

# Non-linear response of PM<sub>2.5</sub> to changes in NO<sub>x</sub> and NH<sub>3</sub> emissions in the Po basin (Italy): consequences for air quality plans

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**Abstract:** Air pollution is one of the main causes of damages to human health in Europe with an estimate of about 380 000 premature deaths per year in the EU28, as the result of exposure to fine particulate matter (PM<sub>2.5</sub>) only. In this work, we focus on one specific region in Europe, the Po basin, a region where chemical regimes are the most complex, showing important non-linear processes, especially those related to interactions between NO<sub>x</sub> and NH<sub>3</sub>. We analyse the ~~sensitivitiessensitivity~~ of PM<sub>2.5</sub> ~~concentration~~ to NO<sub>x</sub> and NH<sub>3</sub> emissions by means of a set of EMEP ~~model~~ simulations performed with different levels of emission reductions, from 25% up to a total switch-off of those emissions. Both single and combined precursor reduction scenarios are applied to determine the most efficient emission reduction strategies and quantify the interactions between NO<sub>x</sub> and NH<sub>3</sub> emission reductions. The results confirmed the peculiarity of secondary PM<sub>2.5</sub> formation in the Po basin, characterised by contrasting chemical regimes within distances of few (hundreds of) kilometres, as well as ~~strong~~ non-linear responses to emission reductions during wintertime. One of the striking results is the ~~slight~~ increase of the PM<sub>2.5</sub> concentration levels when NO<sub>x</sub> emission reductions are applied in NO<sub>x</sub>-rich areas, such as the surroundings of Bergamo. The increased oxidative capacity of the atmosphere is the cause of the increase of PM<sub>2.5</sub> induced by a reduction in NO<sub>x</sub> emission. This process can have contributed to the absence of significant PM<sub>2.5</sub> concentration decrease during the COVID-19 lockdowns in many European cities. It is important to account for this process when designing air quality plans, since it could well lead to transitional increases in PM<sub>2.5</sub> at some locations in winter as NO<sub>x</sub> emission reduction measures are gradually implemented. While PM<sub>2.5</sub> chemical regimes, determined by the relative importance of the NO<sub>x</sub> vs. NH<sub>3</sub> responses to emission reductions, show large variations seasonally and spatially, they ~~are not very sensitive to se variations are less important in terms of~~ moderate (up to 50-60%) emission reductions ~~level~~. ~~Beyond 25% emission reduction strength, responses of PM2.5 concentrations to NOx emission reductions become non-linear in certain areas of the Po basin mainly during wintertime.~~

**Keywords:** urban air pollution, air quality planning, non-linearity, chemical regimes

## 1. Introduction

Air pollution is one of the main causes of damages to human health in Europe with an estimate of about 380 000 premature deaths per year in the EU28, as the result of exposure to fine particulate matter (PM<sub>2.5</sub>) only (EEA, 2020). Many of the exceedances to the EU limit values occur in urban areas where most of the population is exposed.

45 PM<sub>2.5</sub> is partly emitted directly (primary particles) and partly formed through photo-chemical reactions that involve  
gaseous precursors like SO<sub>x</sub>, NO<sub>x</sub>, NH<sub>3</sub> and non-methane volatile organic compounds (NMVOC) to form secondary  
inorganic and organic aerosol (SIA and SOA). The secondary fraction is often dominating the total concentration of  
particulate matter in urban areas as shown by e.g., Beekmann et al. (2015) for the Greater Paris region, De Meij et  
50 al. (2006, 2009) or Larsen et al. (2012) in northern Italy, hence the importance to understand the complex chemical  
processes that lead to its formation. In particular, it is key to identify the precursors involved in these reactions in  
order to target the right sectors of activity in air quality plans to effectively reduce pollution levels. According to the  
EDGAR estimates for 2015-~~(EDGAR, 2020)~~ for Italy, about 90% of the NH<sub>3</sub> is directly emitted in the atmosphere  
by the agriculture sector while SO<sub>x</sub> precursors are predominantly released by the energy production and use  
(industrial) sectors, ~~about 90%~~ (EDGAR ~~estimates for 2015, 2020~~). For NO<sub>x</sub>, emissions are spread among various  
55 sectors, with transport (50%), industry (40%) and agriculture (4%) being the main ones. The gaseous precursors of  
secondary organic aerosols (SOA) include a vast range of ~~high and low volatility~~ NMVOCs among which biogenic  
terpenes and anthropogenic aromatics. The main sources of aromatics in Italy were in 2012 transport (58%), use of  
fuels and solvents (32%), and domestic heating (15 %) ~~(EDGAR, 2020)~~.

Regarding SIA, early works used box models with thermodynamic schemes to address the sensitivity of ammonium  
60 nitrate and sulfate concentrations to gaseous NH<sub>3</sub>, NO<sub>x</sub>, and SO<sub>2</sub> emissions (Watson et al., 1994; Blanchard and  
Hidy, 2003); ~~;~~ Pozzer et al. 2017; Guo et al. 2018; Nenes et al. 2020. These models were later on integrated into  
chemical transport models (CTM), in particular to address the benefit of additional NH<sub>3</sub> emissions reductions in  
addition to already ongoing SO<sub>2</sub> and NO<sub>x</sub> emission reductions. For North America, Makar et al. (2009) simulated  
with a regional CTM that a 30% reduction of ammonia emissions would lead to about 1 µg/m<sup>3</sup> m<sup>-3</sup> reduction in  
65 PM<sub>2.5</sub>. For Europe, Bessagnet et al. (2014) simulated the effects of a 30% NH<sub>3</sub> emission reduction in addition to  
those foreseen by the Gothenburg Protocol for 2030, and found that the G-ratio defined as the ratio between free  
ammonia and total nitrate (Ansari and Pandis, 1998) was a good predictor for the efficiency of NH<sub>3</sub> reductions on  
SIA concentrations. These sensitivities to emission reductions are often governed by complex chemical mechanisms.  
A well-~~described-known~~ phenomenon is the release of free ammonia as a result of decreased SO<sub>2</sub> emissions and  
70 sulfate formation, which allows for the formation of additional particulate nitrate, as described for example by  
Blanchard and Hidy (2003) and Shah et al. (2018) for wintertime PM<sub>2.5</sub> over the Eastern US. For eastern China, Fu  
et al. (2017) and Lachatre et al. (2019) showed both from modelling and satellite observations that ~~this such~~  
processes leads lead to strongly increased ammonia tropospheric columns. Finally, several works compared CTM  
simulations to specific observations. For instance, Pay et al. (2012) showed that the G-ratio was generally  
75 underestimated over Europe, inducing that the Caliope model they used could probably overestimate the efficiency  
of NH<sub>3</sub> emission reductions. Petetin et al. (2016) came to a similar conclusion comparing CHIMERE CTM  
simulations to observations in the Paris region. They ascribed this underestimation to missing NH<sub>3</sub> emissions  
especially during warmer periods.

The formation of SOA results from even more complex reactions involving photo-chemical oxidation (as for SIA),  
80 nitration, fragmentation, and oligomerisation of gaseous precursors or secondary products (Kroll and Seinfeld, 2008;  
Shrivastava et al., 2017). Models generally use simplified parameterizations to calculate the SOA formation yield  
from various classes of parent VOCs (Tsigaridis et al., 2014), and comparison with measurements often show that  
SOA sources are still missing in models (Huang et al., 2020); ~~;~~ Tsimpidi, 2016.

In this work, we focus on one specific region in Europe, the Po basin. In a companion paper (Clappier et al.,  
85 ~~2020~~2021) that analyses PM secondary formation chemical regimes across Europe, the Po basin is clearly identified  
as a peculiar area where the chemical regime distributions are the most complex, showing ~~important~~ non-linear  
processes (Thunis et al., 2013 and 2015; Carnevale, 2020; Bessagnet, 2014), especially those related to interactions  
between NO<sub>x</sub> and NH<sub>3</sub>. The Po basin is also one of the pollution hot spots in Europe where the number of days  
above the limit values prescribed by the European Ambient Air Quality Directives (AAQD) for PM<sub>10</sub> is yet largely  
90 exceeded (EEA, 2020). This situation results from the high emission density in this region and also from the  
geographical setting of the area, in border of the Alps and Apennines mountain ranges that lead to very weak winds  
in the area, favouring the accumulation of atmospheric pollutants.

We focus the present analysis on the NH<sub>3</sub>-NO<sub>x</sub> chemical processes and describe their spatial and seasonal  
variability, which could help to design more effective mitigation strategies. We start by describing the modelling  
95 set-up and detail the series of simulations required to perform our analysis. Section 3 provides a brief overview of

the modelled base-case concentrations. In Section 4, we analyse the sensitivity of  $PM_{2.5}$  concentrations to  $NH_3$  and  $NO_x$  emissions. Section 5 provides an analysis of the non-linearity in  $PM_{2.5}$  response to these emissions while in Section 6, we discuss the implications of these results in terms of mitigation measures and design of air quality plans. Conclusions are finally proposed.

## 100 2. Methodology

### 2.1 Modelling set-up

The modelling study is performed with the EMEP/MSC-W air quality model, version rv4\_17 (Simpson et al., 2012). The emission input consists of gridded annual national emissions ( $SO_2$ ,  $NO$ ,  $NO_2$ ,  $NH_3$ , NMVOC, CO and primary  $PM_{2.5}$ ) at  $0.1 \times 0.1$  degrees resolution, based on data reported every year by parties to the Convention on Long Range Transboundary Air Pollution (CLRTAP). These emissions are provided for 10 anthropogenic source-sectors classified by SNAP (Selected Nomenclature for Air Pollution) codes (EMEP, 2003). Meteorological input data are based on forecasts from the Integrated Forecast System (IFS), a global operational forecasting model from the European Centre for Medium-Range Weather Forecasts (ECMWF). Meteorological fields are retrieved at a  $0.1 \times 0.1$  degree longitude latitude resolution and are interpolated to the  $50 \times 50 km^2$  polar-stereographic grid projection (EMEP, 2011).

The gas-phase chemistry is based on the evolution of the so-called “EMEP scheme” as described in Simpson et al. (2012) and references therein. The chemical scheme couples the sulfur and nitrogen chemistry to the photochemistry using about 140 reactions between 70 species (Andersson-Sköld and Simpson, 1999; Simpson et al. 2012). In the EMEP Status Report 1/2004 (Fagerli et al., 2004) the reactions are described that cover acidification, eutrophication and ammonium chemistry. The aqueous phase chemistry describes the formation of sulfate in clouds via  $SO_2$  oxidation by ozone and  $H_2O_2$  and catalysed by metal ions. An important pathway of particulate nitrate formation is through the hydrolysis of  $N_2O_5$  on wet aerosol surfaces that converts  $NO_x$  into  $HNO_3$ . More information on the chemical equations is given in Simpson et al. (2012), section 7.

The EMEP model has two size fractions for aerosols, fine aerosol ( $PM_{2.5}$ ) and coarse aerosol ( $PM_{10-2.5}$ ). The aerosol components presently accounted for are sulfate ( $SO_4^{2-}$ ), nitrate ( $NO_3^-$ ), ammonium ( $NH_4^+$ ), anthropogenic primary PM and sea salt.

For inorganic aerosols, EMEP uses the MARS equilibrium module to calculate the partitioning between gas and fine-mode aerosol phase in the system of  $SO_4^{2-}$ ,  $HNO_3$ ,  $NO_3^-$ ,  $NH_3$  and  $NH_4^+$  (Binkowski and Shankar, 1995). Aerosol water is calculated to account for particle water within the  $PM_{2.5}$  mass, which depends on the mass of soluble PM fraction and on the type of salt mixture in particles. Sea salt (sodium chloride) and dust components are not accounted for by MARS, which might lead to PM underestimations close to coastal sites and where the dust contribution is important. More information on the gas and aerosol partitioning is given in Simpson et al. (2012), section 7.6.

Regarding secondary organic aerosols (SOA), the EmChem09soa scheme is used, which is a simplified version of the so-called volatility basis set (VBS) approach (Robinson et al., 2007; Donahue et al., 2009). The VBS mechanism is discussed in detail in Bergström et al. (2012). The main differences between the VBS schemes and EmChem09soa is that all primary organic aerosol (POA) emissions are treated as non-volatile in EmChem09soa. This is done to keep the emission totals of both PM and VOC components the same as in the official emission inventories. The semi-volatile biogenic and anthropogenic SOA species are assumed to further oxidise (also known as ageing process) in the atmosphere by OH-reactions. This will lead to a reduction in volatility for the SOA, and thereby increased partitioning to the particle phase. More information on SOA is given in Simpson et al. (2012), section 7.7.

The modelling domain covers the entire Po basin (Figure 1) with a resolution of  $0.1$  by  $0.1$  degree resolution (polar stereographic projection centred at  $60^\circ N$ ) and includes 20 vertical levels. The initial and background concentrations for ozone are based on Logan (1998) climatology, as described in Simpson et al. (2003). For the other species, background/initial conditions are set within the model using functions based on observations (Simpson et al., 2003; and Fagerli et al., 2004). The simulations cover the entire meteorological year 2015. We will not discuss the

validation of the base-case simulation, as this is available in other publications (e.g. Simpson et al., 2012) and regular status reports by EMEP ([https://emep.int/mscw/mscw\\_publications.html](https://emep.int/mscw/mscw_publications.html)).

145 In this work, we simulated a series of 24 scenarios in which NO<sub>x</sub> and NH<sub>3</sub> emissions were reduced independently or simultaneously by 25, 50, 75 and 100% from the base case reference levels. Emission reductions were applied over the entire Po-basin domain for a complete meteorological year (2015).

## 2.2 Spatial and temporal focus

150 Results are generally presented in terms of maps but three locations within the domain were selected for a more detailed analysis. The locations are Bergamo (Be) in the northern part of the domain, Mantova (Ma) in the central eastern part of the Po basin and Bologna (Bo) in its southern part. As described in the following sections, we will see that these locations show very different behaviours in terms of response to emission changes.

155 We also aggregate results into 2 seasons: winter and summer that cover the period from November to February and from April to September, respectively. These two seasons are characteristic of different chemical regimes as illustrated in the following sections. The process to define the temporal bounds of these two seasons is discussed in Annex 1. The two remaining months (March and October) represent transition periods and are not considered in our analysis.

As we only analyse ~~the~~ processes involving inorganic gas-phase precursors, our focus is on secondary inorganic PM<sub>2.5</sub> although most of the results are expressed in terms of total PM<sub>2.5</sub> concentrations. The impact of NO<sub>x</sub> emission reductions on SOA concentration is only briefly discussed in section 5.

## 160 2.3 Indicators

To describe the interactions between NH<sub>3</sub> and NO<sub>x</sub> emissions, we use the relationship proposed by Stein and Alpert (1993) and Thunis and Clappier (2014). This relation expresses the change of concentration resulting from a reduction of both precursors NO<sub>x</sub> and NH<sub>3</sub> simultaneously, as the sum of two single concentration changes and an interaction term, as follows:

$$165 \Delta C_{NO_xNH_3}^\alpha = \Delta C_{NO_x}^\alpha + \Delta C_{NH_3}^\alpha + \hat{C}_{NO_xNH_3}^\alpha \quad (1)$$

Where  $\Delta C$  stands for the PM<sub>2.5</sub> concentration change (reference minus scenario) for a percentage  $\alpha$  emission reduction (thus the term  $\Delta C/\alpha$  is defined positive for a concentration reduction, consecutive to an emission reduction) and  $\hat{C}$  for the interaction term. We then scale each of these terms by the emission reduction ( $\alpha$ ) to generate ~~potentials~~potential impacts (P) (Thunis and Clappier, 2014).

$$170 \frac{\Delta C_{NO_xNH_3}^\alpha}{\alpha} = P_{NO_xNH_3}^\alpha = P_{NO_x}^\alpha + P_{NH_3}^\alpha + \hat{P}_{NO_xNH_3}^\alpha = \frac{\Delta C_{NO_x}^\alpha}{\alpha} + \frac{\Delta C_{NH_3}^\alpha}{\alpha} + \frac{\hat{C}_{NO_xNH_3}^\alpha}{\alpha} \quad (2)$$

175 This division by the factor  $\alpha$  is a ~~mean~~mean to ~~virtually~~-extrapolate virtually the impact resulting from any percentage emission reduction ~~from 1~~ to 100%. ~~Potentials~~Potential impacts facilitate the comparison of concentration changes obtained for different emission reduction levels. Indeed, equal ~~potentials~~potential impacts imply a linear relationship between emission reductions and concentration changes. For example,  $P^\alpha = P^\beta \Rightarrow \Delta C^\alpha = \frac{\alpha}{\beta} \Delta C^\beta$ , for  $\alpha$  and  $\beta$ , two emission reduction levels.

The overall potential impact is therefore the sum of two single ~~potentials~~potential impacts and one interaction term.

A relation between the ~~potentials~~potential impacts of combined emission reductions at two levels of intensity is obtained by writing (2) for two reduction levels  $\alpha$  and  $\beta$  and subtract the two equations. This leads to the following relation:

$$P_{NO_xNH_3}^\beta = P_{NO_xNH_3}^\alpha + \underbrace{P_{NO_x}^\beta - P_{NO_x}^\alpha}_{\hat{P}_{NO_x}^{\beta-\alpha}} + \underbrace{P_{NH_3}^\beta - P_{NH_3}^\alpha}_{\hat{P}_{NH_3}^{\beta-\alpha}} + \underbrace{\hat{P}_{NO_xNH_3}^\beta - \hat{P}_{NO_xNH_3}^\alpha}_{\hat{P}_{NO_xNH_3}^{\beta-\alpha}} \quad (3)$$

180

or equivalently via equation 2:

$$P_{NO_xNH_3}^\beta = P_{NO_x}^\alpha + P_{NH_3}^\alpha + \underbrace{\hat{P}_{NO_xNH_3}^\alpha + \hat{P}_{NO_x}^{\beta-\alpha} + \hat{P}_{NH_3}^{\beta-\alpha} + \hat{P}_{NO_xNH_3}^{\beta-\alpha}}_{\text{non-linear terms}} \quad (4)$$

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While the two first terms on the right hand side of Eq. 4 represent the single ~~potentials~~ potential impacts, the remaining right hand side terms quantify the magnitude of non-linearities.  $\hat{P}_{NO_xNH_3}^\alpha$  quantifies the NO<sub>x</sub>-NH<sub>3</sub> interaction at level  $\alpha$ ,  $\hat{P}_{NO_x}^{\beta-\alpha}$  and  $\hat{P}_{NH_3}^{\beta-\alpha}$  are the single non-linearities associated to NO<sub>x</sub> and NH<sub>3</sub> emissions, respectively, between levels  $\alpha$  and  $\beta$ , whereas  $\hat{P}_{NO_xNH_3}^{\beta-\alpha}$  represents the incremental change of the NO<sub>x</sub>-NH<sub>3</sub> interaction between levels  $\alpha$  and  $\beta$ .

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Information about non-linearity is important to design air quality plans as it informs on the robustness of a given response, i.e. whether or not this response remains valid over a certain range and type of emission reductions. Because air quality models often provide responses for a limited set of scenarios that are then used as a basis to interpolate/extrapolate the responses to other emission reduction levels, robustness shall always be carefully assessed.

In the next section, we present the baseline results in terms of spatial and temporal variations.

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### 3. Baseline concentrations of PM<sub>2.5</sub> and gaseous inorganic precursors

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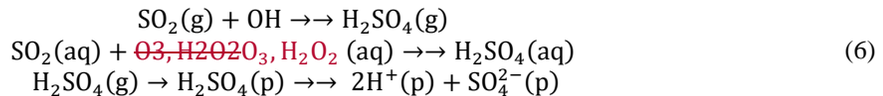
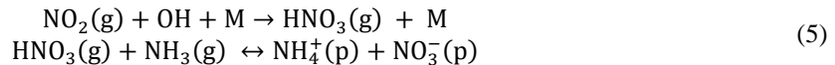
Before analysing the impact of emission changes on concentrations, it is worth having a look at the baseline concentration fields. In Figure 1, the yearly averaged PM<sub>2.5</sub> concentration fields show a widespread pollution plume covering most of the area, with peak values extending in its central part. The maximum modelled yearly values reach 29  $\mu\text{g}/\text{m}^3 \cdot \text{m}^{-3}$ , that represent an average between maximum winter values (maximum of 59  $\mu\text{g}/\text{m}^3 \cdot \text{m}^{-3}$ ) and minimum summer values (17  $\mu\text{g}/\text{m}^3 \cdot \text{m}^{-3}$ ).

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The seasonal fields of PM<sub>2.5</sub> clearly show that high yearly average values mostly result from the winter season contributions when more stable atmospheric conditions lead to stagnant conditions, favouring the accumulation of particulate matter in the area (Pernigotti et al., 2014; Raffaelli et al., 2020). The increased emissions from the residential sector (heating, especially wood burning) ~~also~~ foster this process (Ricciardelli et al., 2017; Hakimzadeh et al., 2020). The Wintertime low temperatures also favour the partitioning of semi-volatile components (e.g. ammonium nitrate) towards the particulate phase. Overall, the relative contribution of secondary inorganic particles (SIA) ranges between 40 and 50%, regardless of the season and is quite homogeneously distributed spatially over the entire area. Strategies targeting SIA have therefore the potential to abate about half of the total PM<sub>2.5</sub> concentration.

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As mentioned earlier, the secondary inorganic fraction of PM<sub>2.5</sub> results from complex atmospheric processes that involve gaseous precursors (mainly SO<sub>2</sub>, NO<sub>x</sub> and NH<sub>3</sub>), that can be summarized by the two following chemical pathways:





215 where (g) means “gas phase”, (aq) aqueous phase, while (p) means “particulate matter”, and the character  $\rightarrow$  symbolizes a chemical pathway that summarizes a set of underlying reactions.

The second of these pathways is generally slower than the first one, the NO<sub>2</sub> oxidation specific time constant being typically some hours to a day, and that of SO<sub>2</sub> one to several days (Seinfeld and Pandis, 2006).

220 The spatial fields for the seasonal average concentrations of these precursors (Figure 2) reflects their emissions spatial patterns resulting in a NO<sub>2</sub>-rich area that comprises Milan plus its northern districts (and to a lesser extent, Turin), while NH<sub>3</sub> is more abundant in the central part of the Po basin, east of Milan, where intensive agriculture practices take place. Finally, high SO<sub>2</sub> concentrations are collocated with the NO<sub>2</sub>-rich areas nearby Milan but with an additional zone around the harbour city of Genova (along the south coast), reflecting the more important SO<sub>2</sub> emissions from the shipping sector there. However, SO<sub>2</sub> concentrations are about one order of magnitude below those of NO<sub>2</sub>. Seasonal variations are well marked for NO<sub>2</sub> and SO<sub>2</sub> concentrations, not so much in terms of minimum and maximum values but rather in terms of spread with an extended high concentration zone covered during wintertime. In contrast, NH<sub>3</sub> concentrations remain very similar in summer and winter, both in terms of values and spatial distribution.

230 In next sections, we simulate a series of emission reduction scenarios to analyse the response of PM<sub>2.5</sub> concentration to single and combined reductions of NH<sub>3</sub> and NO<sub>x</sub>.

#### 4. Analysis of the SIA formation chemical regimes

##### 4.1 Seasonal trends

235 From a strategic point of view, it is important to know whether NO<sub>x</sub> (mostly emitted by the transport, industry and residential sectors) or NH<sub>3</sub> (mostly emitted by agriculture) need to be reduced in priority in order to reach effective results on particulate pollution mitigation. In Clappier et al. (2020, 2021), we have observed a great heterogeneity in SIA formation chemical regimes across the Po-basin, different regimes being present in limited geographical areas. Here we intend to look at these regimes in more detail.

240 To analyse in details the chemical regimes in the Po basin, we compare the two single potential impacts  $P_{NO_x}^{25\%}$ ,  $P_{NH_3}^{25\%}$  obtained for moderate emission reductions of 25% (-). Figure 3 provides a spatial overview of the difference between these two potential impacts ( $P_{NO_x}^{25\%} - P_{NH_3}^{25\%}$ ). This indicator tells whether reductions of NO<sub>x</sub> or NH<sub>3</sub> will lead to the greatest PM<sub>2.5</sub> concentration abatement, i.e. if the regime is rather NO<sub>x</sub>- or NH<sub>3</sub>-sensitive, with positive and negative values, respectively.

- 245 • During summer time (Figure 3 – left), the entire area is under weak NO<sub>x</sub>-sensitive conditions with a maximum intensity in its central part, between Bergamo and Mantova.
- During wintertime (Figure 3 – right), the situation is contrasted with a wide and intense NH<sub>3</sub>-sensitive area that appears around and south-eastwards of Bergamo. This area includes big cities like Milan. Other (not as marked) NH<sub>3</sub> sensitive regime zones appear nearby coastal areas. Most strongly NO<sub>x</sub>-sensitive areas are located in the eastern parts of the domain, north of Bologna and Venice. As expected, NH<sub>3</sub>-sensitive regimes are generally collocated with the NO<sub>2</sub>- and SO<sub>2</sub>- rich areas (Figure 2), whereas NO<sub>x</sub>-sensitive regimes coincide with NH<sub>3</sub>-rich areas. The cases of the three selected cities (Bergamo, Mantova and Bologna) representative of the NH<sub>3</sub>-sensitive, NO<sub>x</sub>-sensitive and neutral regimes, respectively, are further analysed below.

250 The chemical regimes deduced from the results of emission reduction scenarios can be compared with the maps of the G-ratio (Figure 4), defined by Ansari and Pandis (1998) as the ratio between free ammonia (NH<sub>3</sub> and NH<sub>4</sub><sup>+</sup>) and total nitrate (HNO<sub>3</sub> + NO<sub>3</sub><sup>-</sup>) after neutralization of H<sub>2</sub>SO<sub>4</sub>. Values of the G-ratio below 1 indicate a NH<sub>3</sub>-limited chemical regime, while values above 1 characterize a HNO<sub>3</sub>-limited chemical regime.

$$G = \frac{NH_3(g) + NH_4^+(p) - 2SO_4^{2-}(p)}{HNO_3(g) + NO_3^-(p)} \quad (7)$$

We first note that that formulation of the G-ratio, in terms of abundance of free total ammonia and total nitrate, differs from that of the NH<sub>3</sub> and NO<sub>x</sub> emission sensitive chemical regime. The following discussion tends to show how both diagnostics can be used in a complementary way. During summer, the G-ratio values well above unity indicate a HNO<sub>3</sub> limited chemical regime across the Po basin. This corresponds to a NO<sub>x</sub> sensitive chemical regime in this region (Figure 3). Moreover, the location of the G-ratio maximum between Bergamo and Modena/Mantova spatially coincides with the most pronounced NO<sub>x</sub> sensitive regime, and to a maximum of NH<sub>3</sub> concentrations of about 20 µg/m<sup>3</sup> (Figure 2). Indeed, NO<sub>x</sub> emission reductions lead to HNO<sub>3</sub> concentration reductions that is the limiting factor in NH<sub>4</sub>NO<sub>3</sub> formation according to the G-ratio. During winter, the G-ratio still shows large values in the region south-east of Bergamo, but contrary to winter/summer, the chemical regime is clearly NH<sub>3</sub> sensitive (Figure 3). More generally, G-ratio values remain above unity over the whole Po basin while both NO<sub>x</sub> and NH<sub>3</sub> sensitive chemical regimes prevail in different areas. Thus, even if free total ammonia is more abundant, PM abatement is more sensitive to NH<sub>3</sub> than to NO<sub>x</sub> emissions. Therefore, the G-ratio, related to the abundance of total free ammonia and total nitrate, provides information which differs from that obtained by determining the distribution of the NH<sub>3</sub> and NO<sub>x</sub> emission sensitive chemical regimes. These differences illustrate the impossibility to directly use the G-ratio for air quality management, an interesting result in itself. We will further discuss this interesting behaviour later when addressing non-linearity in section 4.3.

#### 4.2 Impact of the emission reduction strength

In this section, we repeat the analysis of Section 4.1 for yearly average concentrations but looking at the step changes of regimes as we progressively remove/reduce emissions from the base case situation. (Figure 5). Chemical regimes are well in place for a 25% level reduction (top left) and are only slightly perturbed from 25 to 50% with a reinforcement of the NO<sub>x</sub> limited regime (top right). Despite this slight change in intensity, the regimes keep therefore the same spatial patterns. From 50% onward, chemical regimes tend to attenuate and reverse themselves from 75% to 100% (bottom right).

In other words, locations that are NH<sub>3</sub>-sensitive for the first steps moderate emission reductions will become NO<sub>x</sub>-sensitive for larger the last steps emission reductions, and vice versa.

#### 4.3 A summarized overview: the PM<sub>2.5</sub> isopleths

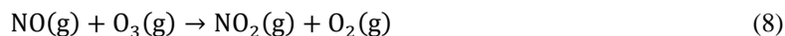
Like isopleth plots that show the variations in the O<sub>3</sub> concentrations as a function of NO<sub>x</sub> and VOC concentrations (Dodge, 1977), similar plots can be created for PM<sub>2.5</sub> concentrations as a function of NO<sub>x</sub> and NH<sub>3</sub> emissions. Simulation results have indeed often been presented as 2D isopleths of PM<sub>2.5</sub> or nitrate as a function of precursor emissions, which allows showing in a comprehensive manner their sensitivity, and also, in a qualitative manner, non-linear effects (for example Watson et al., 1994 over USA; Xing et al., 2018 over the Beijing–Tianjin–Hebei region in China). Figure 6 shows the PM<sub>2.5</sub> isopleths obtained through an interpolation among the 25 simulation concentration values (these 25 simulations correspond to the white square symbols in each isopleth diagram) at the three locations Be, Ma, and Bo previously defined. The X- and Y-axes represent the strengths of the NH<sub>3</sub> and NO<sub>x</sub> emission sources, respectively. With this type of graphical representation, it is possible to visualize the response of PM<sub>2.5</sub> to a NO<sub>x</sub> emission change by moving vertically, the reaction to a NH<sub>3</sub> emission change by moving horizontally or the reaction to a combined NO<sub>x</sub>-NH<sub>3</sub> emission change by moving diagonally. The larger the number of isopleths we cross on the path (high gradient), the larger the expected impact from an emission reduction will be. A simple theoretical model to generate and interpret these isopleths is proposed in Annex 2.

From the analysis of the isopleths, we note the following points:

- In general, the isopleths show a regular pattern with a progressive decrease of PM<sub>2.5</sub> concentration when either the NO<sub>x</sub> or NH<sub>3</sub> emissions are reduced, with the only exception of Bergamo during wintertime, where NO<sub>x</sub> reductions up to 70% lead to a small increase of PM<sub>2.5</sub> (Figure 6 - top left), whatever the reduction in NH<sub>3</sub> emission is. We discuss this particular feature later in this section.

- The diagram areas can be divided into two zones separated by a ridge (dashed line in Figure 6). Above the ridge line, PM<sub>2.5</sub> is more sensitive to NH<sub>3</sub> while below the ridge line, it is more sensitive to NO<sub>x</sub> emission reductions. The orientation of the ridge (tending to vertical or horizontal) informs on the type of chemical regime (NO<sub>x</sub> or NH<sub>3</sub> sensitive, respectively).
- The efficiency of a NO<sub>x</sub> vs NH<sub>3</sub> emission reduction varies across locations. We can compare the efficiency of a given reduction by looking at the horizontal (for NO<sub>x</sub>) and vertical (for NH<sub>3</sub>) gradients. To support this comparison, we included in each diagram dashed oblique lines that connect similar PM<sub>2.5</sub> concentration values for single NO<sub>x</sub> and NH<sub>3</sub> emission reductions. The more vertical are these lines, the larger is the NH<sub>3</sub> abatement impact compared to the NO<sub>x</sub> abatement impact. Conversely, the more horizontal they are, the larger is the NO<sub>x</sub> abatement impact compared to the NH<sub>3</sub> abatement impact. For moderate emission reductions (up to 50%, top right corner), different behaviours are observed: while in Bergamo PM<sub>2.5</sub> is more sensitive to NH<sub>3</sub> reductions, it is more sensitive to NO<sub>x</sub> reductions in Mantova, and equally sensitive to both precursors in Bologna. This corresponds to the spatial patterns of NO<sub>x</sub>- and NH<sub>3</sub>-sensitive regimes depicted in Fig. 3.
- Winter- and summertime isopleths show completely different patterns in Bergamo whereas at the two other locations, they remain similar.
- At Mantova where moderate NO<sub>x</sub> reductions (e.g. 50%) are the most efficient among the 3 sites, NH<sub>3</sub> emission reductions are more efficient than NO<sub>x</sub> emission reductions for larger additional reductions (going for example from 75% to 100%). At Bergamo, NH<sub>3</sub> reductions are the most effective for moderate reductions whereas NO<sub>x</sub> reductions become more effective for larger reductions, as seen by the isopleths spacing. This confirms the findings of reversed chemical regimes for larger additional emission reductions detailed in the previous section and illustrated in Figure 5.

The special pattern of Bergamo's PM<sub>2.5</sub> isopleths during wintertime needs some additional discussion. The increase of the inorganic fraction of PM<sub>2.5</sub> as a response to NO<sub>x</sub> reductions during wintertime has already been noted by several authors (e.g. Le et al., 2020; Sheng et al., 2018). It has been related to an increase in the oxidizing capacity of the atmosphere and in particular to increased ozone levels. This is due to the prevailing titration of O<sub>3</sub> by NO in wintertime high NO<sub>x</sub> conditions and in the absence of photochemical ozone production due to reduced solar radiation (Kleinman et al., 1991).



Accordingly, for a factor of two ~~The impact of NOx emission reductions on the concentration of various pollutants in Bergamo during wintertime is illustrated in Figure 7. As expected, a 50% reduction in NO<sub>x</sub> emissions leads to a decrease, Figure 8 shows in NO<sub>2</sub> concentration (from 47 to 28 μg m<sup>-3</sup>, i.e. a factor of 1.7). In contrast, O<sub>3</sub> concentration increases from 8 to 16 μg m<sup>-3</sup>, roughly a factor of two increase for ozone, while NO<sub>2</sub> decreases by a factor of less than two (1.7).~~ These compensating changes result in a small increase in NO<sub>3</sub> radical production, (Eq. 9), the initial step of the major pathway of wintertime HNO<sub>3</sub> and particulate nitrate formation (Kenagy et al., 2018).



In this pathway, the NO<sub>3</sub> radical formation is followed by combination with NO<sub>2</sub> to form N<sub>2</sub>O<sub>5</sub>, a reversible process, and heterogeneous HNO<sub>3</sub> formation on wet particle surfaces.



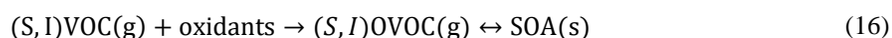
The NO<sub>3</sub> radical has three major rapid sinks, reaction with NO, photolysis, and reaction with ~~NMVOCS~~ NMVOCS especially terpenes.





Reactions 8 to 15 induce additional dependence of HNO<sub>3</sub> formation on NO<sub>x</sub> species on top of (5), but which partly cancel out, as they are both involved in formation and sink processes.

SOA is formed through a series of chemical reactions of gaseous precursors (mainly volatile-, intermediate volatile or semi-volatile organic compounds  $\{(\text{VOCs})\}$ ) with the oxidants O<sub>3</sub>, OH and nitrate radical (NO<sub>3</sub>) (Li et al., 2011).



Putting all the arguments together, it follows that wintertime ammonium nitrate formation over Bergamo is most probably controlled by NO<sub>3</sub> radical formation (9), and its strong non-linearity with respect to NO<sub>x</sub> emissions. The fact that this behaviour is observed in Bergamo and not in Mantova or Bologna is due to the much larger NO<sub>2</sub> levels in the Bergamo - Milano area (above 50 μg/m<sup>3</sup> m<sup>-3</sup> during winter, Fig.2). Such large NO<sub>2</sub> levels are also simulated locally over the Turin area, and also lead to a slightly NH<sub>3</sub> sensitive regime there despite a G-ratio well above unity. Beyond 50% NO<sub>x</sub> reduction, NH<sub>4</sub>NO<sub>3</sub> formation decreases because NO<sub>2</sub> decreases more rapidly than ozone increases up to its maximum (at 75% NO<sub>x</sub> emission reduction, see Figure 7).

~~This non-linearity between~~ The negative response of NH<sub>4</sub>NO<sub>3</sub> to NO<sub>x</sub> emission reductions and HNO<sub>3</sub> formation also in Be during wintertime explains the apparent discrepancies with the analysis of G-ratio indicating stronger sensitivity to HNO<sub>3</sub>, which indicates that NH<sub>4</sub>NO<sub>3</sub> is strongly HNO<sub>3</sub> limited. Simply, sensitivity to the HNO<sub>3</sub> concentration-limited chemical regime cannot be extrapolated to sensitivity to NO<sub>x</sub> emissions in case of the above shown non-linear behaviour-negative response. Total nitrate is less abundant, than free ammonia (defined as NH<sub>3</sub>(g) + NH<sub>4</sub><sup>+</sup>(p) - 2SO<sub>4</sub><sup>2-</sup>(p))<sub>2</sub> but NO<sub>x</sub> emission reductions below up to about 50% do not reduce its concentration, and NH<sub>3</sub> emission reductions are thus more efficient, even if NH<sub>3</sub> is more abundant than HNO<sub>3</sub>. In this respect, the complementary analysis of emission sensitivity and G-ratio gives interesting clues on possible non-linearity in the NO<sub>x</sub> - HNO<sub>3</sub> relationship cannot provide information about negative responses.

At Bergamo during winter, the increase in PM<sub>2.5</sub> (+ 1.8 μg/m<sup>3</sup> m<sup>-3</sup>) arising from a 50% reduction in NO<sub>x</sub> emission also results from an increase in sulfate (+ 0.3 μg/μg m<sup>3</sup> m<sup>-3</sup>) and in SOA (+ 0.6 μg/μg m<sup>3</sup> m<sup>-3</sup>) concentrations. Both sulfate and SOA concentrations are closely related to O<sub>3</sub> concentrations (Figure 7). The sulfate increase is comparable in magnitude to the nitrate increase, even if sulfate levels are much smaller that than nitrate ones. Figure 8 Figure 7 actually shows a strikingly similar response of sulfate, SOA and ozone to NO<sub>x</sub> emission reductions (given that SO<sub>2</sub> and NMVOC emissions are held constant). Indeed the prevailing wintertime aqueous production of H<sub>2</sub>SO<sub>4</sub> requires oxidants and in particular ozone (Le et al., 2020; Sheng et al., 2018). In addition, the formation of SOA in both the gas and particulate phases also requires oxidants (Vahedpour et al., 2011; Huang et al., 2020; Feng et al., 2016; Li et al., 2011; Tsimpidi et al., 2010).

Pinder et al. (2008) also note an oxidant limitation for SIA formation over Eastern United states for the 2000 to 2020 period, but in their simulations, it mainly affects sulfate that increases as a result of NO<sub>x</sub> emission reductions while nitrate decreases. This is due to a more important sulfate to nitrate ratio in eastern US than over the Po basin. Fu et al. (2020) derive from combined measurements and modelling that wintertime nitrate during haze events in the North China Plain (NCP) are nearly insensitive to 30% NO<sub>x</sub> emission reductions, because increased ozone levels increase the NO<sub>x</sub> to HNO<sub>3</sub> conversion efficiency. Following these authors, this conversion also involves the homogeneous HNO<sub>3</sub> formation via the NO<sub>2</sub> + OH. This reaction also could play a role in the Po basin, in addition to the heterogeneous pathway. Also Leung et al. (2020) simulate that wintertime nitrate abatement in the NCP is

385 buffered with respect to emission reductions by increased oxidant build-up, but also by sulfate to nitrate conversion  
by liberating free NH<sub>3</sub> through sulfate concentration reduction, which can then enhance nitrate formation. Womack  
et al. (2019) find an oxidant limitation of nitrate formation over wintertime Utah (USA), and show that nitrate  
concentration diminishes when reducing VOC emissions.

## 5. Analysis of non-linearities

390 Clappier et al. (2020,2021) highlighted the specificities of the Po basin area within Europe. They showed that non-  
linearities are ~~important~~ present in this region. One peculiarity of the Po basin is a marked difference between the  
chemical regimes encountered within a confined area, ~~that~~ which have implications on the linearity of PM<sub>2.5</sub>  
responses to emission changes. In this section, we analyse in more details these non-linearities.

Information on the NO<sub>x</sub>-NH<sub>3</sub> interaction term at the reduction level  $\alpha=25\%$ :  $\hat{P}_{NO_xNH_3}^{25\%} [= P_{NO_xNH_3}^{25\%} - (P_{NO_x}^{25\%} + P_{NH_3}^{25\%})]$   
395 (first non-linear term in Eq. 4) is provided ~~both spatially and as a scatter plot~~ in Figure 8. ~~This~~ At  $\alpha=25\%$ , the  
interaction term is ~~almost constant (~-0.9) over~~ negative (or null) across the entire modelling domain, ~~(most data~~  
~~points are below the 1:1 line)~~ regardless of the chemical regimes ~~(-) and, and averages to approximately -10%, as~~  
~~indicated by the linear fit slope (= 0.9). In relative terms, this interaction term is also almost constant~~ regardless of  
400 the season ~~(only the yearly average is not shown in Figure 9, but the maps for both seasons are almost identical). At~~  
 ~~$\alpha=25\%$ , the interaction terms therefore represents approximately 10% of).~~

~~This negativity can be explained by the fact that a reduction of only NO<sub>x</sub> implies a reduction of both NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>~~  
~~and the same happens when reducing only NH<sub>3</sub>, therefore a simultaneous reduction of both precursors is lower than~~  
~~the sum of the two. the overall concentration change ( $P_{NO_xNH_3}^{25\%}$ ).~~ Single impacts would therefore lead to an  
overestimation (of about 10%) in PM<sub>2.5</sub> reduction if added ~~up~~ to extrapolate linearly the impact ~~of combined 25%~~  
405 ~~NO<sub>x</sub> and NH<sub>3</sub> emission reductions~~ on yearly averaged PM<sub>2.5</sub> concentrations ~~of~~. This result is expected for what  
concerns particulate NH<sub>4</sub>NO<sub>3</sub>, as a ~~combined NO<sub>x</sub>-NH<sub>3</sub> emission reduction at 25%-consequence of the gas/particle~~  
~~equilibrium described in Eq. 5, although non-linear relationships between NO<sub>x</sub> emissions and HNO<sub>3</sub> concentrations~~  
~~also play a role. Qualitatively, this behaviour is explained~~ negative interaction is also highlighted by the ~~hyperbolic~~  
~~shapes of the PM<sub>2.5</sub> isopleths determined for 3 -different sites of the domain (Figure 6). Their hyperbolic shapes~~  
410 ~~indicate this negative non-linearity in the NH<sub>3</sub>-NO<sub>x</sub> interaction.~~ As discussed in Annex 2, linearity would result in  
isopleths parallel to the descending diagonal lines.

When emission reductions increase from 25% to 50%, three additional non-linear terms are generated (three last  
terms in equation 4). Figure 9 and 10 provide an overview of these non-linear terms during wintertime and ~~summer~~  
~~times~~ summertime, respectively. The top left panel of each figure represents their sum, i.e. the total ~~nonlinearity~~ non-  
415 ~~linearity~~ generated between 25 and 50% ~~emission reduction~~. The right column shows the non-linearities  
associated to NO<sub>x</sub> and NH<sub>3</sub> while the bottom left panel reports the non-linear interaction between the two precursors.

Overall, non-linearities are more important during wintertime than during summertime. This is true both in absolute  
and relative (not shown) terms. Non-linearities tend to be the largest in between areas that are characterised by well-  
420 marked NH<sub>3</sub>- or NO<sub>x</sub>-sensitive regimes (indicated by the blue and red drawn contours). This can be explained by the  
fact that when one of the two components (NO<sub>x</sub> or NH<sub>3</sub>) is in large excess (compared to the other one), reductions of  
this compound have then ~~generally~~ only little impact, implying that both single and combined reductions only  
involve one compound and are therefore similar.

During wintertime, the overall non-linearity (Figure 9 top left) is ~~largely~~ dominated by the single NO<sub>x</sub>-related non-  
linearity ( $\hat{P}_{NO_x}^{50-25\%}$ , Figure 9 top right), a singularity in Europe as the Po basin is the only area where this occurs to  
425 this extent (Clappier et al., 2020,2021). In the region of Bergamo, the NO<sub>x</sub> non-linearity remains ~~relatively~~ weak  
despite the peculiar PM<sub>2.5</sub> responses to NO<sub>x</sub> emission reductions (i.e. an increase of PM<sub>2.5</sub> concentrations for NO<sub>x</sub>  
emission reduction up to 50%, Figure 7). In this NH<sub>3</sub> sensitive region, this behaviour can be explained by the strong  
oxidant limitation of HNO<sub>3</sub> formation outlined above (Figure 87). It is worth mentioning that although atypical, this  
behaviour is quasi-linear with PM<sub>2.5</sub> responses that remain proportional to the emission reduction strength ~~(up to~~  
430 ~~50%)~~ but with a negative slope. ~~Similarly to NO<sub>x</sub>, NH<sub>3</sub> single non-linearities (Figure 9, bottom right) are positive but~~

weaker, indicating slightly larger potential impacts for emission reductions in the range 25-50% than in the 0-25% range. Finally, the  $\text{NO}_x\text{-NH}_3$  non-linear interaction terms (Figure 9, bottom left) are mostly negative, i.e. the sum of impacts resulting from single reductions exceeds the impact of indicating that the combined reduction as already observed for the 25% interaction term for 50% emission reductions, is more negative than the corresponding term for 25%, pointing out to a strengthening of the  $\text{NO}_x\text{-NH}_3$  non-linearity when more intense emission reductions are considered.

In relative terms, the overall wintertime non-linearity terms increase by about 30 % when emission reductions increase from 25 and 50% (Figure 11 top, blue points). Note that these non-linearity terms reach larger values in some places of the modelling domain as highlighted by the data point dispersion. -In a previous work, Thunis et al. (2015) quantified the non-linearity of model responses to emission reductions in three areas in Europe, among which the Po Basin. One of their conclusions was that non-linearities remain relatively low for yearly averaged responses. Although the results presented here show important nonlinearities, these occur mainly during wintertime and are limited to specific areas. It is also worth noting that these non-linear responses (for moderate emission reductions up to 50%) only occur in the Po Basin (Clappier et al., 2021).

During summertime, the magnitude of non-linear terms are smaller than during wintertime. The overall non-linearity term (top left) is dominated by the  $\text{NH}_3\text{-NO}_x$  non-linear interaction term ( $\hat{P}_{\text{NO}_x\text{NH}_3}^{50-25\%}$ , Figure 10, bottom left). In summer,  $\text{PM}_{2.5}$  concentration is  $\text{NO}_x$  sensitive in almost all the domain (Figure 3) and emission reductions do not lead to shifts in the chemical regimes. In this case, the interaction terms become more important. The largest non-linear interaction terms appear in the western and northern part of the Po basin. They are negative everywhere in the Po basin, implying again that the sum of the impacts resulting from single interaction terms at 50% is more negative than at 25% emission reduction (reinforcement of the non-linearities when more intense emission reductions are considered). Most negative values appear in the western and northern part of domain.

In relative terms, the mean overall summer time non-linearity term increases by about 10 % when emission reductions increase from 25 and 50% (Figure 11 top, red data points). In contrast to wintertime, non-linearities are approximately constant throughout the domain as highlighted by the very low Root Mean Square (RMS) errors.

Figure 11 shows the increase of the overall non-linear terms for emission reduction increasing steps starting from different points, from 25 to 50%, from 50 to 75%, and finally from 75 to 100%. Regardless of the emission reduction increments step, summer time non-linearities remain small all over the domain with regression slope close to 0.9 and very limited data point dispersions (i.e. low RMSE). Wintertime non-linearities further increase significantly from 50% to 75% reduction levels (regression fit parameter close to 1.21) but tend to stabilize between 75 and 100% reduction level (regression fit parameter close to 1.05). It is interesting to note that potential impacts in winter increase for all segments (all winter points are above the 1:1 line) indicating that the same percentage reduction (25%) gains progressively more impact when more intense reductions are considered.

## 6. Discussion

When designing air quality plans, it is important to identify the key precursors on which to act in priority to hit a specific air quality target, but also to understand the consequences of these choices for various seasons (temporal variations), locations (spatial variability), emission reduction levels (strength) and strategies (combined or single emission reductions). As the information to make this decision is generally incomplete, assessing the robustness of the available model responses is essential. From the results presented here, a few key points appear.

- The seasonal and spatial variabilities in the response of  $\text{PM}_{2.5}$  to the reduction of  $\text{NO}_x$  and  $\text{NH}_3$  emissions are extremely large, with different and sometimes opposite responses to emission changes. Yearly averages do not represent the appropriate time window to evaluate the impact of such emission reductions and a focus on wintertime (November to February) seems to be the right option, especially because concentrations are larger during this period of the year.

- 480 • The responses of PM<sub>2.5</sub> to emission reduction plans that cover the whole area (i.e. uniform emission reductions are applied everywhere in the domain) vary from location to location: opposite responses occur within a few hundreds of kilometres for some reduction levels. In the region of Bergamo, PM<sub>2.5</sub> response to NO<sub>x</sub> emission reductions can be negative, meaning an increase of PM<sub>2.5</sub> when reducing the NO<sub>x</sub> emissions. It is important to combine NO<sub>x</sub> and NH<sub>3</sub> emission reductions in winter, or to go for stronger emission reductions to make sure these unwanted effects are limited.
- 485 • Despite quite important non-linearities, PM<sub>2.5</sub> responses to emission reductions are not chaotic. Indeed, regardless of the emission reduction level, the non-linear terms related to NH<sub>3</sub> emission reduction and to NO<sub>x</sub>-NH<sub>3</sub> interactions are quite uniform spatially. This is not the case of NO<sub>x</sub> emission reduction, for which care must be taken to ensure that the detailed response of PM<sub>2.5</sub> is captured.
- 490 • Although they are location ~~dependentspecific~~, PM<sub>2.5</sub> isopleths represent an interesting tool to assess the impact of different NO<sub>x</sub> and NH<sub>3</sub> emission reductions on PM<sub>2.5</sub> concentration. They indeed allow visualising in one single diagram the impact of any type of reductions on concentrations in a single grid cell (or set of grid cells). We must however remember that these isopleths derive from uniform emission reduction over the whole domain. Comparing sites where PM<sub>2.5</sub> responses to the same “domain-wide” policy are different, it appears challenging to define a single domain-wide policy efficiently reducing PM at all locations.

## 7. Conclusions

500 In this work, we analysed the ~~sensitivitiessensitivity~~ of PM<sub>2.5</sub> to NO<sub>x</sub> and NH<sub>3</sub> emissions by means of a set of EMEP simulations performed with different levels of emission reductions, from 25% up to a total switch-off of those emissions. Both single and combined precursor reduction scenarios were applied to determine the most efficient emission reduction strategies and quantify the interactions between NO<sub>x</sub> and NH<sub>3</sub> emission reductions. ~~TheOur~~ results ~~eenfirmedconfirm~~ the peculiarity of secondary inorganic PM<sub>2.5</sub> formation in the Po basin suggested by Clappier et al. (20202021), characterised by contrasting chemical regimes within distances of few (hundreds of) kilometres, as well as ~~strong~~ non-linear responses to emission reductions during wintertime. One of the striking results is the increase of the PM<sub>2.5</sub> concentration levels when NO<sub>x</sub> emission reductions are applied in NO<sub>x</sub>-rich areas, such as the surroundings of Bergamo. The isopleths in the graphs showing PM<sub>2.5</sub>, nitrate, sulfate, SOA and O<sub>3</sub> concentrations as a function of NH<sub>3</sub> and NO<sub>x</sub> emissions (Figure 7) indicate that the increased oxidative capacity of the atmosphere is the cause of the increase of PM<sub>2.5</sub> induced by a reduction in NO<sub>x</sub> emission ~~of~~ up to -50%. This process can have contributed to the absence of significant PM<sub>2.5</sub> concentration decrease during the COVID-19 lockdowns in many European cities (EEA, 2020; Putaud et al., 2020). It is important to account for this process when designing air quality plans, since it could well lead to transitional increases in PM<sub>2.5</sub> at some locations in winter as NO<sub>x</sub> emission reduction measures are gradually implemented. At this type of location, mitigation measures that would be optimal in the long-term might not be efficient in the short-term.

515 Joint ~~analysisanalyses~~ of PM sensitivity to emissions and the G-ratio ~~and emissions sensitivities~~ can give a clue if a NH<sub>3</sub> sensitive chemical regime is due to either a limitationlack of NH<sub>3</sub> or to a lack of sensitivity of NO<sub>x</sub> emission reduction, due to a non-linear and opposite behaviournegative response of ~~the NO<sub>x</sub>—HNO<sub>3</sub> relationship.~~ concentration to NO<sub>x</sub> emissions. In this latter case, the chemical regime is NH<sub>3</sub> sensitive in terms of NH<sub>3</sub> and NO<sub>x</sub> emission reductions, but it can be HNO<sub>3</sub> limited in terms of G-ratio, as observed for the Bergamo – Milano region. Inversely, a positive-G-ratio greater than 1, indicating HNO<sub>3</sub> limitation to particulate nitrate formation, does not necessarily indicate a larger sensitivity to NO<sub>x</sub> than to NH<sub>3</sub> emissions. Thus, ~~theour~~ results ~~indicateshow~~ the impossibility to directly use the G-ratio for air quality management, an interesting result in itself. ~~WhereasWhile~~ PM<sub>2.5</sub> chemical regimes (determined by the relative importance of the NO<sub>x</sub> vs. NH<sub>3</sub> responses expressed as potentials (i.e. responses projected linearly to a 100% to emission reductions) of NO<sub>x</sub> and NH<sub>3</sub> emission reduction show large variations seasonally and spatially, they are quite insensitive to the emission reduction levels at least up to -50% because the secondary aerosol formation chemical regimes are not modified by those relatively moderate ranges. The response of they are not very sensitive to moderate (up to 50%) emission reductions. Beyond 25% emission reduction strength, responses of PM<sub>2.5</sub> concentrations to NO<sub>x</sub> emission reductions is found to become

530 non-linear in certain areas of the Po basin, ~~especially in winter. However, non-linear terms remain quite constant spatially (although they depend on the emission reduction strength), which suggests that potential approaches to model or parameterize these non-linearities might not be too complex to develop. These parametrisations might be useful to interpolate existing modelling responses to situations that are not explicitly modelled, or for their implementation in source-receptor models, mainly during wintertime.~~

535 Since sulfate concentrations are low in the Po basin, the impact of SO<sub>2</sub> emission reductions was not evaluated here. However, the simulations performed in Clappier et al. (2020/2021) indicate that air quality improvement plans addressing SO<sub>2</sub> emissions may still lead to additional PM<sub>2.5</sub> decreases. Further works should also test if NMVOC emission would further affect the concentration of oxidants, and subsequently of nitrate (and sulfate) during winter. This depends on the fraction of ozone formed photochemically in the Po basin, compared to the one transported from outside by advection or entrainment.

540 Finally, it would be important to compare the results obtained in this work from the EMEPrv4\_17 model with similar results obtained from other models. With its complex setting, the Po basin represents a good test case for such inter-comparisons.

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### Annex 1: Selection of seasons

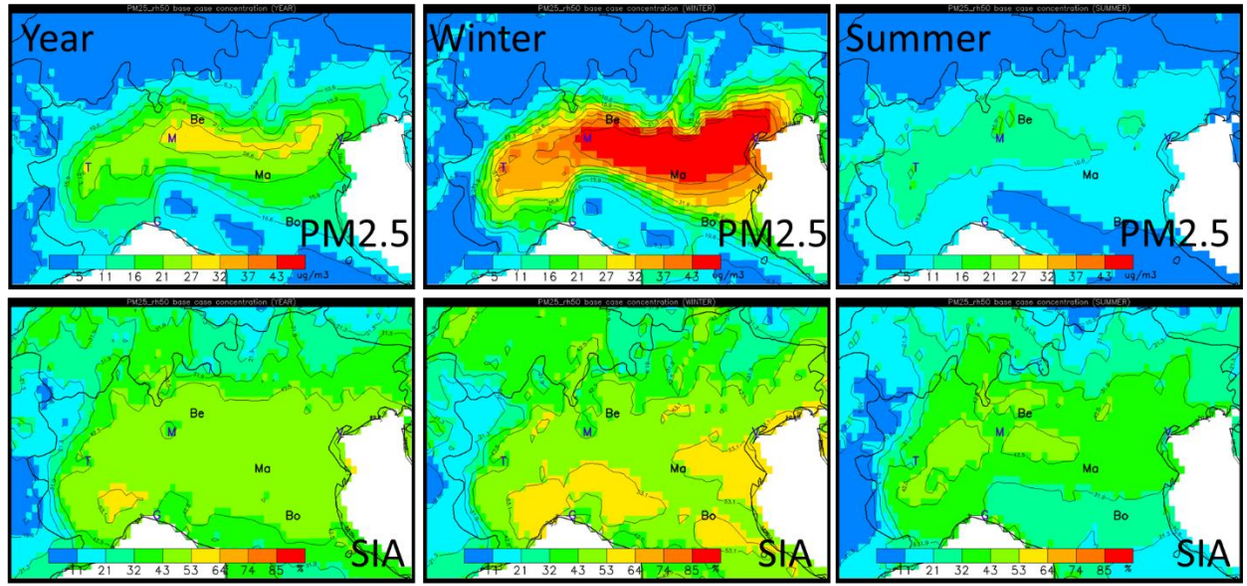
725 Chemical regimes show a great variability with time. To select the extension of the seasons, we analyse the monthly  
behaviour of the NH<sub>3</sub> and NO<sub>x</sub> responses as well as the interaction terms (Figure 12). The analysis is performed for  
two locations: Bergamo which lies in a NH<sub>3</sub> sensitive zone and Mantova which lies within a NO<sub>x</sub> sensitive zone. On  
these plots, we identify two major seasons with a consistent behaviour: winter from November to February and  
summer, from April to September. These two seasons are similar at both locations. The two remaining months,  
730 ~~April~~ March and October are transition months and are not considered in the analysis. It is interesting to note that  
these two transition periods correspond more or less to the switching time between the NO<sub>x</sub> and NH<sub>3</sub> concentration  
time profiles (Figure 12 - right panel). On the latter figures, the SO<sub>2</sub> and NO<sub>2</sub> temporal evolutions are almost  
identical, in contrast to NH<sub>3</sub>.

### Annex 2: A theoretical example for the isopleths

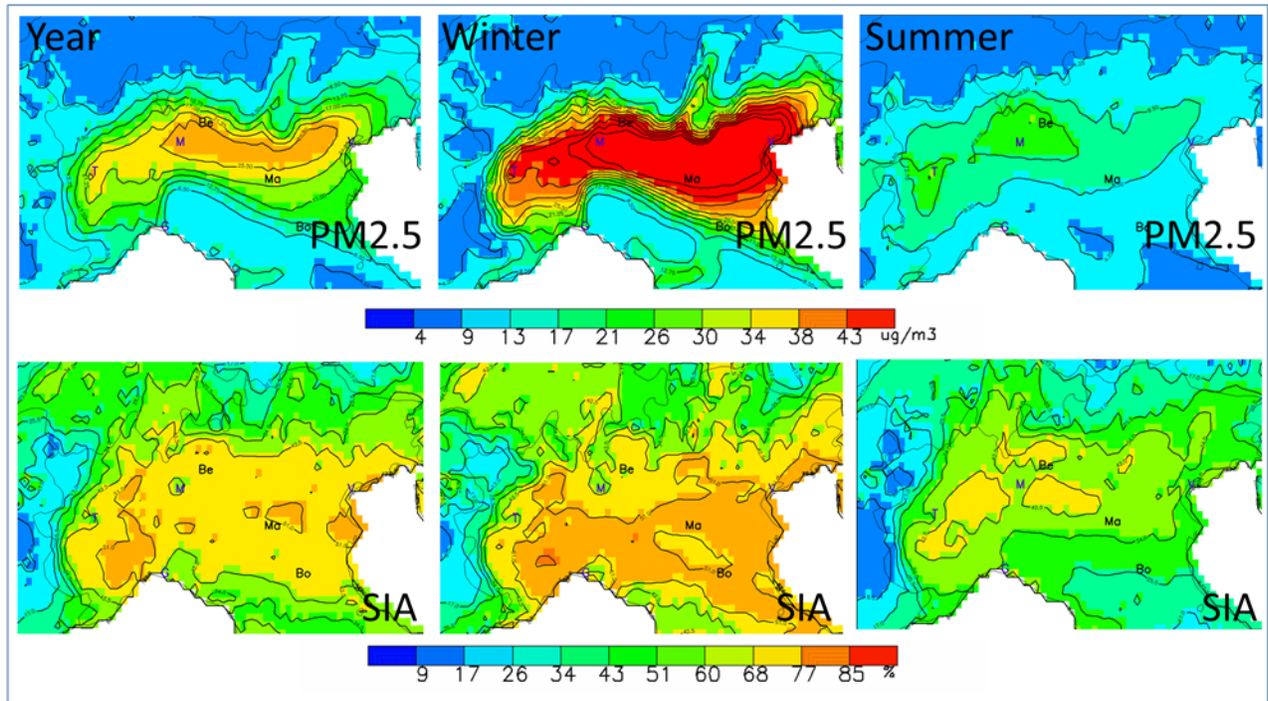
735 To facilitate the interpretation of the isopleths diagrams, we use a simple theoretical example that mimics the  
complex reactions process schematized by equations (5) and (6) above. Our simplified process is described by the  
following relation:  $C_c(x, y) = \min(E_A[x], E_B[y])$  where  $C_c$  is the concentration of a compound “c” that is given by  
the minimum between two emitted species A and B. The concentration depends on the strengths of these emissions,  
specified by the parameters x and y. For each emission strength (x or y), the two emission species are proportional to  
740 their full-scale value (100%):  $E_A(x) = xE_A[100]$  and  $E_B(y) = yE_B[100]$ , respectively. If we choose  $E_A(100) \gg$   
 $E_B(100)$ , we create a B-sensitive environment (Figure 13 - left column) and inversely (Figure 13 - right column). If  
we select mixed situations, representing for example an average of days, during which we alternate between A-  
sensitive and B-sensitive regimes, we obtain the two bottom isopleth diagrams that represent cases where a larger  
number of A-sensitive (right) or B-sensitive (left) events are recorded. Although extremely simple, these diagrams  
745 illustrate properties that are observed on the real test cases.

Let’s take the example of a A-sensitive regime. Similar observations can be made in the case of a B-sensitive  
regime. We note the following points:

- 750 • The diagram area can roughly be divided into two zones separated by a ridge: a top-left triangle where the  
sensitivity to emission reductions of species “AB” dominate and a bottom-right triangle where the sensitivity to  
BA dominates. The slope of the ridge (larger or less than one) informs on the type of regime.
- 755 • In the case of a single A-sensitive day (top right) with  $E_A E_B(100) = 2E_B 2E_A(100)$ , the concentration of compound  
C remains unchanged for emission reductions of AB up to 50% while its concentration react in a linear way to  
emission changes of BA from 0 to 100%. ~~As concentration must drop to zero for full reductions~~ Between the  
base case and a reduction level of either 50%, the A or B, a lower gradient for B than A for small emission  
760 reductions implies a reversed behaviour (is therefore larger than the B-gradient for B than A) for . This implies  
that the B-gradient is larger reductions than the A-gradient between the 50 and 100% reduction levels because  
we know that for a 100% reduction of A or B, the concentration must be zero. In our simple example, the  
gradient of B is zero from 0 to 50% but is twice as large as A ~~from 0~~ between 50 to 50% emission  
strength 100%.
- While the combination of several events (e.g. days) characterised by different regimes leads to smoother  
isopleths (bottom), the same characteristics can be noted. In particular, the inclination (tending to the horizontal  
or vertical) provides information on the type of chemical regime.

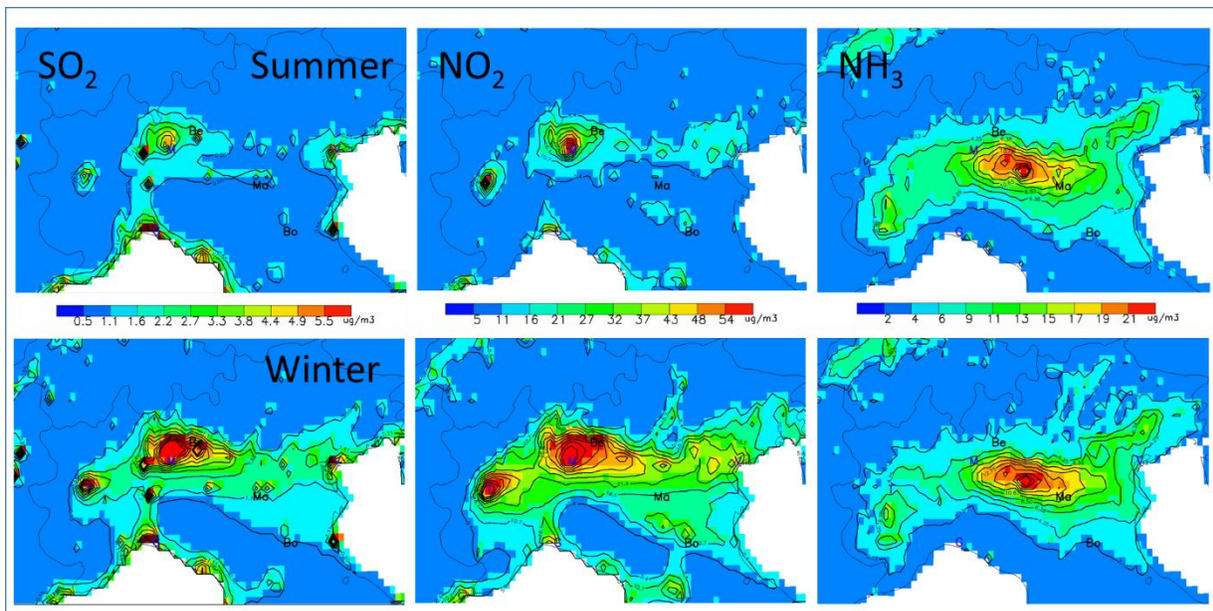


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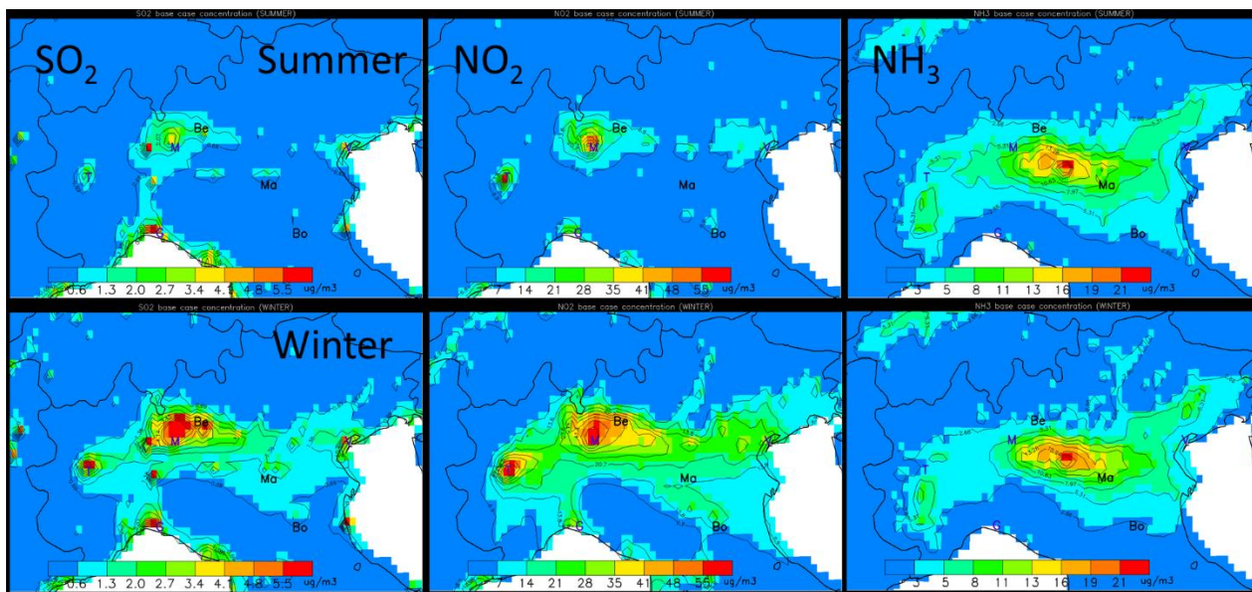
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Figure 1: Seasonal variations of the base case PM<sub>2.5</sub> (in  $\mu\text{g m}^{-3}$ , top row) and SIA (in % of the PM<sub>2.5</sub> concentration, bottom row). The symbols: Be, Ma and Bo represent the locations selected for more detailed analysis: Bergamo (Be), Mantova (Ma) and Bologna (Bo). Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.



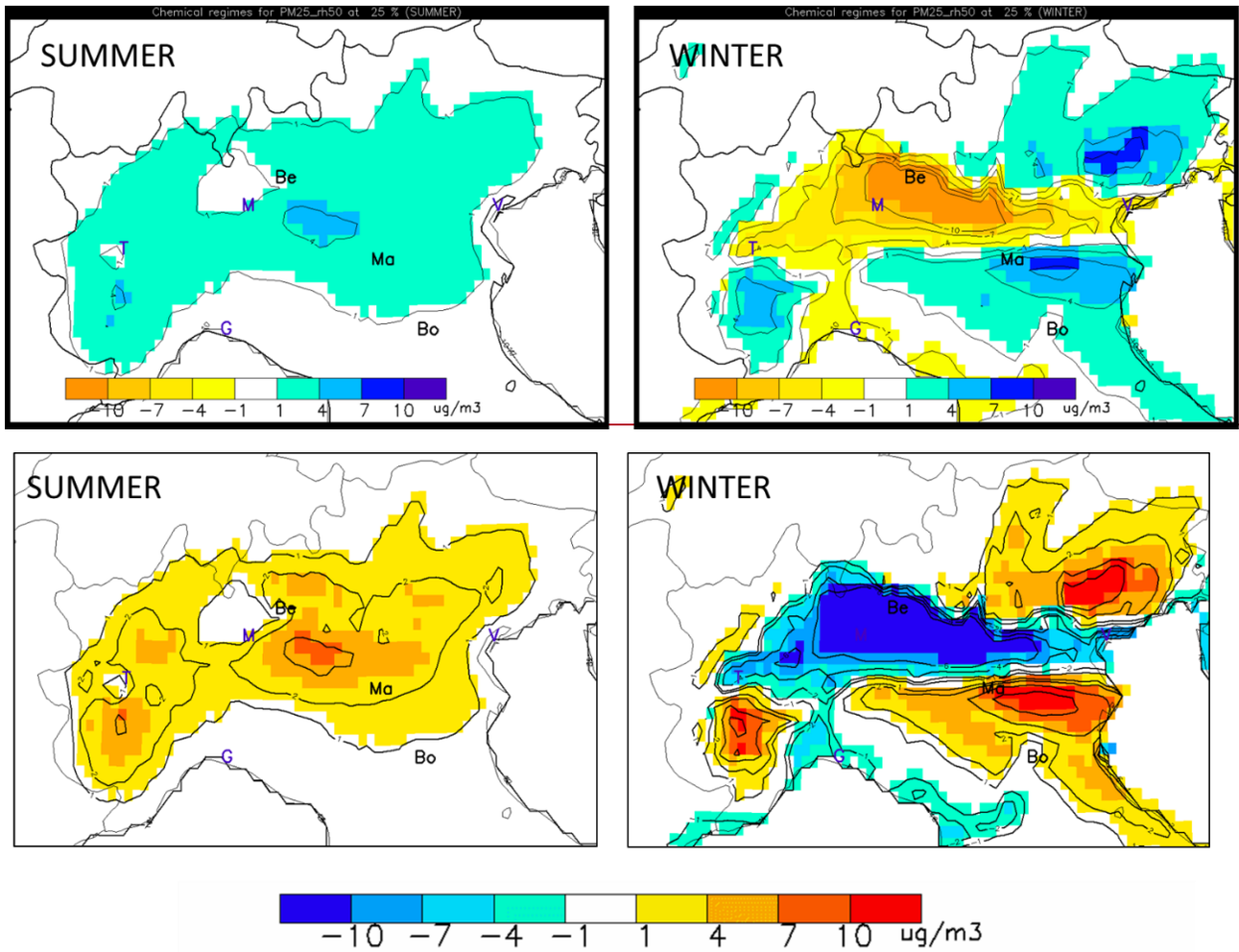
**Figure 2: Summer (top) and winter (bottom) concentration base-case fields for SO<sub>2</sub> (left column), NO<sub>2</sub> (central column) and NH<sub>3</sub> (right column) expressed in in  $\mu\text{g m}^{-3}$ . The symbols: Be, Ma and Bo represent the locations selected for more detailed analysis: Bergamo (Be), Mantova (Ma) and Bologna (Bo). Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.**

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**Figure 2: Summer (top) and winter (bottom) concentration base-case fields for SO<sub>2</sub> (left column), NO<sub>2</sub> (central column) and NH<sub>3</sub> (right column). The symbols: Be, Ma and Bo represent the locations selected for more detailed analysis: Bergamo (Be), Mantova (Ma) and Bologna (Bo). Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.**

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Figure 3: Winter (right) and Summer (left) chemical regimes obtained ~~at~~ from an emission reduction level of  $\alpha=25\%$ . The maps represent the  $P_{NO_x}^{25\%} - P_{NH_3}^{25\%}$  (unit:  $\mu\text{g}/\text{m}^3 \text{ m}^{-3}$ ) indicator that shows the  $\text{NO}_x$ - and  $\text{NH}_3$ -sensitive areas in ~~blue~~ blue and yellow/red and blue, respectively. The symbols Be, Ma and Bo indicate the location of Bergamo, Mantova and Bologna, respectively. Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.

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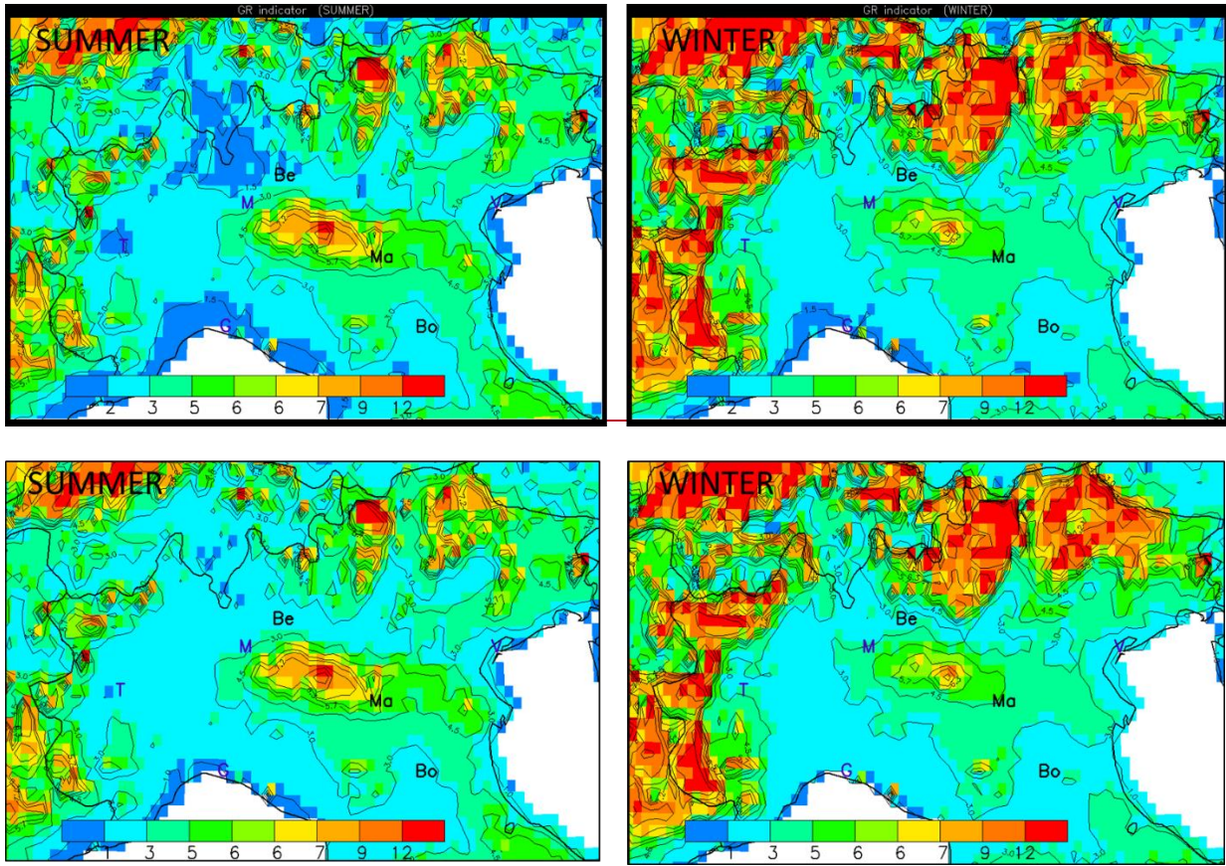
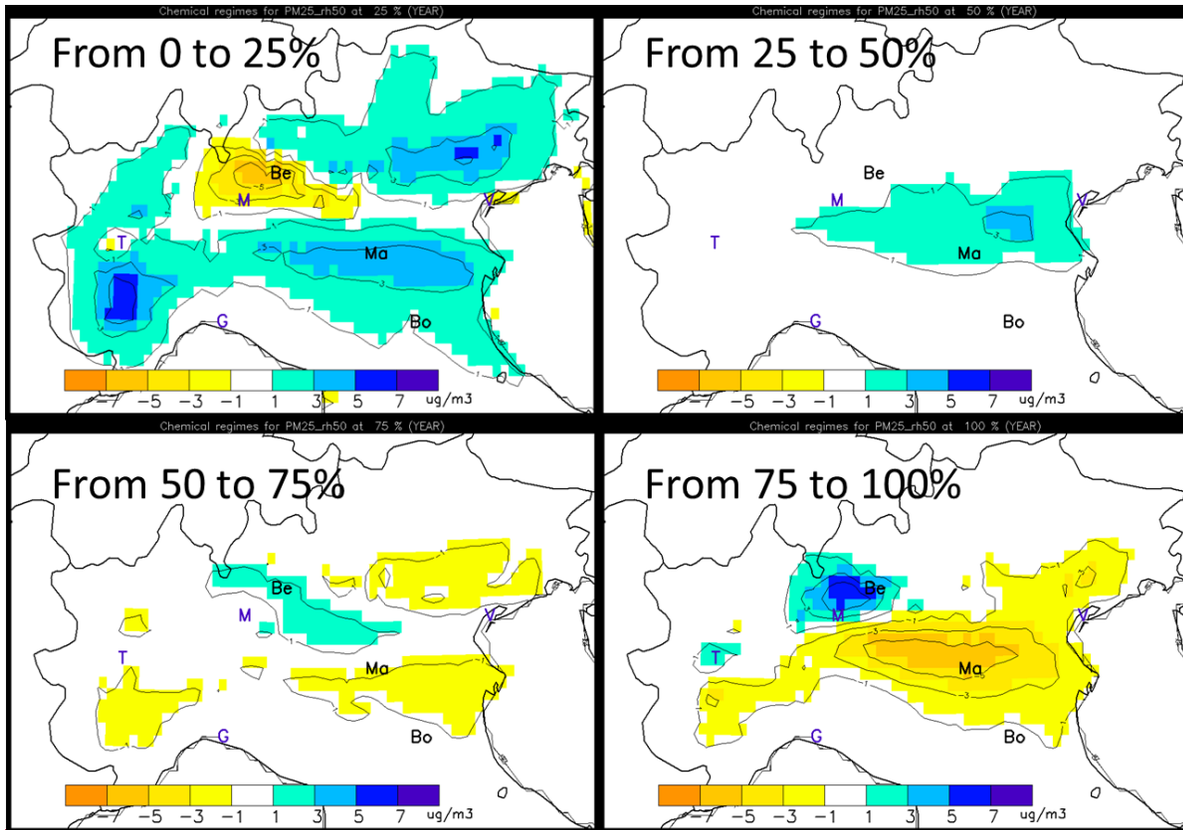
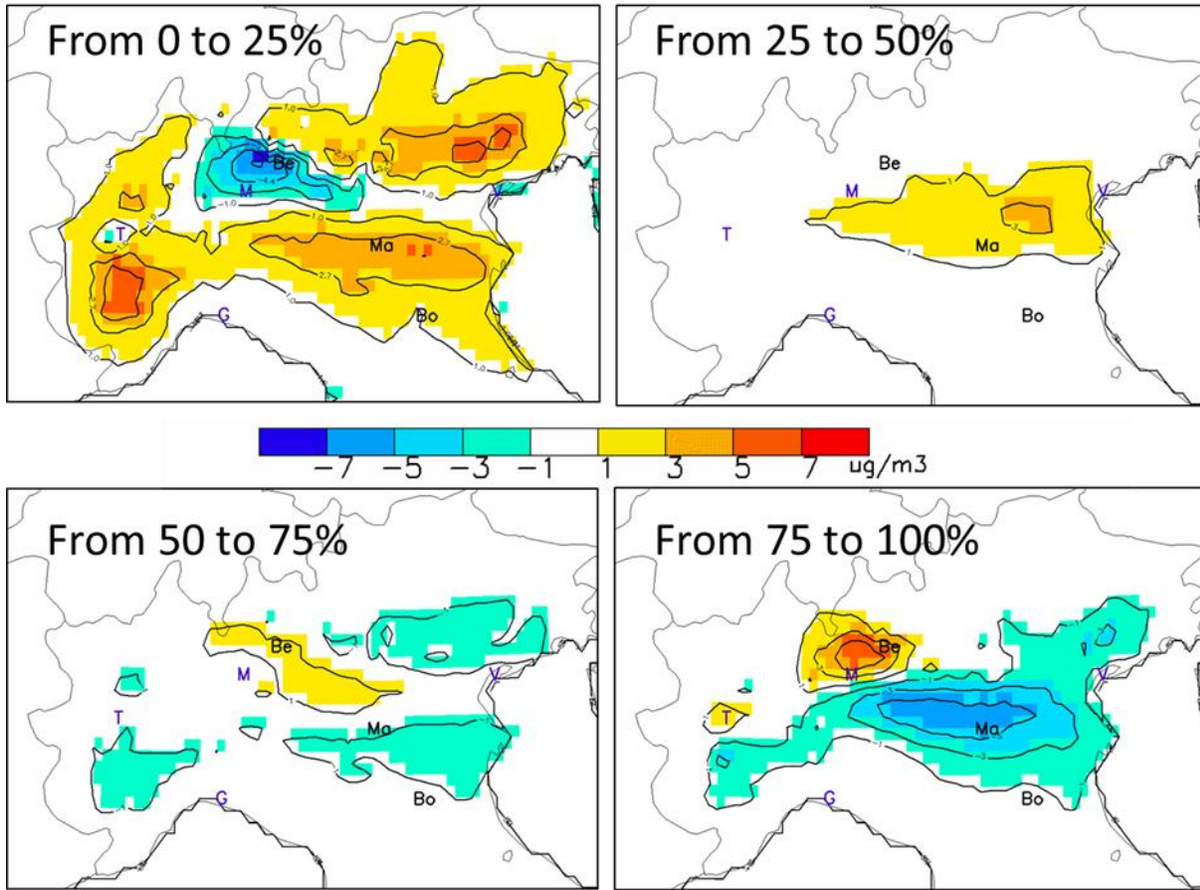


Figure 4: G-ratio for winter (right) and summer (left) times. The symbols Be, Ma and Bo indicate the location of Bergamo, Mantova and Bologna, respectively. Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.

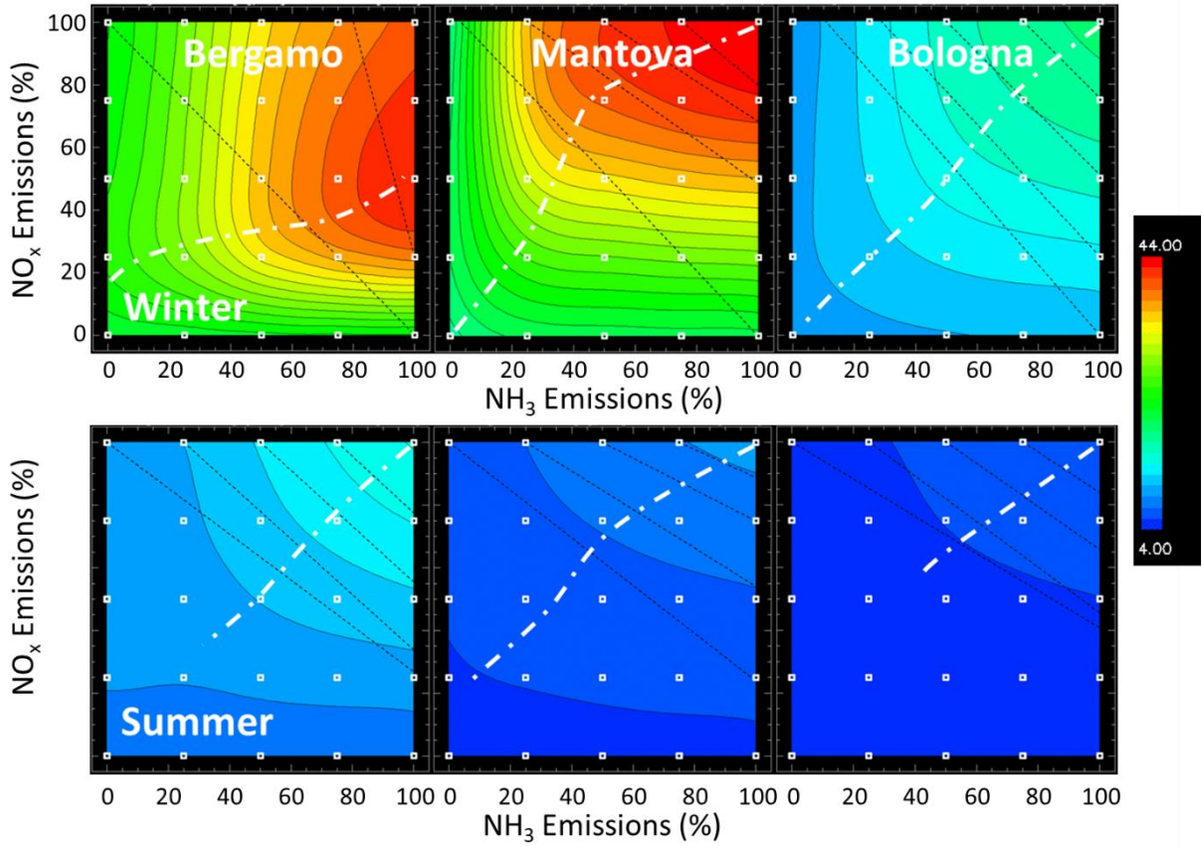
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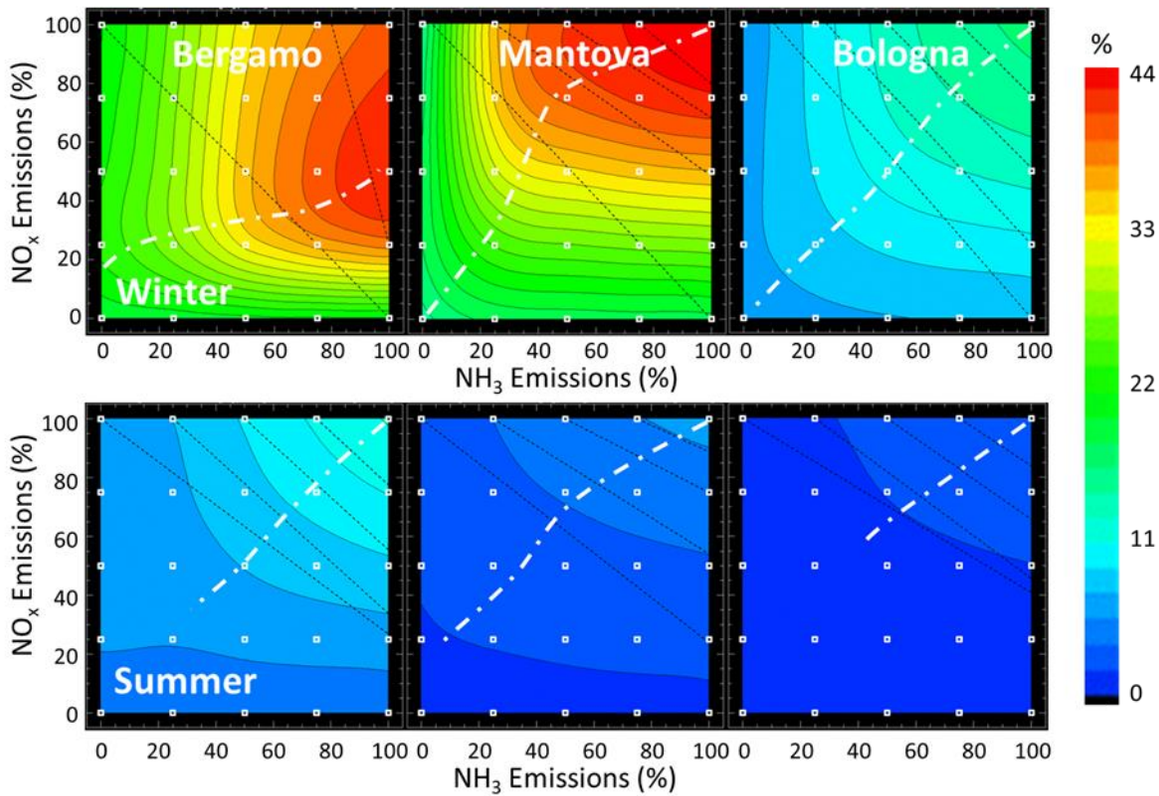




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Figure 5: Yearly averaged chemical regimes obtained ~~for~~from a 25% emission reduction starting at different levels of emissions corresponding ~~to~~to  $\alpha=0, 25, 50$  and 75. The maps represent the  $(P_{NO_x} - P_{NH_3})$  between the starting and ending levels (unit:  $\mu\text{g}/\text{m}^3 \text{ m}^{-3}$ ) showing the NO<sub>x</sub>- and NH<sub>3</sub>-sensitive areas in ~~blue and~~ yellow/~~red and~~ blue, respectively. The symbols Be, Ma and Bo indicate the location of Bergamo, Mantova and Bologna, respectively. Other cities are indicated by their first letter for convenience: Venice, Milan, Turin and Genova.

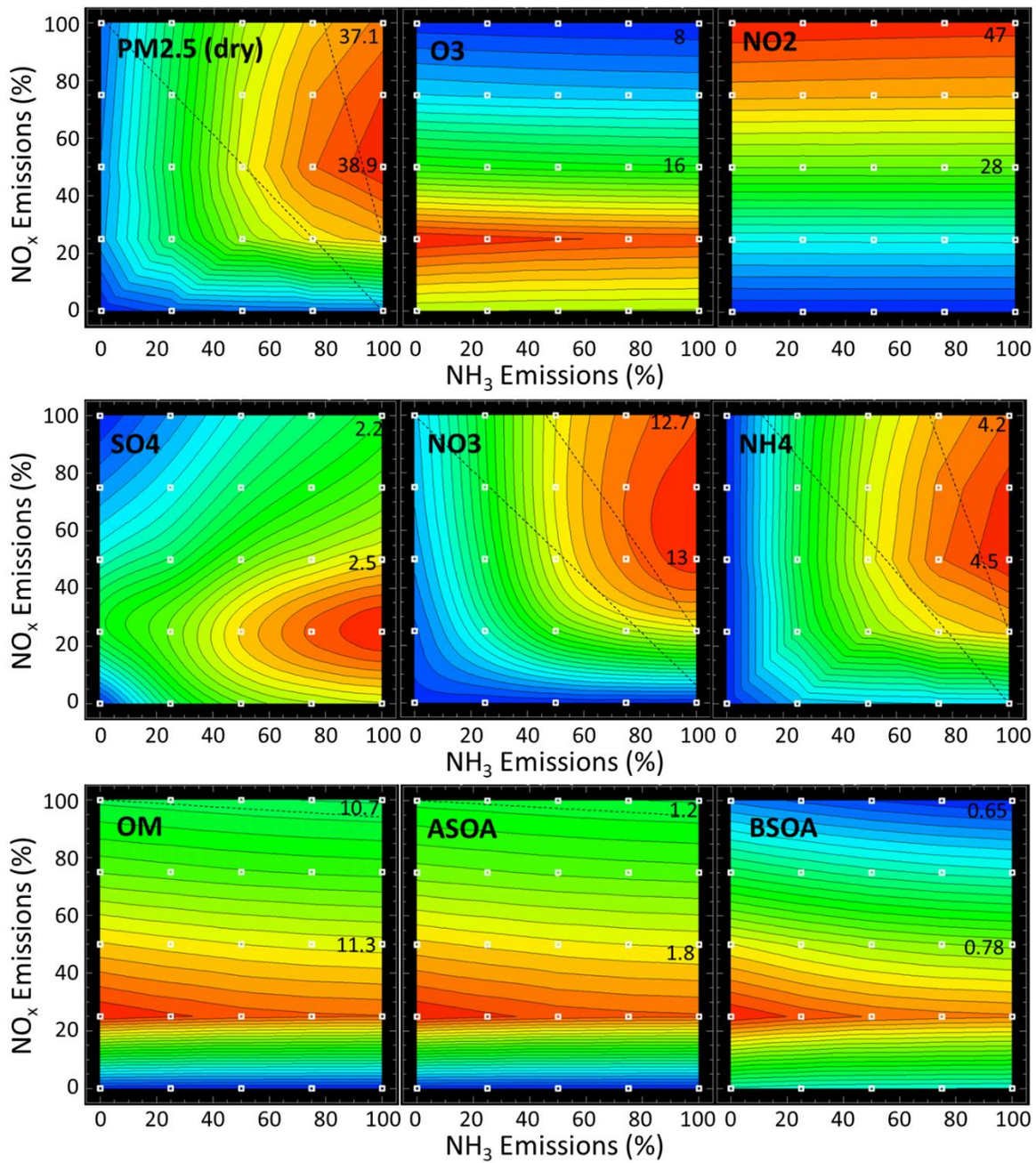


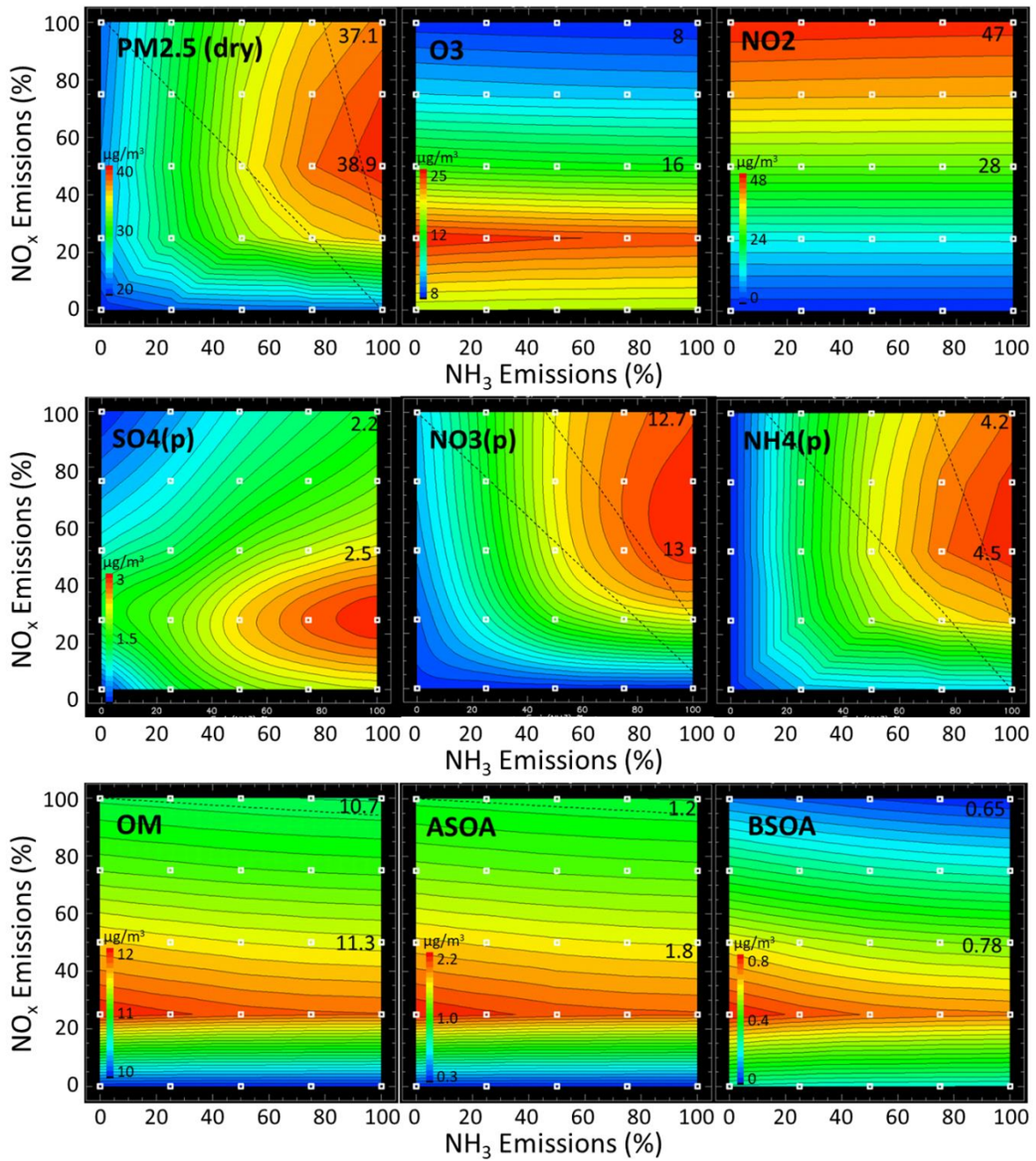


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Figure 6: PM<sub>2.5</sub> isopleths during Winter (top row) and Summer (bottom row) at the three locations of interest (see maps). PM<sub>2.5</sub> concentrations are expressed in  $\mu\text{g m}^{-3}$  as a function of the intensity of the NO<sub>x</sub> (Y-axis) and NH<sub>3</sub> (X-axis) emissions, respectively. The overlaid dashed oblique lines on each diagram connect similar PM<sub>2.5</sub> concentration values for single NO<sub>x</sub> and NH<sub>3</sub> reductions. The more vertical are these lines, the larger is the NH<sub>3</sub> abatement impact compared to the NO<sub>x</sub> abatement impact; The more horizontal they are, the larger is the NO<sub>x</sub> abatement impact compared to the NH<sub>3</sub> abatement impact. The dashed line delineates the ridge between the NH<sub>3</sub> and NO<sub>x</sub> sensitive areas.

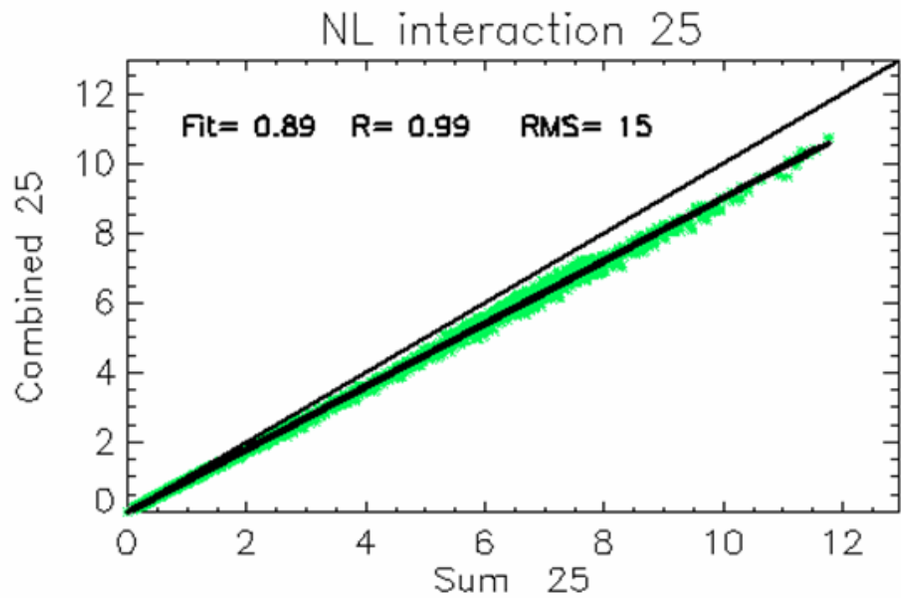
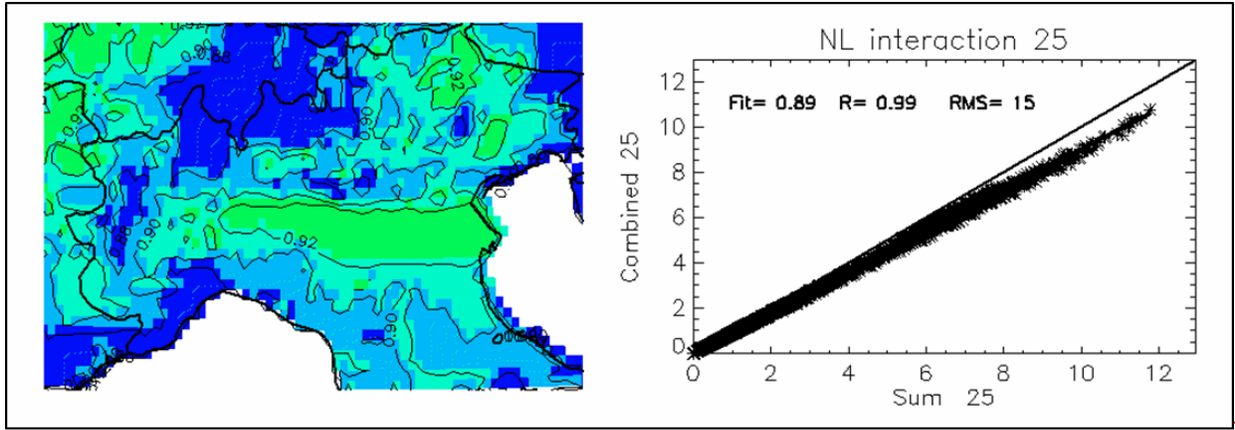
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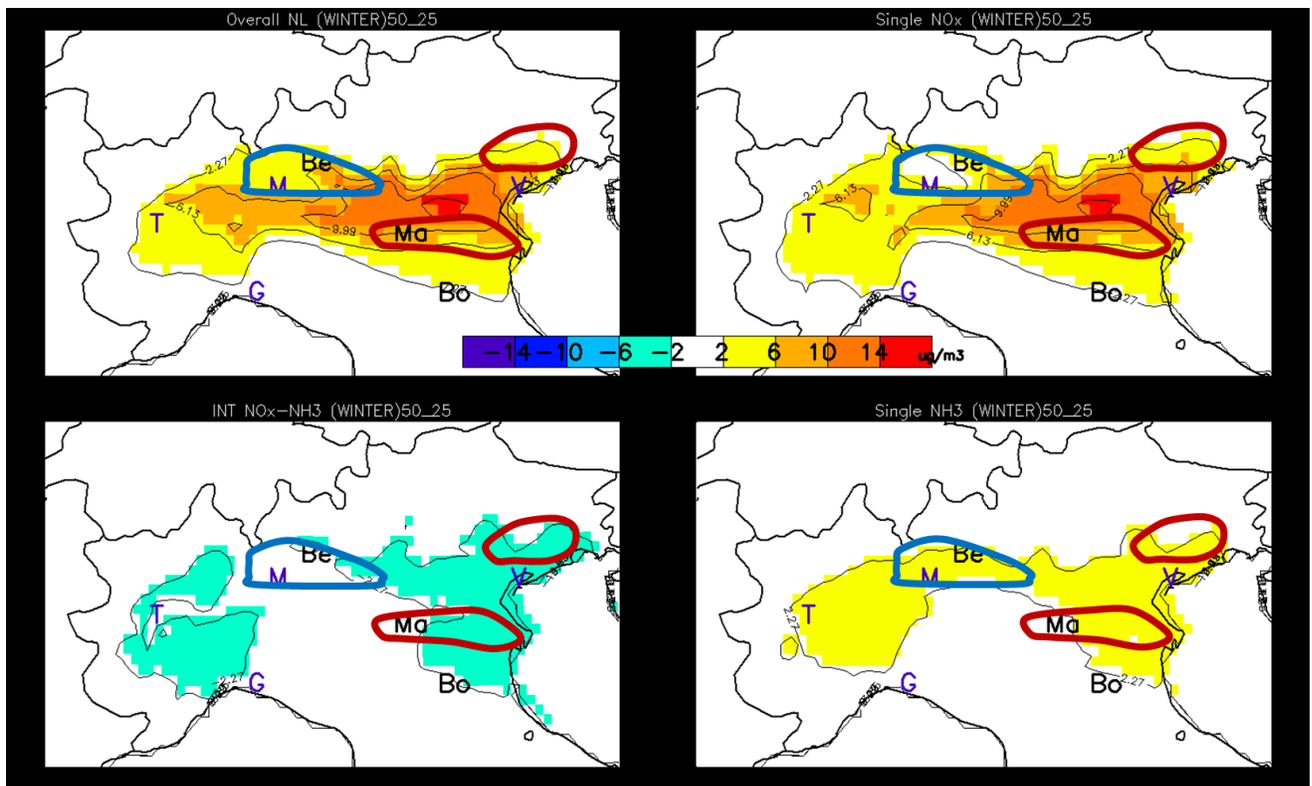
Figure 7: Wintertime isopleths in Bergamo for the species: dry PM<sub>2.5</sub>, O<sub>3</sub>, NO<sub>2</sub>, sulfate (SO<sub>4</sub>), nitrate (NO<sub>3</sub>), ammonium (NH<sub>4</sub>), organic matter (OM), anthropogenic and biogenic secondary aerosols (ASOA and BSOA, respectively). The two numbers on the vertical axis indicate concentration values for the base case and at the 50% NO<sub>x</sub> emission level.



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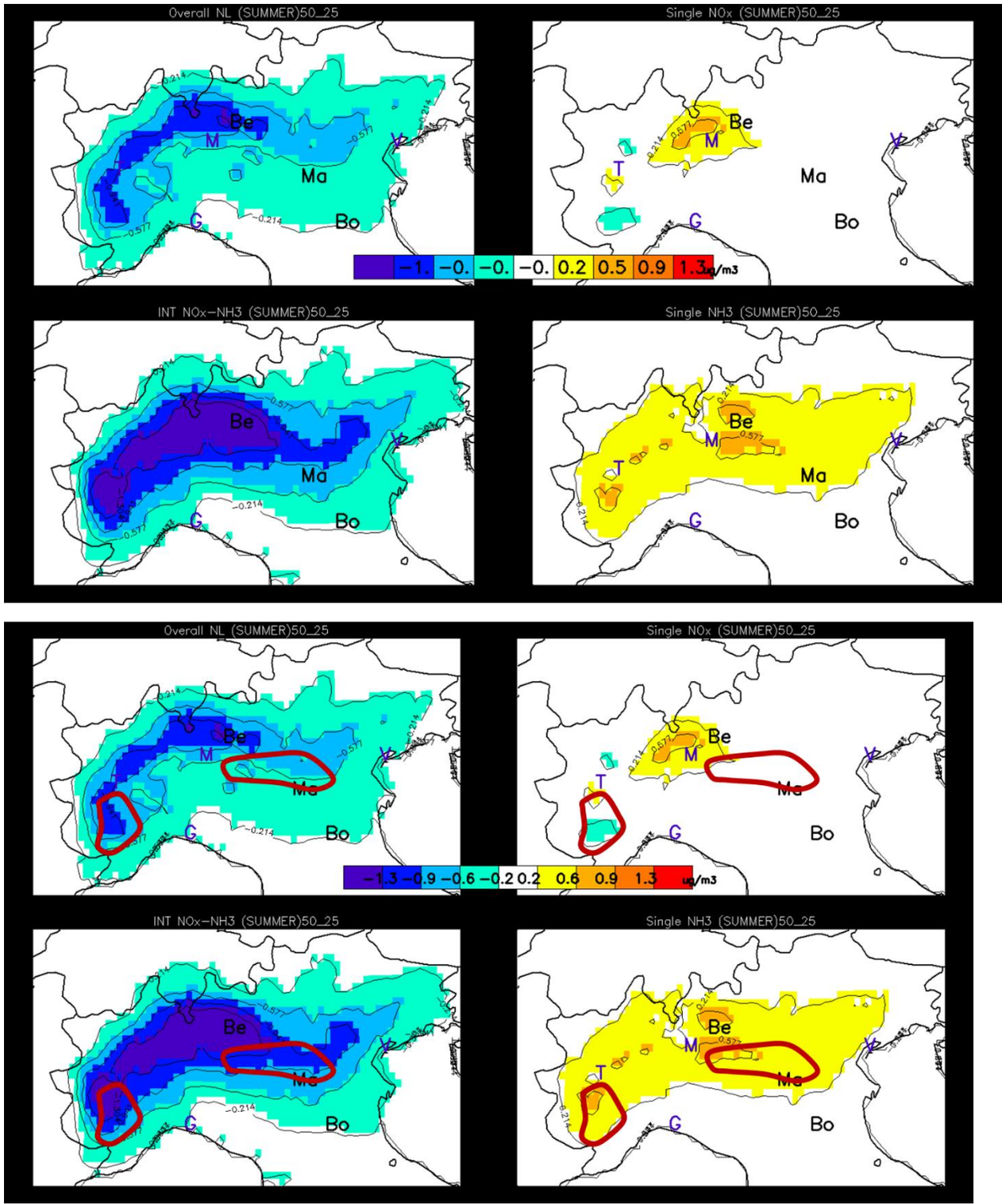
Figure 8: Yearly mean non-linear interaction term for a 25% reduction in NO<sub>x</sub> and NH<sub>3</sub>. The **map (left) shows the ratio  $\frac{p_{NO_xNH_3}^{25\%}}{p_{NO_x}^{25\%} + p_{NH_3}^{25\%}}$  whereas the scatter plot compares the sum of the potential impacts of the two single reductions (X-axis) with the potential impact of the combined emission reduction ( $\mu\text{g}/\text{m}^3\text{unit: } \mu\text{g m}^{-3}$ ) (Y-axis). Departure from the 1:1 line quantifies the overall non-linearity. Each data point represents the yearly average values for a grid cell in the modelling domain.**

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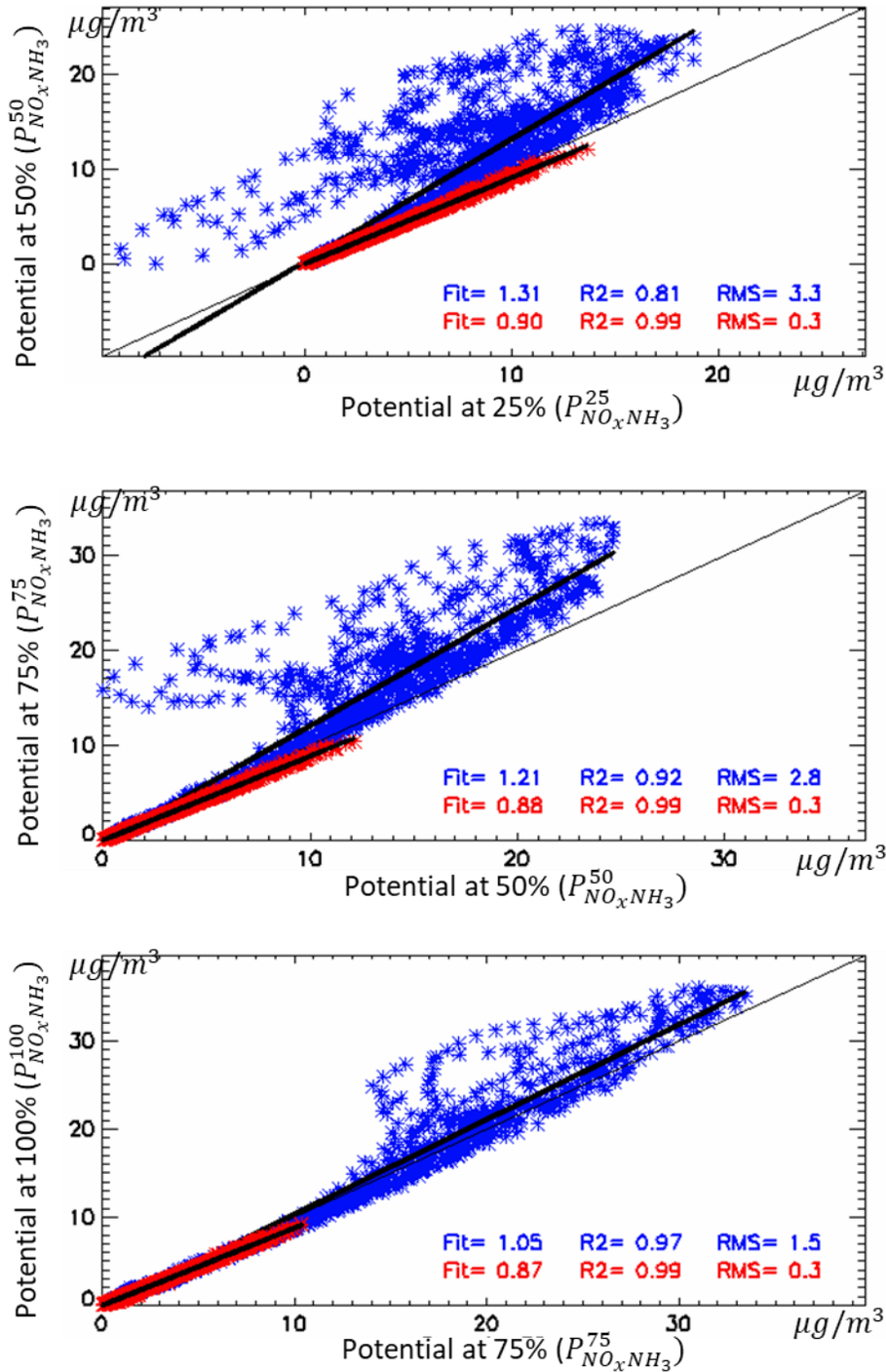
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Figure 9: Wintertime maps of the overall non-linearity term (top left) and of its components: expressed as potential impacts between 25 and 50%: the single NO<sub>x</sub> non-linearity term (SNO<sub>x</sub>—top right), the single NH<sub>3</sub> non-linearity term (SNH<sub>3</sub>—bottom right) and the NO<sub>x</sub>-NH<sub>3</sub> interaction term (INO<sub>x</sub>-NH<sub>3</sub>—bottom left). The three locations of interest: Bergamo, Mantova and Bologna are indicated by their first two letters while other cities are indicated with their first letter for convenience (Venice, Milan, Turin and Genova). The hand drawn contours roughly indicate the central position of the NH<sub>3</sub> (blue) and NO<sub>x</sub> (red) sensitive regime areas (see Figure 3 - right).



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Figure 10: same as Figure 9 but for summertime.



840 Figure 11: Changes of the overall non-linearity terms from 25 to 50% (top), from 50 to 75% (middle) and from 75 to 100% (bottom) in  $\text{NO}_x$  and  $\text{NH}_3$ . The overall non-linearity is visualised as the distance from the 1:1 diagonal, i.e. the difference between the overall potential impact at two levels of emission reduction, X- and Y-axis. Each point represents one land grid cell within the domain for wintertime (blue) and summer time (red). The “fit” parameter indicates the slope of the regression line while R2 and RMS provide information on the coefficient of determination and the root mean square error, respectively.

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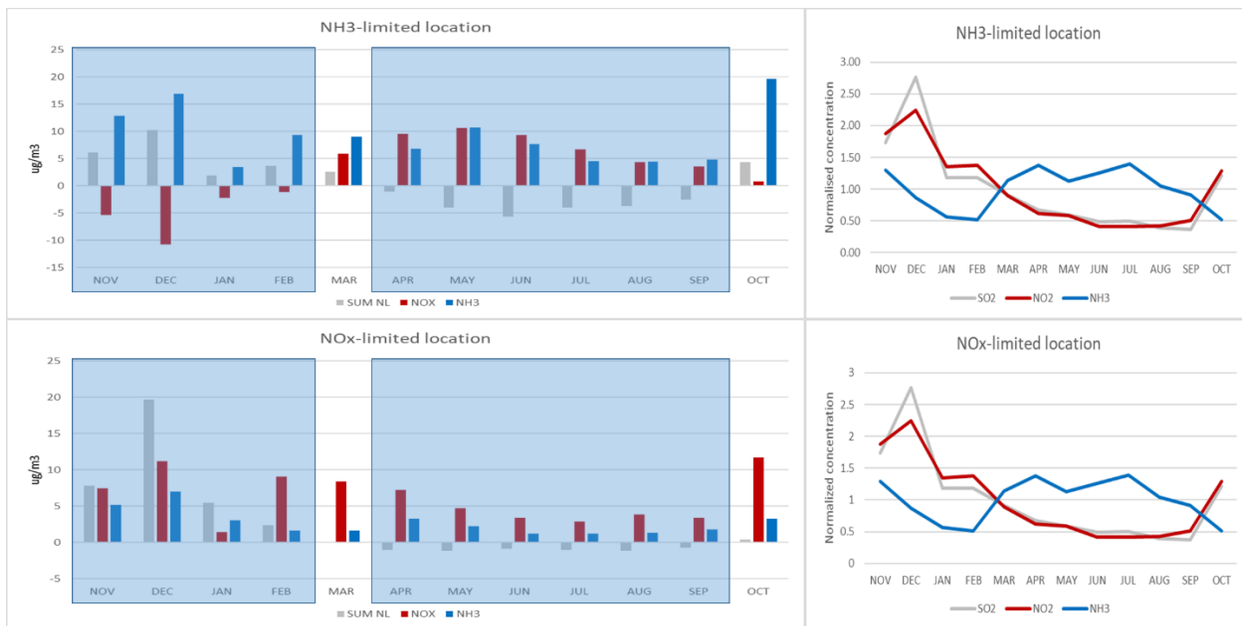
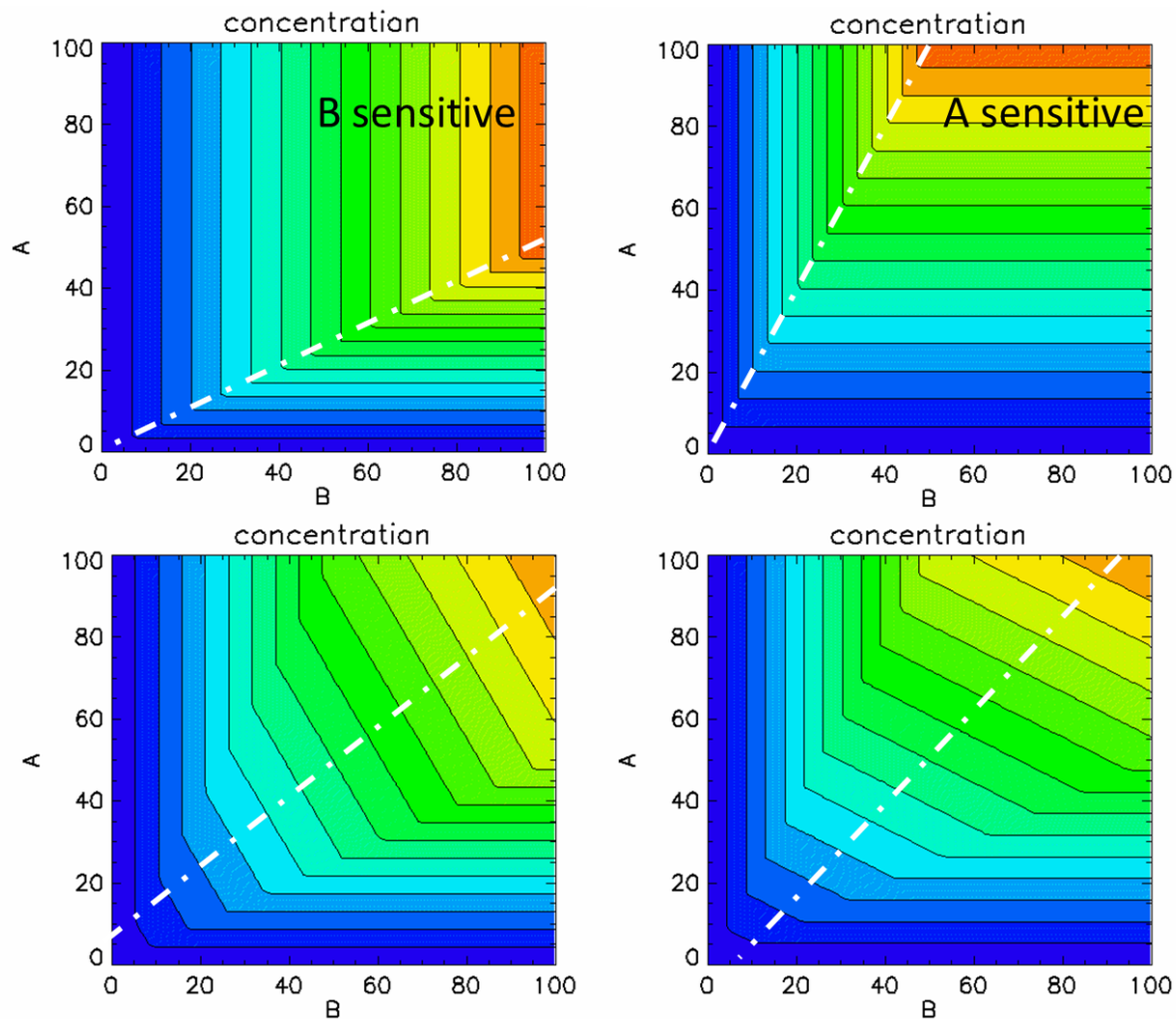


Figure 12: Monthly averaged responses to  $\text{NH}_3$  (blue),  $\text{NO}_x$  (red) reductions (25%) and interaction terms (grey) at two locations: Bergamo (top) and Mantova (bottom). The right panel shows the monthly evolution of the concentrations of  $\text{NO}_2$ ,  $\text{NH}_3$  and  $\text{SO}_2$  at those two locations. Note that concentrations are normalised by their average values.



850 **Figure 13: Isopleths for a simple theoretical system consisting of two emission precursors (A and B) competing through non-linear reactions to the concentration of a pollutant. See details in the text.**

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