

participated in the EMEP and laboratory intercomparison and the laboratories generally have a satisfactory quality.

2.2 Nitrogen concentrations in air

Altogether fourteen stations have delivered data for nitrogen dioxide (NO_2) and thirteen total reduced nitrogen ($\text{NH}_3+\text{NH}_4^+$) and total nitrate ($\text{HNO}_3+\text{NO}_3^-$). Annual averages of the different nitrogen species are presented in Figure 2.2. The lowest concentrations for all the three nitrogen species were reported at the northernmost Swedish site (SE05) in 2013. The concentrations were 0.16, 0.04 and 0.08 $\mu\text{g N}/\text{m}^3$ for respectively $\text{NH}_3+\text{NH}_4^+$, $\text{HNO}_3+\text{NO}_3^-$ and NO_2 at this site. Highest concentrations of nitrogen in air were found at the Danish sites with annual concentration means of 2.78 $\mu\text{gN}/\text{m}^3$ for NO_2 , and 0.85 $\mu\text{gN}/\text{m}^3$ for sum nitrate at Risö (DK12), and 1.93 $\mu\text{gN}/\text{m}^3$ for sum ammonium at Tange (DK03). Data for particulate nitrate and ammonium from the German site (DE09) is not included in the figures below, because it is not comparable with the sums shown there, though the data are given in the annex. Details for monthly and annual concentrations for all the sites are found in the annex, table A.1.

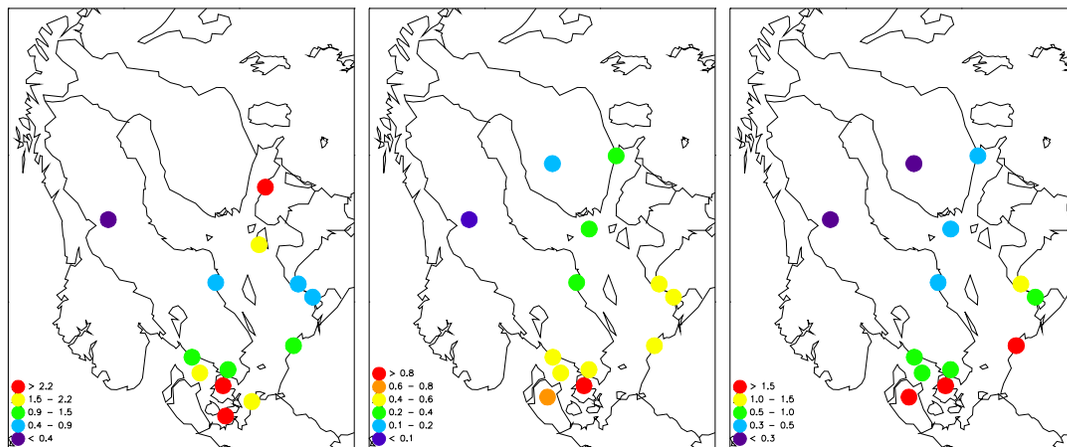


Figure 2.2. Concentrations of left: NO_2 in air, middle: total nitrate ($\text{HNO}_3+\text{NO}_3^-$) and right: total reduced nitrogen ($\text{NH}_3+\text{NH}_4^+$) in 2013 Unit: $\mu\text{g N}/\text{m}^3$.

There is a tendency of increasing concentrations from north to south and towards west. This concentration gradients reflect the varied influence of traffic (ship as well as cars) and agricultural activities. A similar gradient can also be noticed in Figure 2.3-2.5 displaying the station averages of $\text{NH}_3+\text{NH}_4^+$, $\text{HNO}_3+\text{NO}_3^-$ and NO_2 observations across six sub-basins

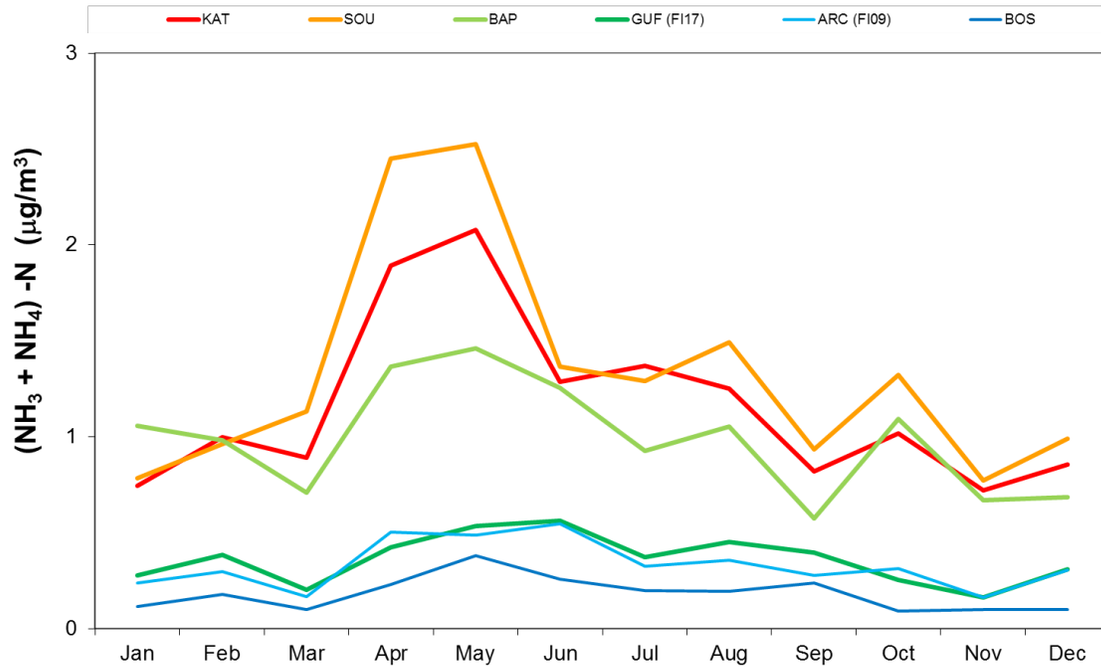


Figure 2.3. Monthly total reduced nitrogen ($\text{NH}_3 + \text{NH}_4$) concentrations in the air in 2013

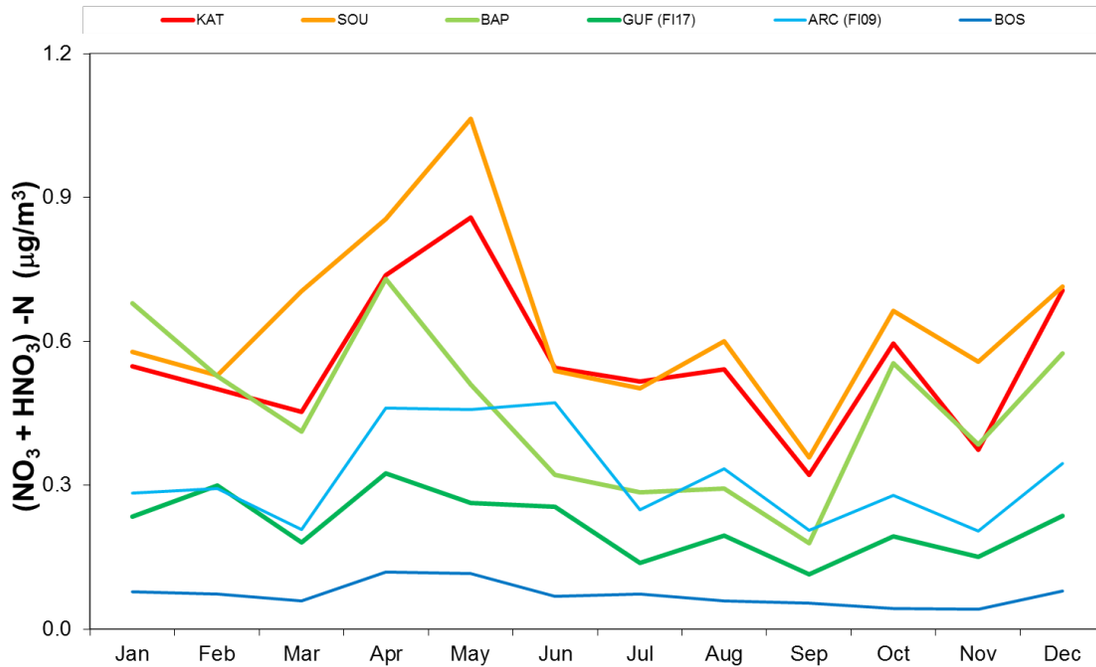


Figure 2.4. Monthly total oxidized nitrate ($\text{HNO}_3 + \text{NO}_3$)-N concentrations in the air in 2013.

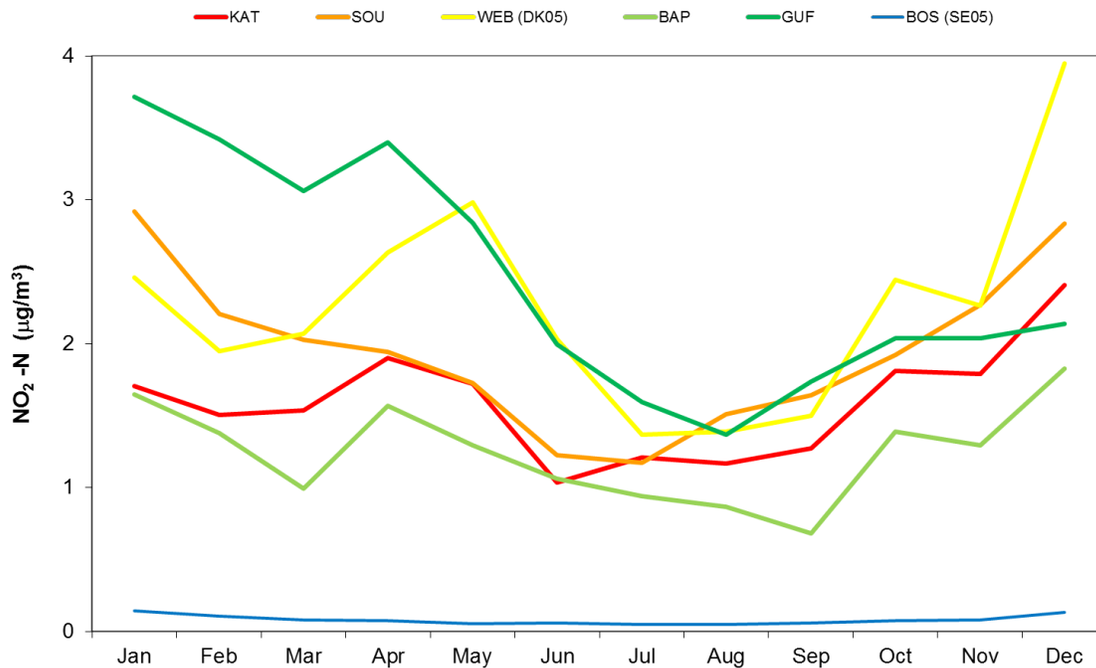


Figure 2.5. Monthly NO₂ concentrations in the air in 2013

Observations of the total reduced nitrogen (NH₃+NH₄⁺), show a seasonal pattern similar for most the sub-basins with highest concentrations during April when the fertilizing is most important. Agricultural activities (natural fertilizer) are the main source for NH₃+NH₄⁺. But also high concentrations are seen in late autumn, maybe due to some fall fertilization, but more likely due to longer residence time of NH₄NO₃ in the atmosphere (see below).

Total nitrate (HNO₃+NO₃⁻) concentration show elevated levels in the spring late autumn; and generally somewhat higher concentrations in winter than summer. NO₂ is reacting photo-chemically and the reaction product is total nitrate. This reaction is mostly dominating during spring and summer. However, total nitrate is dominated by particulate nitrate in the cold season, which has a higher residence time in the atmosphere than nitric acid. In the summer, more of total nitrate consists of nitric acid, which is dry deposited very fast. This effect of fast removal of sum nitrate seems to be relatively more important than higher production in summer, causing summer minima. Concentrations of NO₂ also show, not unexpectedly, a temporal pattern with a winter maxima/summer minima. During winter the atmospheric residence time is longer due to low photo-chemically activity and reduced vertical mixing.

2.3 Nitrogen in precipitation

Altogether sixteen stations have delivered data for ammonium and nitrate in precipitation. Stations from seven sub-basins have delivered data for ammonium and nitrate in precipitation. Annual averages of the two nitrogen species are presented in Figure 2.6.

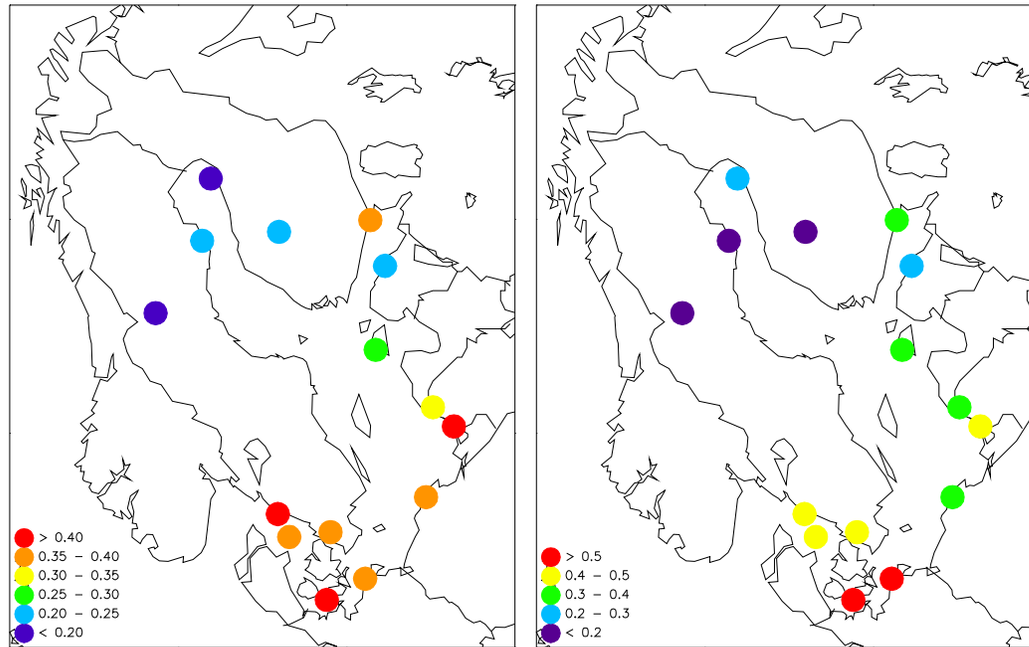


Figure 2.6. Concentrations of left: nitrate (NO_3^-), and right: ammonium (NH_4^+) in precipitation in 2013. Units: mg N/l.

The yearly mean concentrations in precipitation have been calculated from daily or weekly reported values as precipitation-weighted averages. A south-north gradient similar to air can also be seen for nitrogen in precipitation with higher concentrations in the south, and also a west-east gradient is seen. Lowest concentrations for both nitrogen species were seen at SE05 with 0.07 and 0.08 mg N/L for nitrate and ammonium respectively. The highest levels are found at DE09 and DK05 for ammonium (0.66 and 0.62 mg N/L) and DK05 and LT15 for nitrate (0.41 mg N/L). Figure 2.7 displays the station average monthly depositions of oxidized and reduced nitrogen across the regions given.

There is no clear seasonal patterns for the nitrogen wet deposition as for airborne components. The spatial pattern persists, however, with clearly decreasing depositions with progression northwards. For example, the northern regions typically receive half the deposition of reduced nitrogen supplied to southern areas.

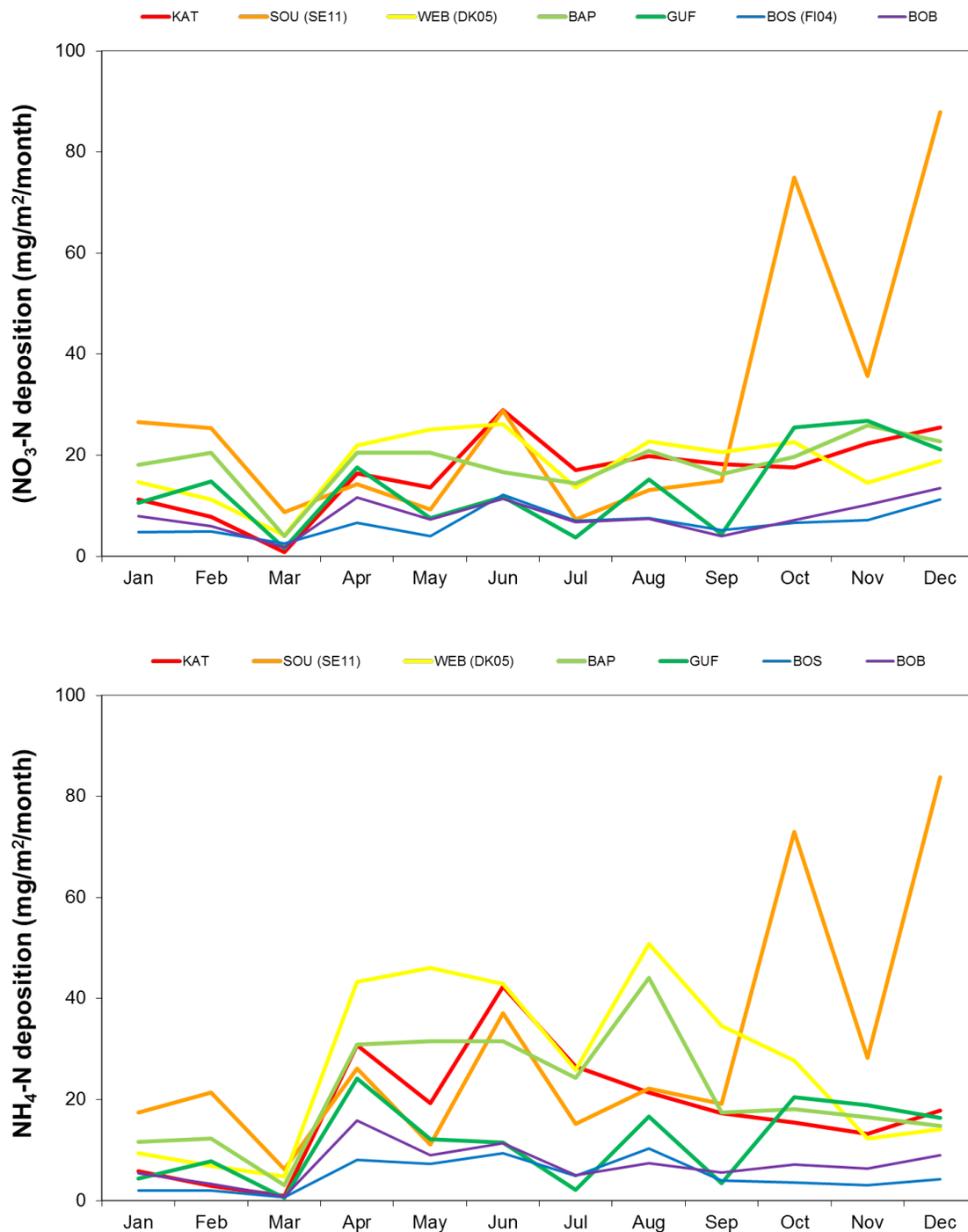


Figure 2.7. Monthly nitrogen depositions in 2013 averaged for the sub-basins. Top: nitrate (NO_3^-), and bottom: reduced nitrogen (NH_4^+).

2.4 Heavy metals in the air

Altogether twelve stations have delivered heavy metal data. Only four sites have reported data for elemental Hg in air, and three of these sites were Swedish. Annual averages of Cd and Pb are presented in Figure 2.8. The lowest concentrations of Cd is seen at the Swedish site SE05 (0.01 ng/m³) while the highest concentration was seen at LV10 with 0.1 ng/m³. For lead, the highest concentrations were also observed at DE09 with 2.8 ng/m³; and the lowest level was at SE05 with 0.3 ng/m³. For elemental mercury the concentrations ranged from 1.37 (SE05) to 1.65 (DE09).

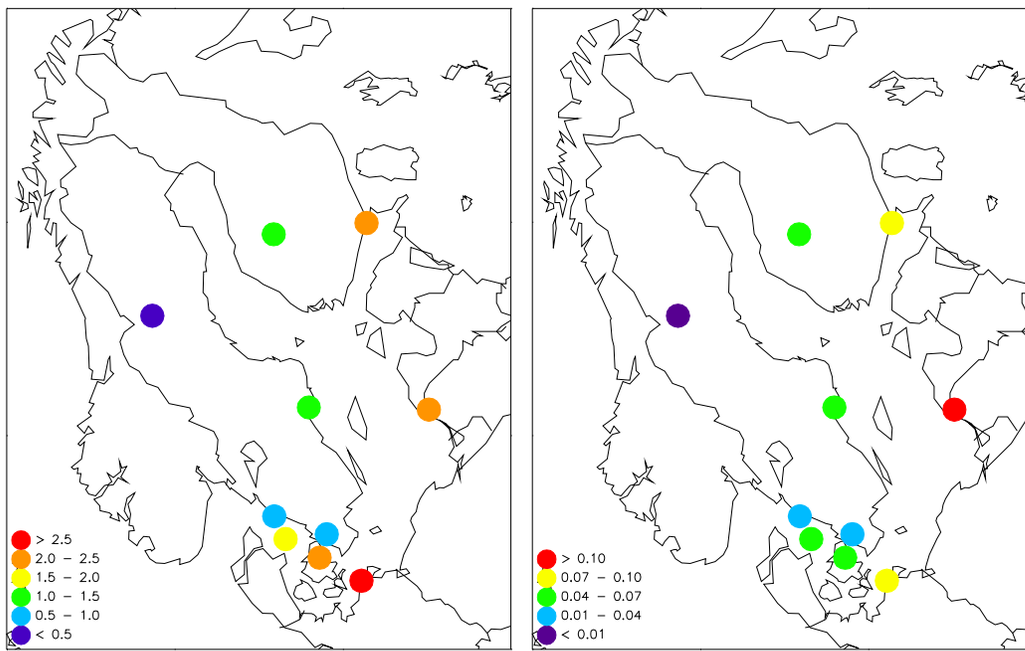


Figure 2.8. Concentrations of left: lead (Pb) and right: cadmium (Cd) in aerosol in air in 2013. Units: ng/m³.

There are insufficient stations to reasonably represent regional patterns; hence, the station data itself is presented here for some of the sites (Fig. 2.9). From this, most of the sites show a winter maximum for Cd and Pb. This is probably due longer atmospheric residence time in winter and reduced vertical mixing.

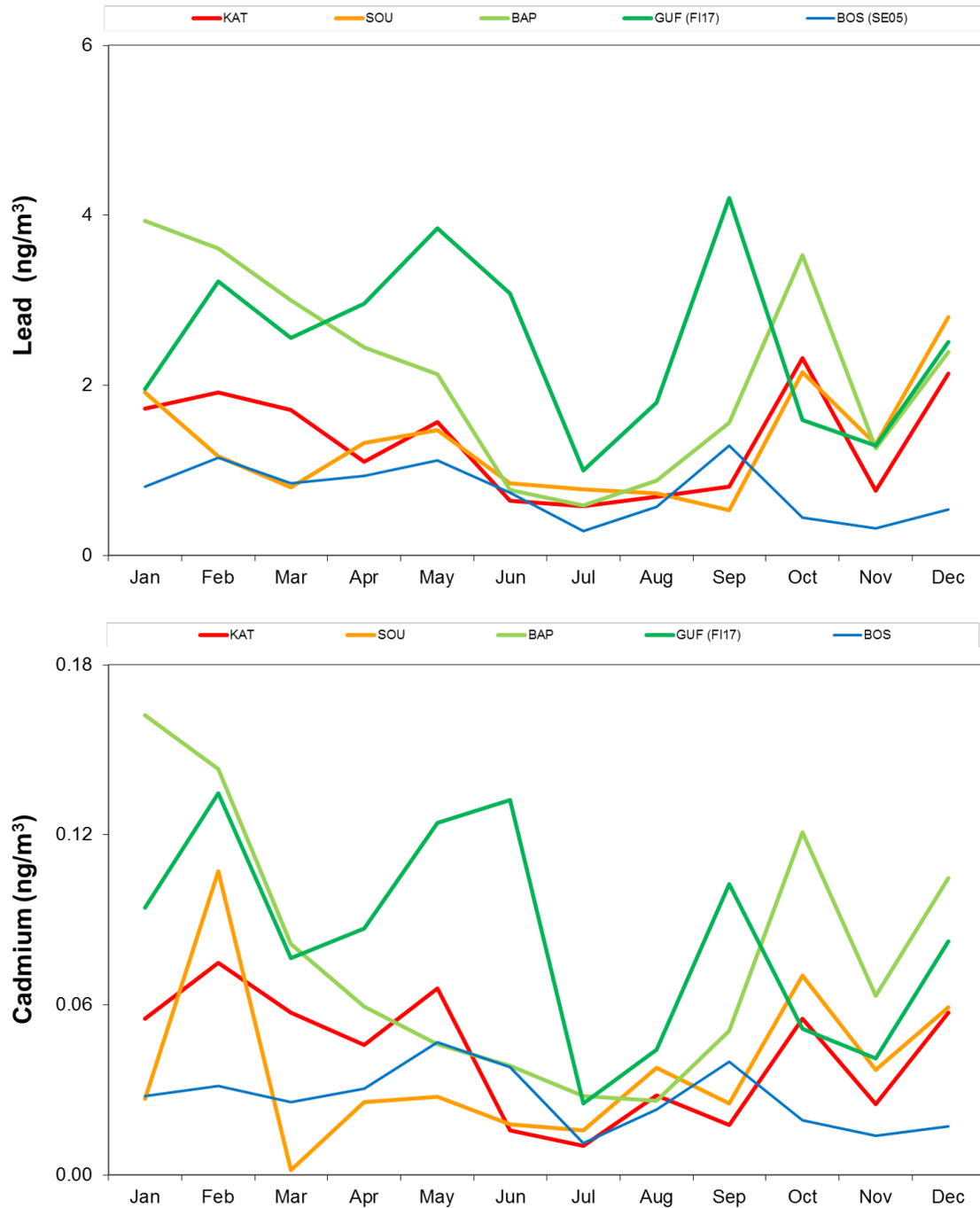


Figure 2.9. Monthly concentrations in air in 2013 averaged for the sub-basins: Top: cadmium, bottom: lead.

Hg concentrations at the four sites has no clear seasonal pattern, though a tendency for a weak winter maxima, Figure 2.10.

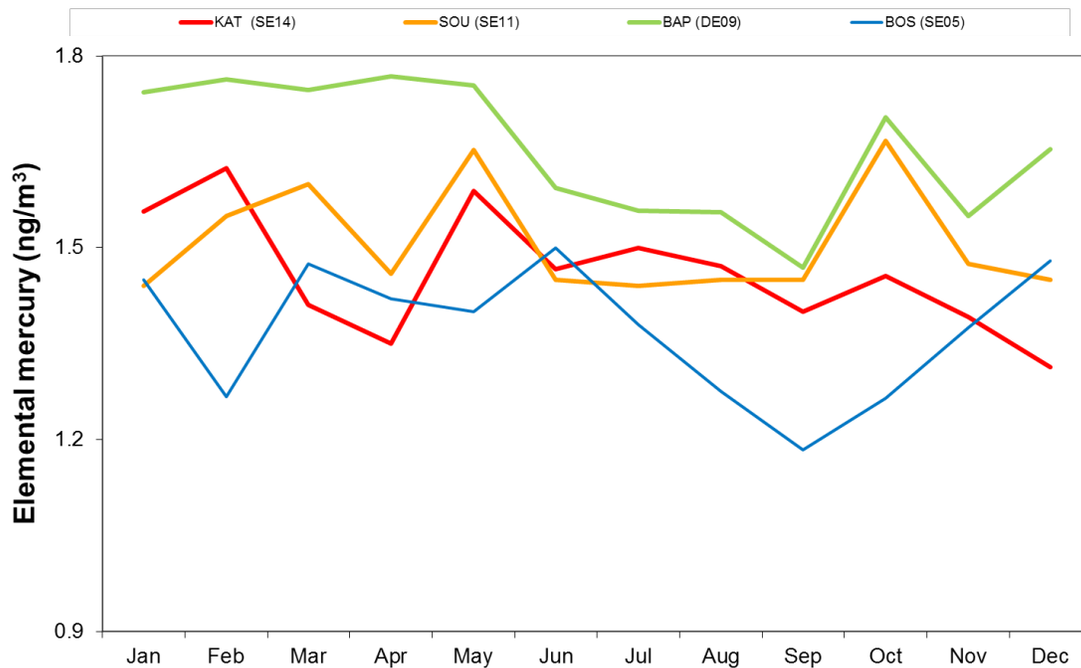


Figure 2.10. Monthly concentrations of Hg in air in 2013 for the four sites representing different the sub-basins.

2.5 Heavy metals in precipitation

Twelve stations have delivered data for Cd and Pb in precipitation, and six have delivered data for Hg in precipitation. Annual averages of Cd and Pb are presented in Figure 2.11. The yearly mean concentrations in precipitation have been calculated from weekly or monthly reported values as precipitation-weighted averages. The lowest concentrations were seen at the Estonian site, but here there are problems with the detection limit especially for Cd. Besides EE09, the lowest concentrations for Cd and Pb in precipitation were reported for the Finnish site FI53 with 0.02 and 0.4 $\mu\text{g/l}$, respectively. For lead, similar low concentrations were seen at SE14, and for cadmium at DK08. The highest concentration of both Cd and Pb were measured at DK05 with 0.10 $\mu\text{g/l}$ and 2.0 $\mu\text{g/l}$, respectively. SE05 observed unusual high lead concentrations in 2013 (1.5 $\mu\text{g/l}$). This is mainly due to very high concentration in December. The reason for this is unknown.

For mercury in precipitation, the highest levels are seen at SE11 and SE14 with 8 ng/L and the lowest at LV10 with 4.5 ng/L.

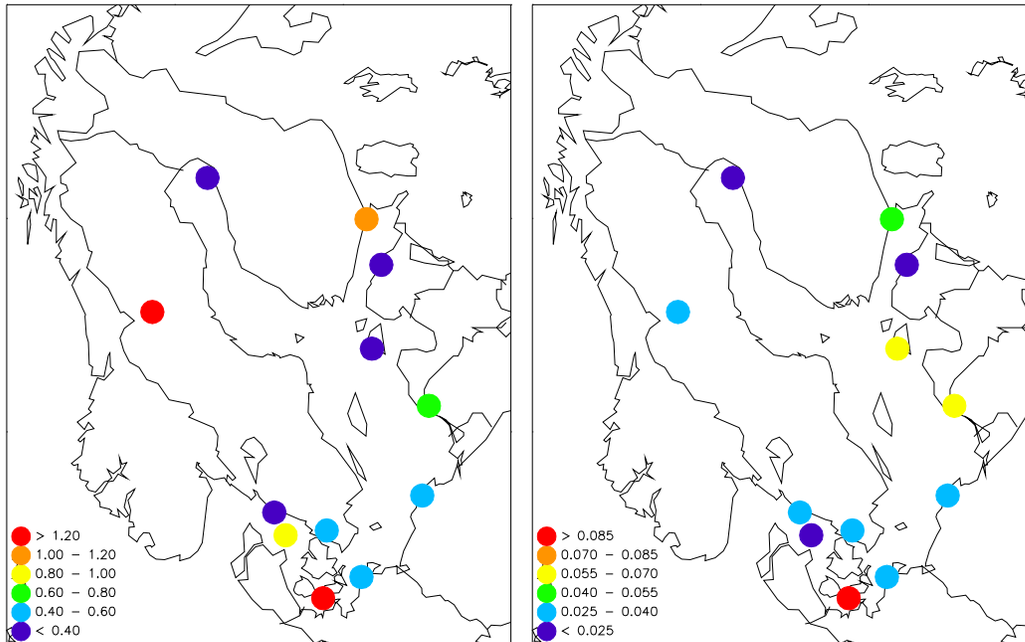


Figure 2.11. Concentrations of left: lead (Pb), right: cadmium (Cd) in precipitation in 2012. Units: µg/l.

2.6 Conclusions for Chapter 2

- Measurement data was reported from eighteen HELCOM stations in 2013, but few sites have a complete measurements program with measurements in both air and precipitation.
- There is a general tendency of decreasing concentrations from south to north for all relevant species; and for many species an east west gradient.
- Many of the components measured in air show a winter maxima due to longer atmospheric residence time.
- The seasonal patterns in precipitation are not as strong as for airborne components. This is due to the presence of the precipitation effect. Though the highest deposition of reduced nitrogen is seen in summer due to enhanced agricultural activity