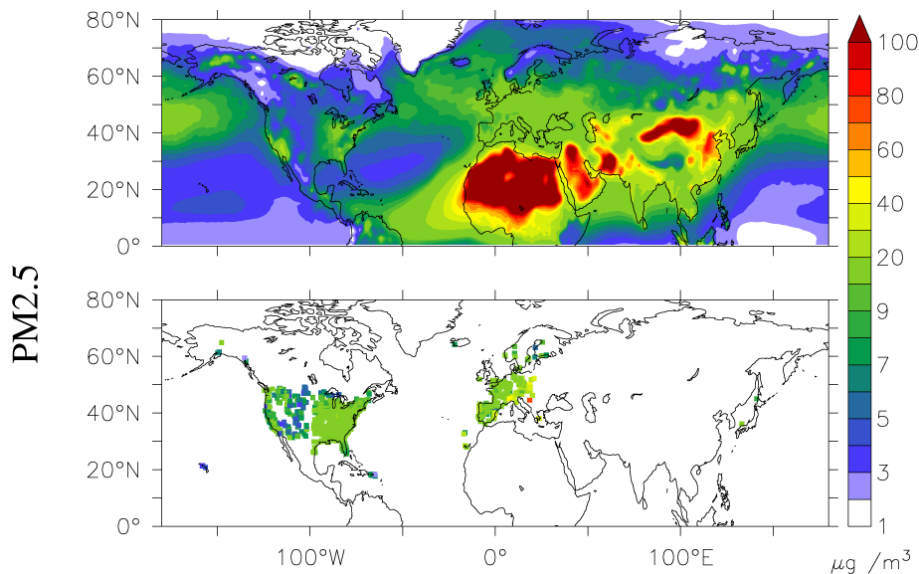


Fig. 2. UPPER PANEL: simulated annual concentrations of PM_{2.5} in $\mu\text{g m}^{-3}$ for the year 2005. LOWER PANEL: annual mean in $\mu\text{g m}^{-3}$ of observed PM_{2.5} for the year 2005 from the EPA, EEA and EANET network.

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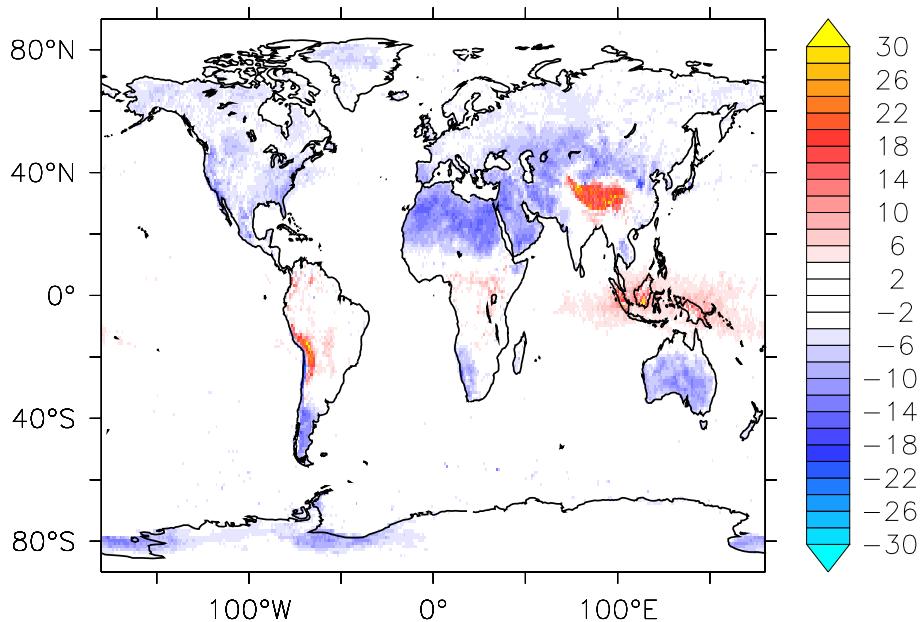


Fig. 3. Annual mean carbon monoxide relative difference (in %) between model results and MOPITT satellite observations near the surface. The model CO was calculated on the MOPITT kernel.

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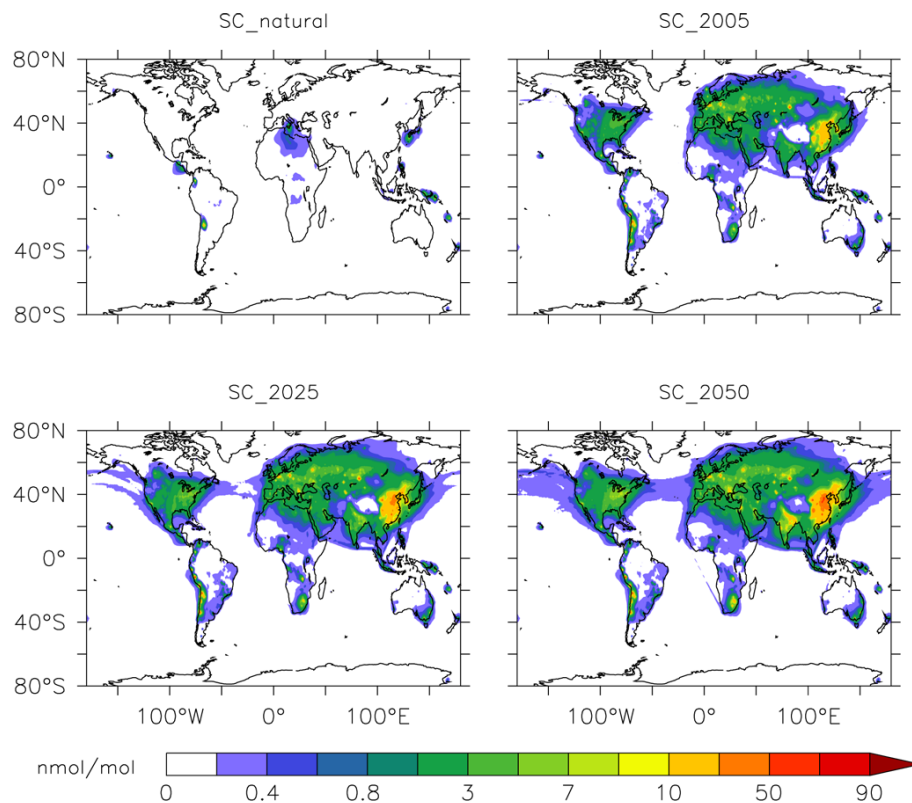


Fig. 4. Model calculated annual mean near-surface sulphur dioxide in nmol mol^{-1} from simulation *SC_natural*, *SC_2005*, *SC_2025* and *SC_2050*.

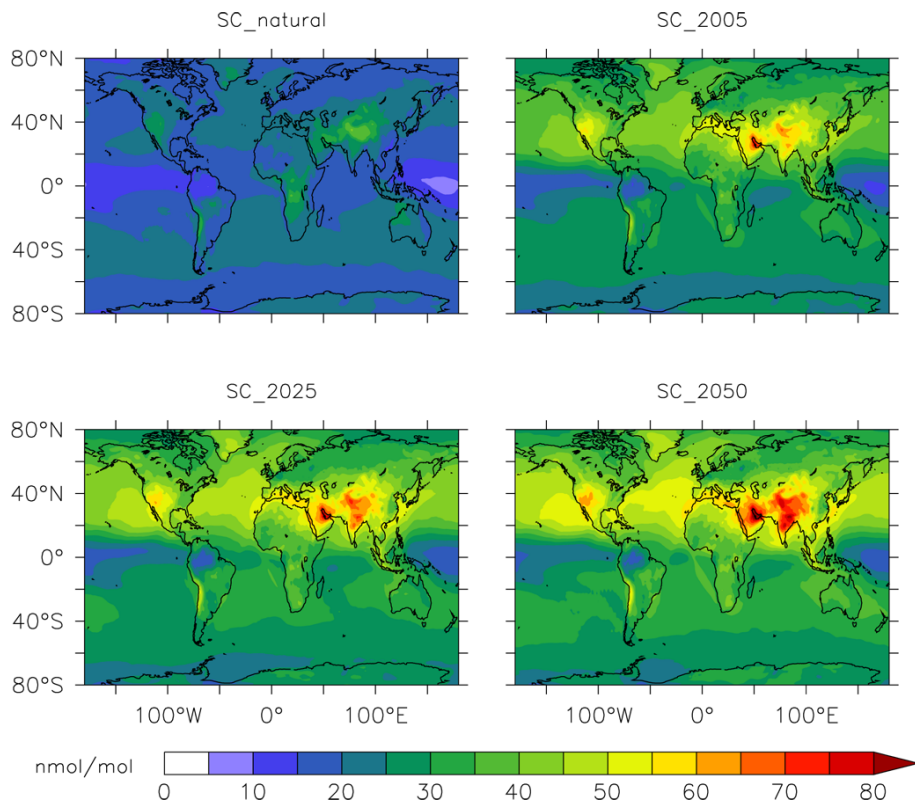


Fig. 5. Model calculated annual mean near-surface ozone in nmol mol^{-1} from simulation *SC_natural*, *SC_2005*, *SC_2025* and *SC_2050*.

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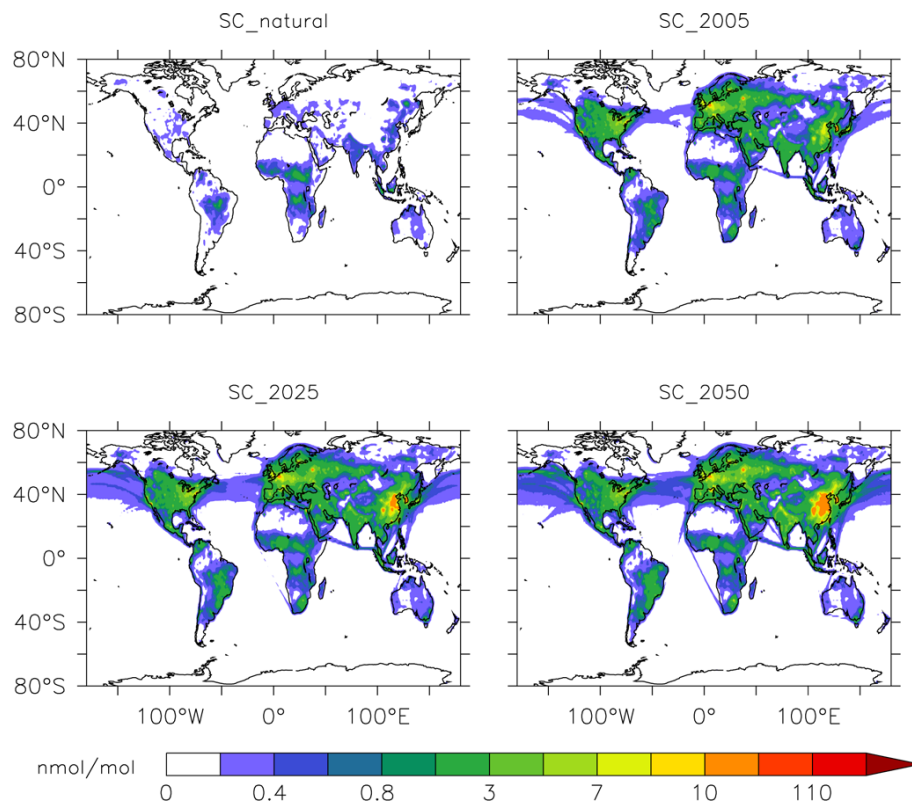


Fig. 6. Model calculated annual mean near-surface nitrogen dioxide in nmol mol^{-1} from simulation *SC_natural*, *SC_2005*, *SC_2025* and *SC_2050*.

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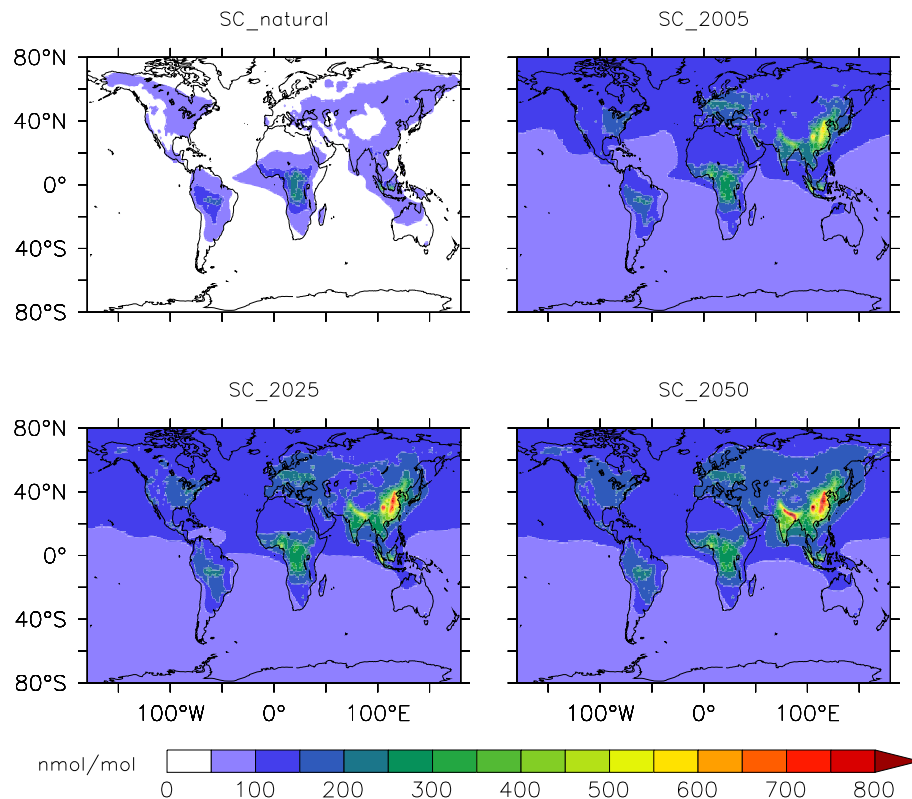


Fig. 7. Model calculated annual mean near-surface carbon monoxide in nmol mol^{-1} from simulation *SC_natural*, *SC_2005*, *SC_2025* and *SC_2050*.

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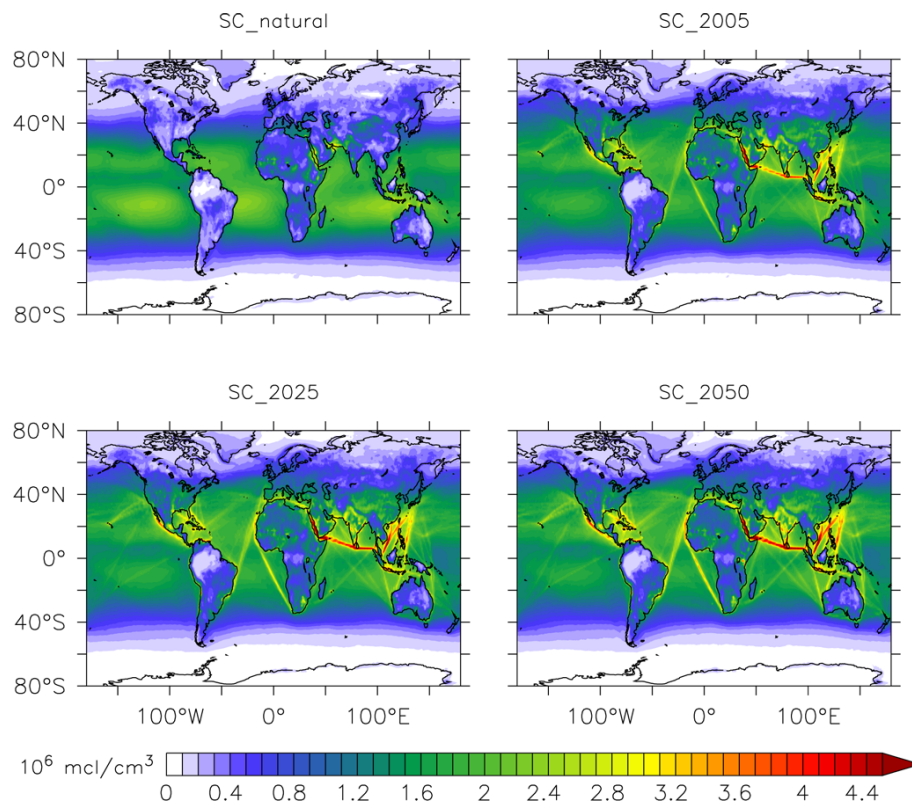


Fig. 8. Model calculated annual mean surface hydroxyl radical concentration in 10^6 cm^{-3} from simulation *SC_natural*, *SC_2005*, *SC_2025* and *SC_2050*.

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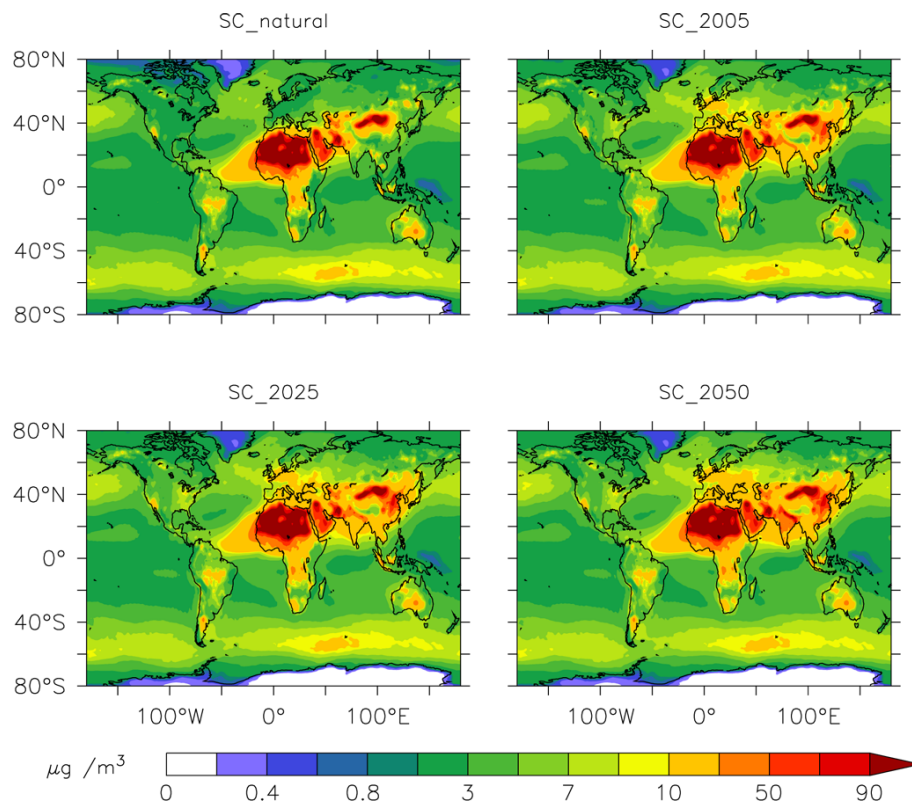


Fig. 9. Model calculated annual mean near-surface PM_{2.5} in $\mu\text{g m}^{-3}$ from simulation SC_{natural}, SC₂₀₀₅, SC₂₀₂₅ and SC₂₀₅₀.

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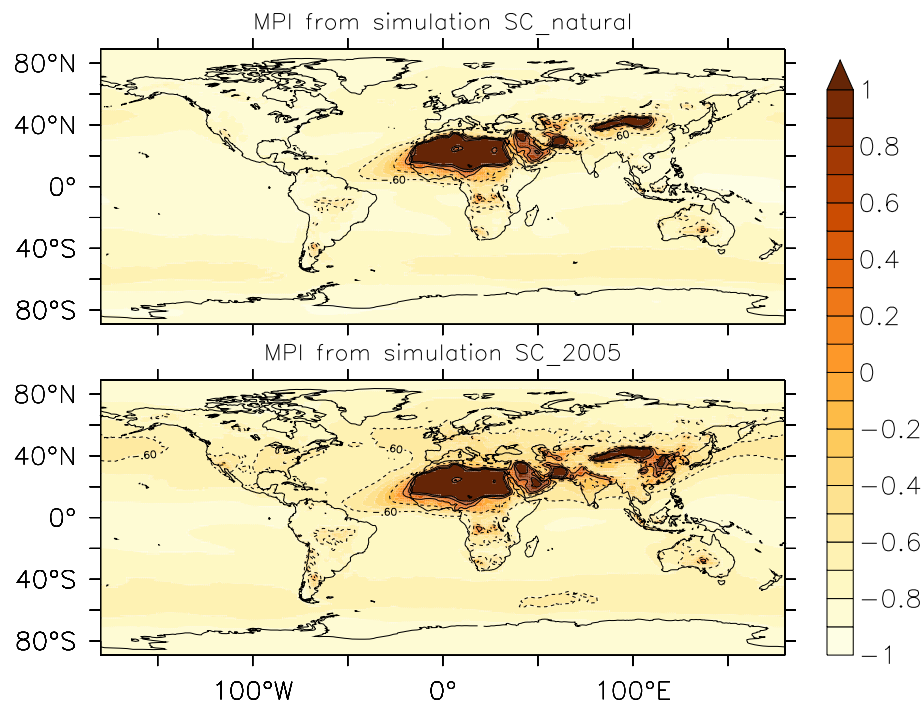


Fig. 10. Multi Pollutant Index (MPI) as calculated from simulation *SC_natural* and from simulation *SC_2005*. The contours are shown at multiples of 0.4.

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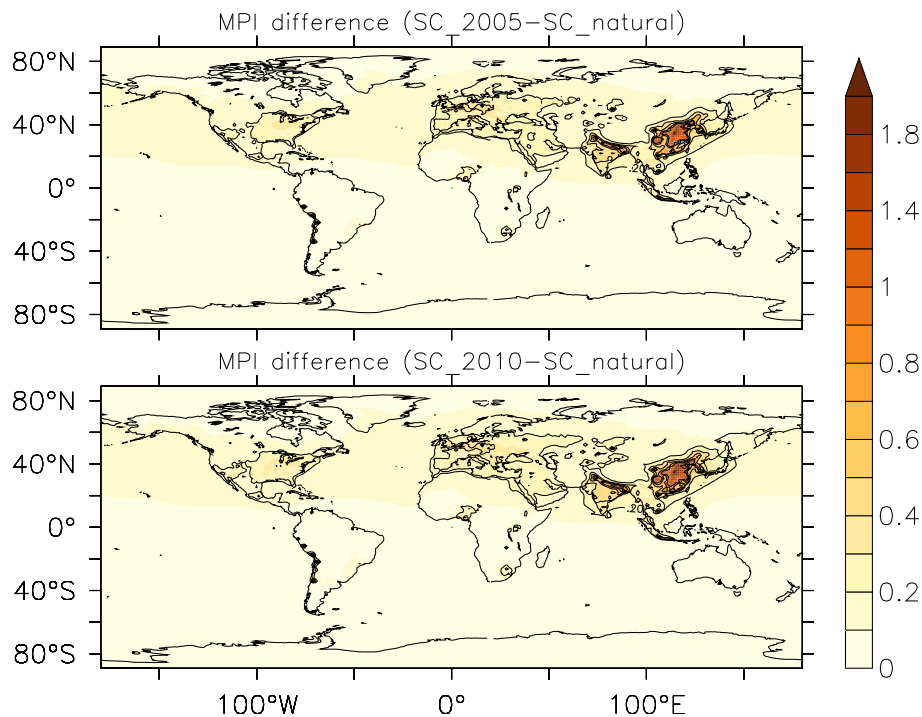


Fig. 11. Difference between the MPI estimated from simulations *SC_2005/SC_2010* and the MPI calculated from simulation *SC_natural*. The contours are shown at multiples of 0.2 up to the level of 1.

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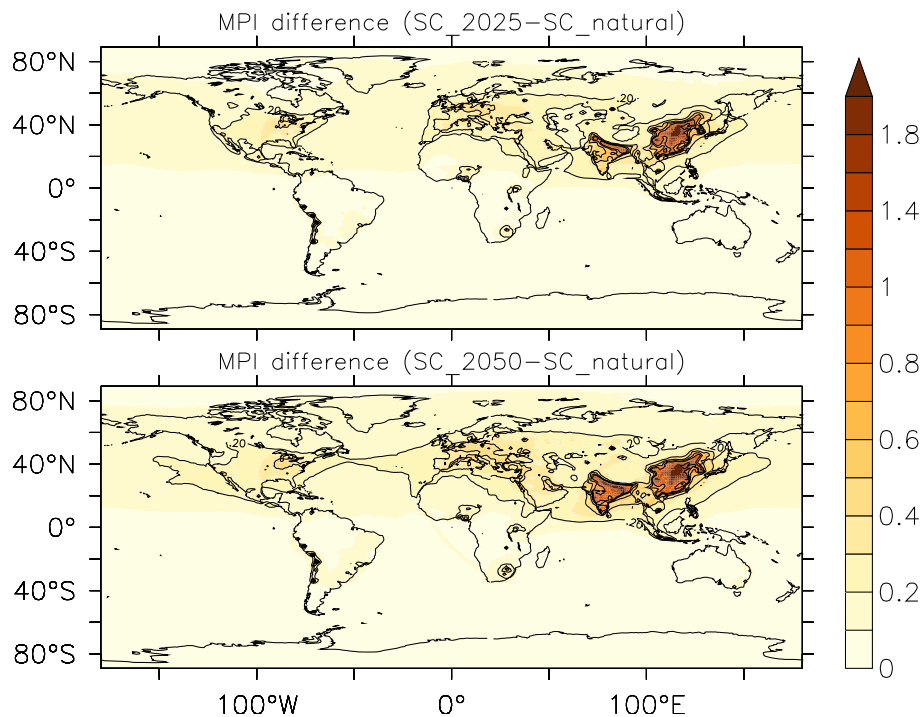


Fig. 12. Difference between the MPI estimated from simulations *SC_2025/SC_2050* and the MPI calculated from simulation *SC_natural*. The contours are shown at multiples of 0.2 up to the level of 1.

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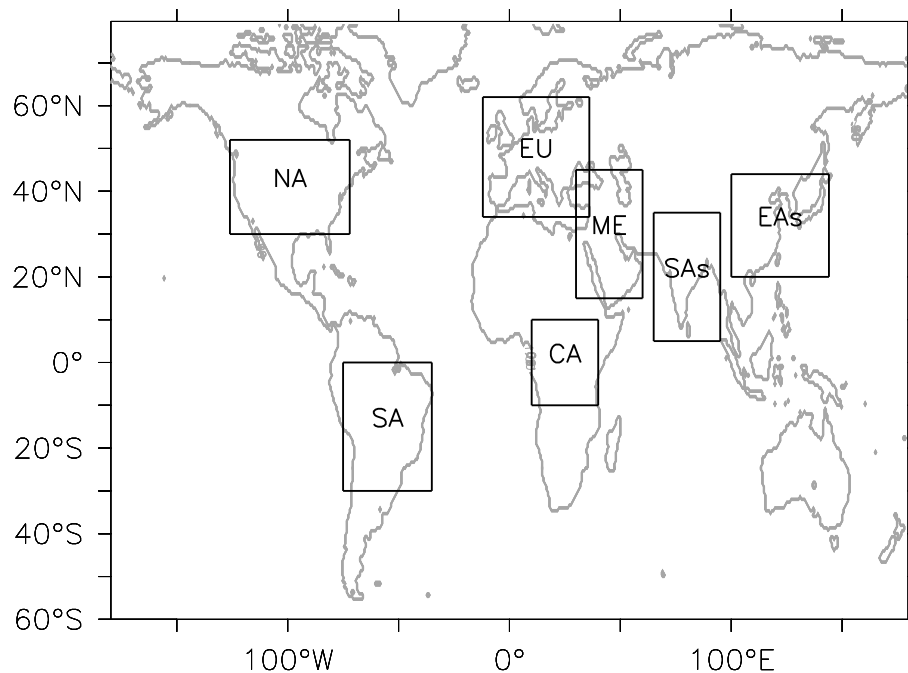


Fig. 13. Location and extension of the geographical regions used in the PCMPI calculation. The regions are: North America (NA), Europe (EU), East Asia (EAs), South Asia (SAs), Central Africa (CA) and South America (SA).

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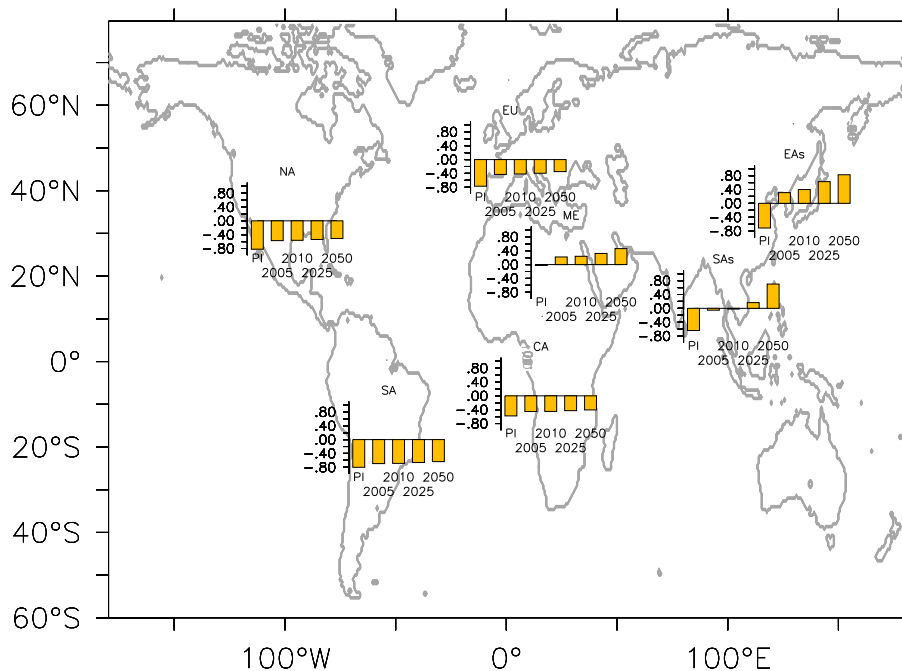


Fig. 14. Per Capita Multi Pollutant index values for different regions. This figure summarises Table 4. The regions are: North America (NA), South America (SA), Europe (EU), Middle East (ME), Central Africa (CE), South Asia (SAs) and East Asia (EAs).

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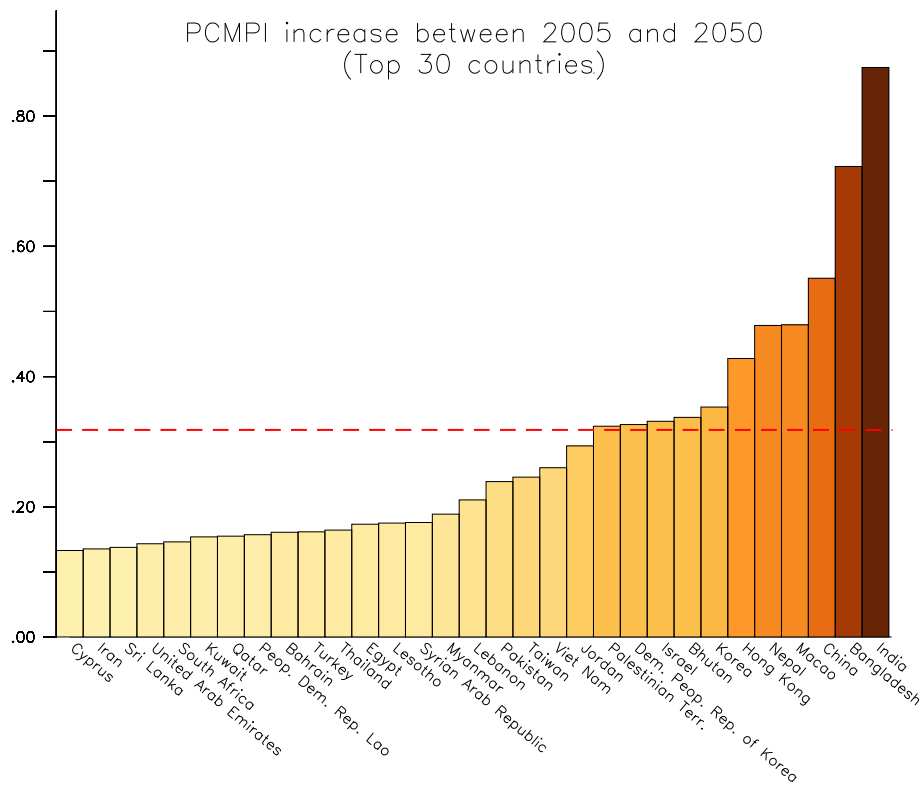


Fig. 15. PCMPI changes between 2005 and 2050 for different countries (30 countries with the strongest changes). The red dashed line depicts the global PCMPI increase between 2005 and 2050.

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