

Global distribution of atmospheric phosphorus sources, concentrations and deposition rates, and anthropogenic impacts

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[1] A worldwide compilation of atmospheric total phosphorus (TP) and phosphate (PO₄) concentration and deposition flux observations are combined with transport model simulations to derive the global distribution of concentrations and deposition fluxes of TP and PO₄. Our results suggest that mineral aerosols are the dominant source of TP on a global scale (82%), with primary biogenic particles (12%) and combustion sources (5%) important in nondusty regions. Globally averaged anthropogenic inputs are estimated to be ~5 and 15% for TP and PO₄, respectively, and may contribute as much as 50% to the deposition over the oligotrophic ocean where productivity may be phosphorus-limited. There is a net loss of TP from many (but not all) land ecosystems and a net gain of TP by the oceans (560 Gg P a⁻¹). More measurements of atmospheric TP and PO₄ will assist in reducing uncertainties in our understanding of the role that atmospheric phosphorus may play in global biogeochemistry.

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1. Introduction

[2] Phosphorus is an important nutrient for terrestrial and ocean ecosystems, rivaling carbon and nitrogen in importance [e.g., *Schlesinger*, 1997]. Atmospheric total phosphorus (TP) transport is fundamentally different than carbon and nitrogen in that it does not have a stable gaseous phase in the Earth's atmosphere. Thus, TP is mainly restricted to aerosols [e.g., *Graham and Duce*, 1979]. As a result, perturbations to atmospheric TP have been much less than either atmospheric carbon (which is likely to double in the next few decades) or fixed nitrogen (which has already doubled) [e.g., *Schlesinger*, 1997]. Nonetheless, atmospheric TP is being altered. Recent studies have highlighted the role of combustion, primary biogenic and biomass burning

aerosols as significant sources of TP [e.g., *Avila et al.*, 1998; *Echalar et al.*, 1995; *Ferek et al.*, 1998; *Graham et al.*, 2003; *Mahowald et al.*, 2005a; *Baker et al.*, 2006c]. These increases may thus strongly impact terrestrial and oceanic biogeochemistry. For instance, there is evidence that some tropical forests and savanna ecosystems are phosphorus-limited because of the age of their soil [e.g., *Vitousek*, 1984; *Okin et al.*, 2008], and these ecosystems may represent a significant fraction of global net primary productivity [*Field et al.*, 1996]. Ocean ecosystems are commonly believed to be phosphorus-limited on longer, geologic timescales based on residence time arguments [*Falkowski et al.*, 1998], as well as on shorter timescales in some regions [e.g., *Wu et al.*, 2000; *Mills et al.*, 2004; *Krishnamurthy et al.*, 2007]. It should also be noted that as

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humans continue to preferentially increase nitrogen deposition [e.g., Galloway *et al.*, 2003], ecosystems may shift from nitrogen limitation to limitation by other nutrients (e.g., phosphorus) or trace elements.

[3] While model based deposition maps of important nutrients, such as nitrogen [Dentener *et al.*, 2006] or iron [Mahowald *et al.*, 2005b] are available, there has been no similar synthesis of global atmospheric TP since the groundbreaking work of Graham and Duce [1979]. Therefore, a major purpose of this study is to develop a preliminary model that adequately depicts atmospheric TP and bioavailable phosphorus (phosphate or PO_4) concentrations and deposition worldwide. Expanding upon the Amazon phosphorus study of Mahowald *et al.* [2005a], we explicitly model several additional sources of TP, such as biofuels, fossil fuels and volcanic activity. The goals of this paper are to (1) compile available measurements on the global distribution of atmospheric TP and PO_4 , (2) identify the sources of TP and PO_4 , (3) use a preliminary model of TP and PO_4 to provide global maps of TP and PO_4 deposition, (4) understand how humans are perturbing the deposition of atmospheric TP, and (5) highlight gaps in our understanding and prioritize future research.

2. Methodology

2.1. Observations

2.1.1. Description of Data

[4] Most atmospheric TP is in the form of aerosols [Graham and Duce, 1979]. Yet TP is rarely measured in aerosols on a routine basis. Concentrations of TP in the atmosphere are much lower relative to other macronutrients [e.g., Schlesinger, 1997], ranging from only 2–200 ng m^{-3} , thus making detection difficult. The available data fall broadly into two groups of methodologies, the first involving techniques likely to detect TP in the aerosol (e.g., PIXE and XRF) and others that detect a soluble fraction. Measurement of the soluble fraction of TP by ion chromatography (IC) and spectrophotometric methods is sometimes referred to as phosphate (PO_4) or dissolved inorganic phosphorus. IC measures PO_4 and all hydrated forms (e.g., phosphoric acid) [House *et al.*, 1998]. The spectrophotometric method however, requires acidification to pH ~ 1 [Koroleff, 1983]. Thus, this technique measures PO_4 as well as those organic P compounds that are easily hydrolyzed under acidic conditions (such as simple sugars). Although individual studies may show clear relationships between atmospheric TP and inorganic phosphorus [e.g., Chen *et al.*, 2006], there appears to be no significant relationships in the measurements used here [e.g., Baker *et al.*, 2006a]. This may be partly due to issues of sampling, storage and extraction of PO_4 [House *et al.*, 1998].

[5] In this study, we compile high-resolution data sets of TP measurements from across the world (see Table 1 for location, and reference and Table S1 for measurement technique, fine or coarse mode, and other detailed information).¹ Most of these measurements have been published

elsewhere, so methodological details are not reviewed in detail. Readers are directed to the original paper for more comprehensive information. We include 30 long-term stations (Figure 1), and 8 cruises for our detailed analysis. These measurements represent over 7500 individual data points, and most sites include measurements of other elements or chemical species, which provide insight into TP and PO_4 sources.

[6] We also compiled literature values of concentration and deposition measurements (for example, from Graham and Duce [1979]) (Figure 1 and Tables S2a–S2c). Note that some of this data represents one sample at a particular time, and others represent long-term averages. The latitude, longitude, phosphorus value, and citation are shown in detail in the online supplement for phosphorus concentrations, deposition and phosphate concentrations.

[7] Some of the measurements used here segregate aerosols into coarse and fine fractions, usually using 1–2.5 μm in diameter as the division between the aerosol fractions. Most of the measurements, however, only consider aerosols $< 10 \mu\text{m}$ in diameter, while at Erdemli, no maximum cutoff size was used. Significant cycling of phosphorus occurs within ecosystems, and on land this cycling occurs through leaf fall. This may be why studies of phosphorus deposition using moss on land show large spatial variability [e.g., Gombert *et al.*, 2004] (note that this is not included in the comparisons). Therefore we do not show deposition comparisons in the main paper, but rather in the online supplement (Figure S1). The modeled deposition could also be in error, but it is difficult to assess using measurements that includes larger particles. We do have three deposition sites with long-term measurements for which we conduct a detailed comparison between model and observations (see Table 1 for more details): a terrestrial site at Ashiu, Japan, which measured wet plus dry deposition and where they attempted to remove large particles; and two sites with PO_4 concentration measurements in rainwater (Erdemli, Turkey and Tel-Shikmona, Israel).

[8] We make the assumption that measurements of total inorganic phosphorus (TIP) measure most of the TP for our analysis consistent with Chen *et al.* [2006], since the observations at Erdemli, Turkey, Finokalia, Crete, and Tel-Shikmona, Israel and the cruises taken in the Mediterranean are for TIP. This assumption does not appear to be a significant source of uncertainty in our results (section 3), as the data from these stations are consistent with the trends observed with the other stations.

2.1.2. Phosphorus Sources

[9] In order to estimate various sources of TP within the aerosol observations, we compare TP temporal variability at each site with the temporal variability in other measured species that are indicative of specific aerosol source functions. This is conducted using both correlation and regression analysis for each species versus TP as well as by Principal Component Analysis (PCA). Generally, PCA gives similar results to the correlation analysis, but in many cases was difficult to interpret in a consistent manner across many different stations. Furthermore, phosphorus often was isolated in the PCA analysis, providing us no information about sources. We therefore adopt the following methodology.

¹Auxiliary materials are available in the HTML. doi:10.1029/2008GB003240.

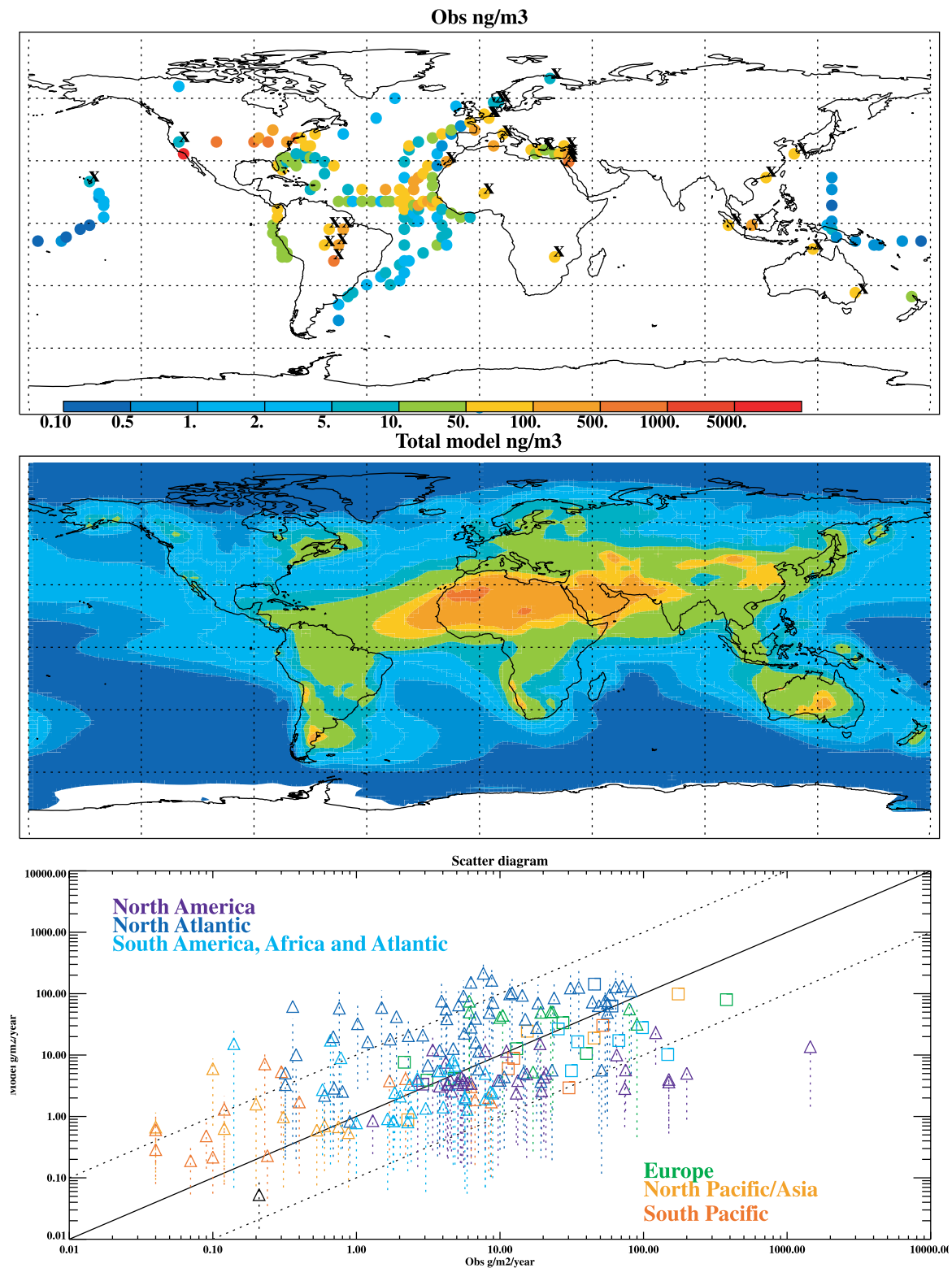


Figure 1. Global distribution of total phosphorus (TP) concentrations based on (top) direct observation (ng m^{-3}) (Tables 1 and S1) and (middle) model. The same color scheme is used for both panels. The locations of the stations with detailed long-term information (Figures 2 and 3) are shown with an “x.” (bottom) A scatterplot between the observations and model estimates. Regions are depicted by color, with squares representing the stations with long-term measurements. In the scatterplot the colored dotted lines are estimates of the error in the model-data comparison when the model represents an annual mean, while the data is taken on 1–2 days (triangles). The methodology is discussed in more detail in section 3.2.

Table 1. Location, Number of Observations and Citation for Each of the Detailed Data Sets Used for Analysis in This Study

	Latitude	Longitude	Observations	Citation
Sevettijarvi, Finland	69.4	28.5	256	<i>Virkkula et al.</i> [1999]
Skreadalen, Norway	58.5	6.4	784	<i>Maenhaut et al.</i> [1997b]
Birkenes, Norway	58.2	8.2	750	<i>Maenhaut et al.</i> [1997b]
Waasmunster, Belgium	51.1	4.1	142	<i>Maenhaut and Cafmeyer</i> [1998]
Gent, Belgium	51.0	3.4	117	<i>Maenhaut and Cafmeyer</i> [1998]
Corsica, France	42.0	9.0	901	<i>Bergametti et al.</i> [1992a]
Lake Tahoe, United States	39.0	240.0	39	A. Paytan and Y. Chen (personal communication, 2007)
Erdemli, Turkey	36.3	34.5	195	<i>Markaki et al.</i> [2003]
Finokalia, Crete	35.2	25.8	115	<i>Markaki et al.</i> [2003]
Cheju, Korea	33.5	126.5	21	<i>Cohen et al.</i> [2004]
Tel Shikmona, Israel	32.8	35.0	72	<i>Herut et al.</i> [1999, 2002, 2005], <i>Carbo et al.</i> [2005], <i>Kocak et al.</i> [2005]
Sde Boker, Israel	30.9	34.8	1511	<i>Maenhaut et al.</i> [1996a] <i>Maenhaut et al.</i> [1997a] <i>Andreae et al.</i> [2002] <i>Derimian et al.</i> [2006]
Eilat, Israel	29.5	34.8	135	<i>Chen et al.</i> [2008]
Canary Islands	29.2	346.5	19	<i>Bergametti et al.</i> [1989]
Hong Kong	22.3	114.2	40	<i>Cohen et al.</i> [2004]
Hawaii, United States	20.0	204.0	139	<i>Benitez-Nelson et al.</i> [2003]
Tillabery, Mali	14.2	1.5	10	<i>Bergametti et al.</i> [1992b]
Pontianak, Kalimantan (Indonesia)	-0.5	109.3	136	<i>Maenhaut et al.</i> [2002b]
Bukit Tinggi, Sumatra (Indonesia)	-0.3	100.3	207	<i>Maenhaut et al.</i> [2002b]
Balbina, Brazil	-2.0	300.0	345	<i>Artaxo et al.</i> [2002]
Santarem, Brazil	-2.0	305.0	290	<i>Artaxo et al.</i> [2002]
Alta Floresta, Brazil	-10.0	303.0	204	<i>Maenhaut et al.</i> [2001] <i>Maenhaut et al.</i> [2002a]
Rondonia, Brazil	-11.0	298.0	87	<i>Fuzzi et al.</i> [2007]
Jabiru, northern Australia	-12.7	132.9	97	<i>Maenhaut et al.</i> [2000b] <i>Vanderzalm et al.</i> [2003]
Rukomechi, Zimbabwe	-16.0	29.5	494	<i>Maenhaut et al.</i> [2000a], <i>Nyanganyura et al.</i> [2007]
Cuiaba, Brazil	-17.3	301.6	178	<i>Maenhaut et al.</i> [1996b] <i>Maenhaut et al.</i> [1999]
Liverpool, Sydney Australia	-33.9	150.9	379	D. Cohen (personal communication, 2007)
<i>Cruises</i>				
PS-Atlantic			18	<i>Losno et al.</i> [1992]
UEA PS-Atlantic			9	<i>Baker et al.</i> [2006b]
UEA JCR-Atlantic			25	<i>Baker et al.</i> [2006a], <i>Baker et al.</i> [2006c]
UEA M55-Atlantic			28	<i>Baker et al.</i> [2006a]
UEA AMT12, AMT13-Atlantic			46	<i>Baker et al.</i> [2006b]
Mantra MP01-09 Atlantic and Pacific			171	<i>Chen</i> [2004]
Aegeo, Mediterranean			8	<i>Carbo et al.</i> [2005]
Meteor, Mediterranean			11	<i>Carbo et al.</i> [2005]
<i>Deposition Data</i>				
Tel Shikmona, Israel	32.8	35.0	29	<i>Herut et al.</i> [1999, 2002], <i>Carbo et al.</i> [2005], <i>Kocak et al.</i> [2005]
Erdemli, Turkey	36.3	34.5	171	<i>Ozsoy</i> [2003]
Ashiu, Japan	35.0	135.0	157	<i>Tsukuda et al.</i> [2005], <i>Tsukuda et al.</i> [2006]

We correlate the TP in the observations at an individual station to the other species (either elements or chemicals) measured. For species which have a statistically significant correlation at the 95% confidence limit, we determine the slope of the relationship between TP and that species. We then multiply the slope by the average species concentration and divide by the average TP amount. This allows us to estimate what fraction of the average observed TP is from a specific element or source [e.g., *Mahowald et al.*, 2005a]. Since there are often several species that represent the same type of aerosol, this will also provide a range of values for the observations. Thus, the range in values presented in the results do not include an error estimate, but rather simply represent a range of % contributions based on different species. Table 2 describes the types of aerosols, and what species we use as indicators for each aerosol type. For marine regions, we use non-sea-salt K, S, and Mg as tracers based on Na.

[10] There is substantial K in biomass burning aerosols as well as primary biogenic particles and desert dust. However, we can separate these sources by size, e.g., fine K is related to biomass burning [e.g., *Echalar et al.*, 1995], while coarse K and Mg is related to primary biogenic particles in non-dust-dominated regions [e.g., *Graham et al.*, 2003]. Inevitably there are uncertainties in these derived sources from this analysis, which will hamper our ability to be definitive about the sources of TP in the atmosphere. Nonetheless, they represent a useful starting point and comparison to model

Table 2. Identification of Different Aerosols

Type of Aerosol	Species Used to Identify
Mineral aerosols	Al, Fe, Ca, Si
Combustion	Black carbon (BC), S, V, SO ₄ , NO ₃
Primary Biogenic Particles	K, Mg
Marine aerosols	Na, Cl

results. For deducing sources of TP, we exclude all measurements which are below the detection limit. For calculating monthly mean values, however, excluding low values will tend to overestimate the mean. Therefore, for monthly mean value calculations, we assume that all values below the detection limit are 50% of the detection limit, if provided.

2.2. Model Description

[11] The chemical transport model used to simulate TP is the Model of Atmospheric Transport and Chemistry (MATCH, version 4.2) [Rasch *et al.*, 1997], driven by the National Center for Environmental Prediction/National Center for Atmospheric Research (NCEP/NCAR) reanalysis data set [Kistler *et al.*, 2001; Mahowald *et al.*, 1997]. The horizontal resolution of the model is T62 ($\sim 1.8 \times 1.8$ degree), and there are 28 vertical levels from the surface to 10 mb. Both dry and wet deposition are included as loss processes for all aerosols [Rasch *et al.*, 2000]. We provide more details about the simulation of dust and other TP containing aerosols in section 2.2.1. Model simulations were conducted for the year 2000, with 1 month of spin-up. Generally, there is very little information about the size distribution of phosphorus, beyond fine or coarse, but differences in size make a large difference to how far aerosols are carried and where they are deposited. We explicitly include information on the assumptions we make regarding the size distribution because of its importance to our final calculation. Because of the large uncertainty in the sources, interannual variability in meteorology is less important for this analysis.

2.2.1. Mineral Aerosol Sources

[12] For the simulation of mineral aerosols or dust, we use the Dust Entrainment and Deposition model [Zender *et al.*, 2003] within MATCH driven by NCEP winds. This combination has been extensively compared to observational data in previous studies [Mahowald *et al.*, 2002; Luo *et al.*, 2003; Hand *et al.*, 2004; Luo *et al.*, 2005]. We choose a globally constant particle size distribution at the sources, but allow four size bins to be transported separately and evolve [Mahowald *et al.*, 2002; Luo *et al.*, 2003; Zender *et al.*, 2003]. The fraction of the mass assumed to go into the different size bins is the following: 0.1 for the class 0.1–1 μm , and 0.3 for the classes 1–2.5, 2.5–5.0, and 5.0–10 μm , where the sizes are the diameters of the particles. Within each bin we assume log normal distributions in aerosol sizes [Zender *et al.*, 2003]. As shown by Alfaro *et al.* [1998], the size distributions of emitted dust vary with soil type and wind speed, and the segregation chosen here is reasonable, although comparisons have suggested it contains too large a fraction of fine particles [e.g., Hand *et al.*, 2004].

[13] Because mineral aerosols are the dominant form of TP, it is important to know the abundance of TP in mineral aerosols. The average crustal amount of TP is 700 ppm [Taylor and McClellan, 1995], but there is significant variability in this value. In the Mediterranean, Carbo *et al.* [2005] report most values between 900 and 1300 ppm. G. Bergametti and collaborators (personal communication, 2007) collected aerosols near source regions and used the slope of the aerosol TP to Al obtain ~ 720 ppm near the

Sahara, 940 ppm in fertilized agricultural areas in Spain, and 1090 ppm in Asia. It is currently not clear how to create a global map of phosphorus in soils and atmospheric TP in desert dust. Okin *et al.* [2004] presents a map of soil phosphorus, which suggests that the Chinese source regions should have lower phosphorus than the African source regions, but this contradicts the available TP in the aerosol measurements above. In order to predict the soil source, we also need to know the phosphorus soil size distribution and what pedogenic and anthropogenic processes, if any, concentrate or reduce the phosphorus in the soils in the size fraction entrained into the atmosphere. Better understanding TP concentrations in dust is beyond the scope of this study, but worthy of further work. Overall, these measurements suggest that TP in dust can vary by at least 30%. For this study, we will assume TP in dust is 720 ppm, a typical value for the most important dust source (North Africa). Sensitivity studies where we increase this fraction uniformly by 30% show that this increase does not qualitatively change the results of this study. This is because where the atmosphere is dusty, dust will dominate by a large fraction, other sources are only important where dust concentrations are low.

2.2.2. Combustion Sources

[14] Spatially explicit emissions of TP from industrial combustion sources were calculated using the Speciated Particulate Emission Wizard (SPEW). This program calculates emissions by combining fuel consumption data given by the *International Energy Agency* [1998a, 1998b], tabulated emission factors, and estimated technology prevalence. Bond *et al.* [2004, 2007] describe this program and the resulting inventories of black and organic carbon. Information added to SPEW for this study included the fraction of aerosols with diameters above 1 μm (supermicron), removal efficiencies for supermicron aerosols by particulate controls, and fractions of TP in submicron and supermicron aerosols.

[15] Percentages of TP found in fine and coarse particulate matter emitted by various sources are given in Table 3. Our goal in this paper is to assess whether combustion sources of TP are potentially important. Thus, we estimate TP emission fluxes toward the high end of reasonable estimates. These emission factors were multiplied by the mass of fine and coarse PM emissions calculated by SPEW. For coal combustion in stokers or residential applications, fine particulate matter is primarily carbonaceous [Flagan and Friedlander, 1978]. We assumed that mineral matter made up all material that was not black carbon (BC), organic carbon, or hydrogen and oxygen associated with the carbon. We then multiplied the mineral matter emission rate by the fractions in Table 3. Coarse particles from coal combustion are usually mineral matter [Flagan and Friedlander, 1978], and we applied the elemental fractions to the entire coarse-mode emission.

[16] For the phosphorus combustion sources, we assume that the submicron TP is transported similarly to BC in the model, but the coarse-mode TP is distributed into the dust size bins in the following fashion for simplicity: 50% in bin 3 and 50% in bin 4. TP is assumed to be emitted in the model

Table 3. Percentages of Phosphorus in Fine and Coarse Particulate Matter Emitted by Various Combustion Sources

Source	Citation	Fine P	Coarse P	Notes
<i>Fossil Fuels</i>				
Coal combustion	<i>Olmez et al.</i> [1988]	0.62%	0.22%	
	<i>Smith et al.</i> [1979]	<0.3–1.3%	-	
	<i>Mamane et al.</i> [1986]	0.69%	0.22%	
	Value used	1.5%	0.2%	
Oil boiler	<i>Olmez et al.</i> [1988]	1.25%	1.95%	
	<i>Mamane et al.</i> [1986]	1%	-	
	<i>Hildemann et al.</i> [1991]	0.17%	-	
	Value used	1%	0.2%	
Gasoline engines	<i>Kleeman et al.</i> [2000]	-	-	a
Diesel engines	<i>Kleeman et al.</i> [2000]	-	-	a
<i>Waste</i>				
Incinerator	<i>Olmez et al.</i> [1988]	0.63%	0.9%	

^aP not reported; we assumed that they would have been if significant values were measured.

with a ratio of P/BC in fine mode of 0.0029 and P/BC in coarse mode of 0.02 [Mahowald *et al.*, 2005a].

[17] Fine-mode biomass burning aerosols are simulated following the methodology following *Rasch et al.* [2001], with the monthly mean biomass burning source from *van der Werf et al.* [2003], as described next. We assume the TP is associated with BC aerosols, which are initially treated as hydrophobic, but convert to hydrophilic on a 1.2 day e-folding timescale (e-folding timescale is one over the first-order rate constant or the time over which the constituent will be reduced to e^{-1} of its original magnitude). Hydrophobic material is assumed to not be removed by precipitation, while hydrophilic material is. To match the observations in the Amazon, coarse-mode biomass burning aerosols are assumed to have a source that is 20% of the fine-mode source [Mahowald *et al.*, 2005a]. Coarse-mode biomass burning aerosols are modeled similar to the coarse-mode desert dust aerosols using two size bins (2.5–5 and 5–10 μm). There is evidence for the entrainment of soil material during fires, which could be responsible for the coarse-mode phosphorus seen from biomass burning.

[18] For biofuels, there are no data regarding emission factors, yet biofuel emissions should be similar to that of biomass burning. For this study, we use the spatial distribution of emissions from the SPEW model for BC from biofuels, and the P/BC ratios from biomass burning, as described above.

2.2.3. Primary Biogenic Aerosols

[19] Primary biogenic aerosols are biotic particles <10 μm that are emitted directly to the atmosphere, such as small bits of plants or spores. There is no clearly defined measurement method for primary biogenic aerosols. Because this aerosol species is poorly understood, these modeling results should therefore be considered tentative. To simulate primary biogenic aerosols we follow *Mahowald et al.* [2005a] and assume that primary biogenic aerosol emissions are spatially proportional to the annually averaged above-ground biomass (in aboveground C/m^2) from *van der Werf et al.* [2003]. We further assume that the source is constant year-round. To obtain the magnitude of the source, we match the values observed in the Amazon [Mahowald *et al.*,

2005a]. This results in an emission factor of 8.9×10^{-18} gP (g aboveground C^*s^{-1})⁻¹ [Mahowald *et al.*, 2005a]. The assumption of year-round biomass release of primary biogenic particles is likely reasonable in the tropics; it may be inappropriate at higher latitudes. In the absence of further information we make the simplest assumption and evaluate it in sections 3.1 and 3.2. For primary biogenic particles, we assume that they are split equally into the two coarse size fractions, since we have no better information at this time.

2.2.4. Sea Salt Aerosols

[20] MATCH does not contain a prognostic sea salt aerosol parameterization that allows us to estimate sea salt aerosol deposition. We use simulations for sea salt aerosols from the Community Atmosphere Model (CAM3) as described and compared to observations by *Mahowald et al.* [2006]. These results are interpolated from the 2.8° CAM3 simulation to 1.8° resolution of the MATCH simulations. The contribution of sea salt to TP is generally thought to be small [Graham and Duce, 1979], thus any errors due to model inconsistencies or resolution should not significantly impact our results. Because PO_4 concentrations in seawater, and hence in sea salt, vary widely, we use the map of surface PO_4 concentrations in seawater from the National Oceanic and Atmospheric Administration's Oceanographic Data Center (NODC) (<http://www.nodc.noaa.gov/OC5>). These values are between 0.05 and 3.5 μM of P. In addition, we assume 0.2 μM of dissolved organic P [Karl and Bjorkman, 2002], although the values in the Mediterranean are likely to be lower. In seawater, average sodium concentrations are 10.781 g Na (kg water)⁻¹ [Quinby-Hunt and Turekian, 1983]. Since sodium is about 30% of sea salt aerosols by weight [Seinfeld and Pandis, 1998], we can obtain a rough spatial distribution of TP in sea salts by assuming that the sea salts deposited have the same TP as the sea salts generated in that grid box. Size distributions are included in the calculation as described by *Mahowald et al.* [2006]. There is some debate on whether there is enrichment of phosphorus in sea salt aerosols [e.g., Graham and Duce, 1982; Hunter, 1997], but in the data contained in this study, we see no evidence for enhanced phosphorus in sea salt aerosols. Regardless, this merits further study.

2.2.5. Volcanic Aerosols

[21] TP-containing compounds in volcanic aerosols have been postulated to stimulate ocean productivity [e.g., Frogner *et al.*, 2001], however actual estimates of the amount of TP emitted from volcanoes are few. Estimates of sulfur emissions from volcanoes are readily available on gridded global data sets [Spiro *et al.*, 1992], so we can use those estimates to place an upper bound on TP emissions from volcanoes. The majority of sulfur in basaltic volcanoes exists as a separate fluid phase. As an approximation, we assume that there is three times as much TP as sulfur in most lava rocks, but the TP is much less likely to volatilize or form small acids than the sulfur dioxide formed in lava flows. In Hawaii, *Sansone et al.* [2002] measured aerosol plume to basalt concentration ratio of 0.002 for TP, which we use here to estimate the amount of TP that is converted to aerosol. A similar value was found for Mt. Etna [Bergametti *et al.*, 1984]. We evaluate these assumptions in section 3.2. While *Duggen et al.* [2007] suggest that during eruptive events

volcanic ash can contribute significant nutrients, we focus here on average or background volcanic activity and not episodic volcanic events. We assume that the volcanic phosphorus is in the fine aerosol mode, as it should be formed at high temperatures.

2.2.6. Model Phosphate

[22] Over ocean regions, only the soluble aerosol fraction may be bioavailable, because of the low residence time of the aerosols in the ocean surface mixed layer. The solubility of TP varies greatly depending on the type of aerosol. In general, the soluble fraction is considered to be PO_4 , and we assume that this is true here as well. Observations suggest between 7 and 100% of phosphorus is soluble [e.g., *Graham and Duce*, 1982; *Chen et al.*, 1985; *Bergametti et al.*, 1992b; *Herut et al.*, 1999, 2002; *Ridame and Guieu*, 2002]. In the Mediterranean, the leachable fraction of inorganic P is 8–25%, 10% for dust storms, and 50–100% for ambient European aerosols [*Bergametti et al.*, 1992b; *Herut et al.*, 1999, 2002; *Ridame and Guieu*, 2002]. On transects of the Atlantic Ocean, average aerosol phosphorus solubility is $\sim 32\%$, with Saharan phosphorus solubility averaging $\sim 10\%$ [*Baker et al.*, 2006a, 2006c]. On the basis of these measurements, we assume that mineral aerosols are 10% soluble P (PO_4), while other sources of TP are assumed to be 50% soluble, with the exception of that from volcanoes, which we assume are 100% soluble phosphorus. We should note that TP is unlikely to be 50% soluble PO_4 , however, nonsoluble P, such as primary biogenic material is likely to be at least partially bioavailable as well [e.g., see *Benitez-Nelson*, 2000; *Karl and Bjorkman*, 2002]. The solubility of aerosol phosphorus is clearly uncertain. Thus, it deserves further study as the soluble aerosol fraction is likely to be the most biogeochemically reactive, especially in aquatic systems.

3. Results and Discussion

3.1. Analysis of Observations

[23] The global distribution of our atmospheric TP measurement compilation show higher concentrations over land or close to the North African dust region, with smaller concentrations in more remote marine environments (Figure 1). There is a large concentration range from 0.002 ng m^{-3} (over ocean regions from *Graham and Duce* [1979]) to 1450 ng m^{-3} (urban airshed in the United States from *Graham and Duce* [1979]; see Table S2a or Figure 1). Notice that these extreme values come from relatively old measurements, and may be inconsistent with newer measurements. The newer studies have spans from 0.12 ng m^{-3} (*Losno et al.* [1992] over South Atlantic ocean) to 378 ng m^{-3} at Sde Boker, Israel [*Maenhaut et al.*, 1996a, 1997a; *Derimian et al.*, 2006]. Measurements on a particular day at Sde Boker, Israel, however, exceed 8000 ng m^{-3} .

[24] The seasonal cycle of TP at the long-term observational sites is shown in Figure 2. These results suggest that there is a strong seasonal cycle in TP concentrations at many locations, having a maximum in the summer and fall in midlatitudes, with a smaller seasonal cycle in the tropics. The source of this seasonality, however, is less clear. Some

sites in the tropical forests of Brazil (Alta Floresta, Cuiaba and Santarem) show strong TP seasonality, associated with biomass burning [e.g., *Mahowald et al.*, 2005a], whereas those near dust sources, show seasonality associated with the dust cycle (not shown). In contrast, primary biogenic particle estimates (based on observed K and Mg seasonal cycles) suggests that this source is low in temporal variability relative to either dust or combustion sources (not shown). Thus, our earlier assumption of no seasonal cycle in the primary biogenic particles is likely reasonable for a first approximation of TP distributions. Also shown in Figure 2 are seasonal cycles in deposition of TP in Japan, which suggest higher deposition during the spring time that are consistent with outbreaks of natural and anthropogenic aerosols from China.

[25] The results of our source apportionment study for all of the observations where measurements were available are shown in Figure 3 for the coarse (Figure 3a), fine (Figure 3b), and total (Figure 3c) aerosol fractions. Although sources were recalculated in a consistent manner for comparison across all the measurements, they tend to remain consistent with the sources identified in the original published papers (see Table 1). For this analysis, we use multiple species, if available, to estimate the fraction of TP from different sources. This provides a range of estimates of the strength of each source, as opposed to true uncertainties (see section 2.1.2 and Table 1). We show in Figure 3 the maximum and minimum estimate for each source derived from the correlation and slope analysis with other elements. If only one line is shown, there is only one estimate. If no line is shown, there is no data. Similar to the results from *Graham et al.* [2003], much of the coarse-mode TP in the Amazon and Indonesia appears to come from primary biogenic particles. This appears to also be true at the midlatitude and high-latitude sites in northern Europe (Finland, Norway, and Belgium) and Asia (Korea) (Sevetti-jarvi, Skreadalen, Birkenes, Waasmunster, Gent and Cheju), but less true at sites close in proximity to anthropogenic (Hong Kong) or dusty areas (Sde Boker, Israel). Most of the stations with only one mode of aerosols collected (Figure 3c) are dominated by dust, and they are predominately in the dust regions close to North Africa. Combustion is an important source for both the fine and coarse fractions of most stations (sometimes exceeding 50%). This includes the coarse fraction in remote areas of the Amazon, Australia, Indonesia and Zimbabwe, and are likely due to biomass burning. In the fine mode, combustion aerosols may even be potentially important in Sde Boker, Israel, where a phosphate processing facility was located nearby. The source apportionment study in the deposition measurements at Ashiu, Japan suggests that 20–35% of the TP is derived from combustion sources, similar to *Tsukuda et al.* [2006]. The UEA ocean data sets [*Baker et al.*, 2006a, 2006c] are consistent with significant anthropogenic sources in both the Northern and Southern hemispheres (Figures 3a and 3b). Unfortunately, there is still substantial ambiguity in our derived sources of TP. As such, these results should be treated with caution. Nonetheless, results are consistent with atmospheric TP coming from nondust sources over large regions of the land and ocean.

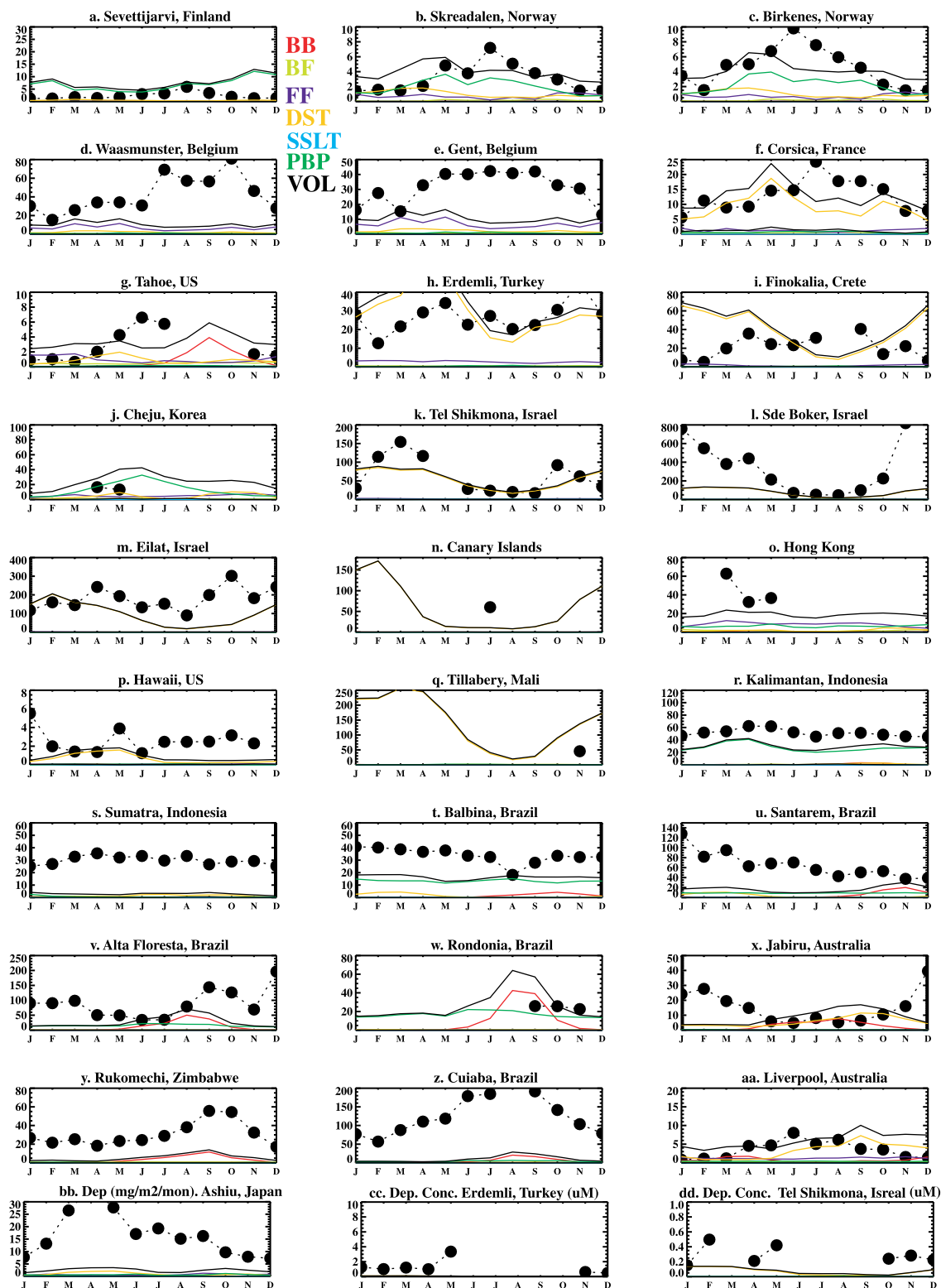


Figure 2. Seasonal cycle of TP at each station (ng m^{-3}). The locations of these stations are shown in Table 1 and Figure 1. The dots and dotted lines represent observations, while the solid lines are estimates from model simulations. The color scheme is as follows: BB, biomass burning, red; BF, biofuels, light green; FF, fossil fuels, purple; dust, yellow; SSLT, sea salts, blue; PBP, primary biogenic particles, bright green; VOL, volcanics, black; and the total model estimate is also in black. Seasonal cycle of TP in deposition ($\text{mg m}^{-2} \text{month}^{-1}$) is shown at Ashiu, Japan (bb), and the concentration of phosphate in deposition (μM) is shown at Erdemli, Turkey (cc) and Tel Shikmona, Israel (dd).

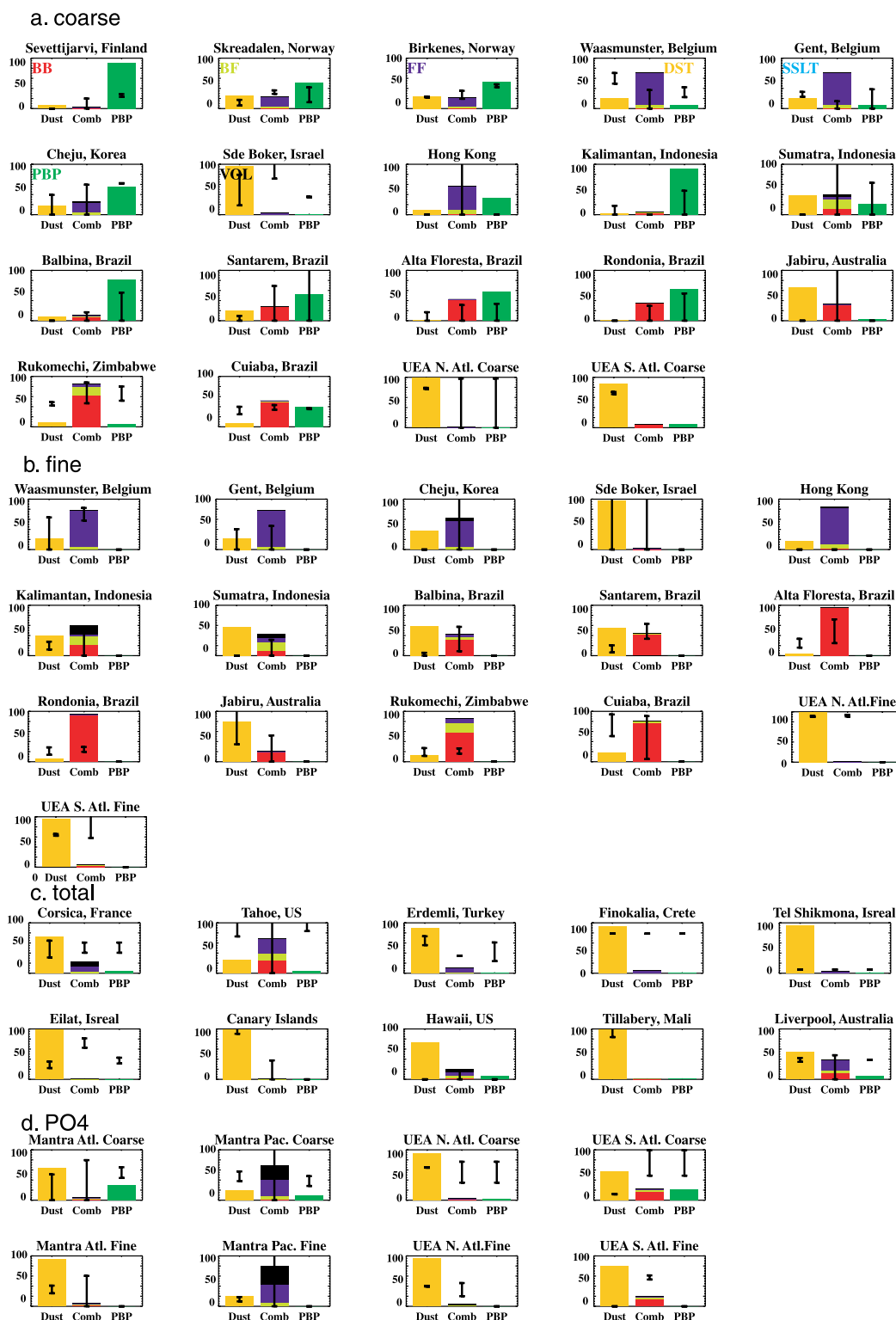


Figure 3. The source apportionment (percentages) for TP in (a) coarse, (b) fine, and (c) total particles at each station and for (d) PO₄. The first column is for mineral aerosol (dust, yellow), the second column is for combustion aerosols (Comb), and the third column is for primary biogenic particles (PBP, green). The filled bars are the model estimates and are represented by the same color scheme as presented in Figure 2. For combustion aerosols, please note that the model estimate is the sum of four different components: biomass burning (BB, red); fossil fuels (FF, purple); biofuels (BF, light green); and volcanoes (VOL, black). The vertical lines on each of the filled bars represent the minimum and maximum estimates derived from the observations (see section 2.1.2 for an explanation).

[26] PO_4 observations are shown in Figure 4. Most of the PO_4 observations are over oceans, and show substantial spatial variability, not necessarily correlated with high-dust regions. This variability could also be due to temporal changes, as the observations for PO_4 tend to be from cruises and represent 1 day aerosol concentrations. The source apportionment analysis (Figure 3d) for the PO_4 observations from the Mantra cruises [Chen, 2004] and UEA data sets [Baker et al., 2006a, 2006b] suggests that PO_4 has large contributions from dust in the coarse and fine mode over most oceans (Figure 3d). However, the measurements are also consistent with a large fraction of the PO_4 coming from combustion sources. As discussed in section 2.1.2, the source apportionment analysis is only a correlative analysis, and so cannot provide conclusive evidence for one source over another. Rather it broadly argues that dust, combustion, and primary biogenic particles may be important sources of PO_4 over the oceans.

3.2. Modeling Results

[27] The aerosols simulated in our model have been extensively and successfully compared against available observations in previous papers [e.g., Luo et al., 2003; Hand et al., 2004; Luo et al., 2005, 2008]. Such validation has not been done previously for the primary biogenic particles and the volcanic sources, so we briefly discuss them here. There are no direct measurements of primary biogenic particles, but our results appear consistent with current assumptions of the distribution of primary biogenic particles as discussed in the online supplement. For volcanic sources, the assumptions in this paper result in a TP source from Hawaiian volcanic activity of $43,000 \text{ kg a}^{-1}$, while Sansone et al. [2002] estimated that only 234 kg a^{-1} of total dissolved TP (which may be smaller than total TP) is emitted, consistent with other estimates [Benitez-Nelson, 2000]. For Mt. Etna, Bergametti et al. [1984] found approximately $76,000 \text{ kg a}^{-1}$ emissions of P (although some days have much higher emissions) while our modeled source predicts $99,000 \text{ kg a}^{-1}$ for that same volcano. This analysis suggests that our model results are a reasonable upper bound on volcanic emissions of TP, and our results (Figure 5) suggest that volcanoes are not important (<10% contribution to annual aerosol concentration) at most locations using these estimates.

[28] This first TP global modeling study appears to capture the major trends in the TP distribution over several orders of magnitude, but differs from observations at many of the sites by more than 1 order of magnitude (Figure 1). The correlation coefficient (calculated using the log of concentrations in order to weight the high and low ends more equally) between the model and observations indicates moderate correlations that are highly statistically significant ($r = 0.64$). The model underestimates atmospheric TP concentrations over North America and Europe, and overestimates it over parts of the remote Pacific region. In Figure 1, the triangles show locations where we are comparing model annual averages to observations taken on a cruise for only 1–2 days (triangles). Because of the high variability in the deposition of aerosols, the annual average will be higher than most of the daily averaged values

collected. In order to account for this in the scatterplot comparisons, we show the range of values representing the median 66% of the daily averaged model concentration as an error bar on the model annual mean (dotted vertical colored lines) at each location.

[29] The model has significant difficulty in capturing the seasonal cycle of TP (Figure 2). For example, the model has a spring maximum with higher concentrations in the summer in rural northern European stations (Skreadalen and Birkenes, Norway), while the observations show the reverse. The model appears to underestimate TP at some of the more human impacted sites by more than a factor of 2 (Waasmunster and Gent, Belgium and Santarem, Brazil). It also overestimates TP at Svettijarvi, Finland and Cheju, Korea (due to too many primary biogenic particles, as discussed in section 3.2), and reverses the seasonality at Svettijarvi, Finland (model peaks in winter, observations in summer). In dusty regions, the model does not capture the seasonal cycle at Finokalia, Crete or Sde Boker, Israel, but does better at Erdemli, Turkey (where total suspended particulate was measured) and Tel Shikmona, Israel. Combined, this suggests that the model does not simulate the seasonal cycle of dust in the Mediterranean. Notice, however, that the model annual average concentration is similar to the observed TP concentrations at these Mediterranean stations (except for Sde Boker, Israel) using our assumed dust composition of 720 ppm TP, although it has been suggested that this region should have a higher P dust value [Carbo et al., 2005]. In contrast, in the tropics (Sumatra, Zimbabwe and Cuiaba, Brazil) the model underestimates TP concentrations, but reproduces the TP seasonal cycle. Since the source apportionment study (see section 3.1) suggests that observations are dominated by combustion, the model TP concentrations likely underestimate the biomass burning source at these stations.

[30] The seasonal cycle in TP deposition appears to be captured at Ashiu, Japan, but again it is unclear how well the model is capturing the seasonal cycle in PO_4 concentrations in the Mediterranean (Figure 2). Overall, the model underestimates the magnitude of TP deposition; consistent with significant deposition derived from local sources that may include particles $> 10 \mu\text{m}$, which are not included in our model. Please note, most of the sources included in the model do not have a seasonal cycle (primary biogenic particles, fossil fuels, biofuels), and inclusion would likely improve simulations in future studies.

[31] Figure 3 also shows the model estimates of the source apportionments (colored bars). The model appears to be consistent with the broad range of estimates from observations at many stations for the coarse mode. The model correctly predicts that dust dominates the TP cycle at stations where this has been observed (Tillabery, Mali Erdemli, Turkey, Finokalia, Crete and Corsica, France), but underestimates the TP dust contribution in urbanized areas (Waasmunster and Gent, Belgium). While these regions are far from the dominant natural sources of dust (North Africa), they may be impacted by large sources of 'fugitive' dust from processes like construction or road dust within industrialized regions. Measurements are ambiguous

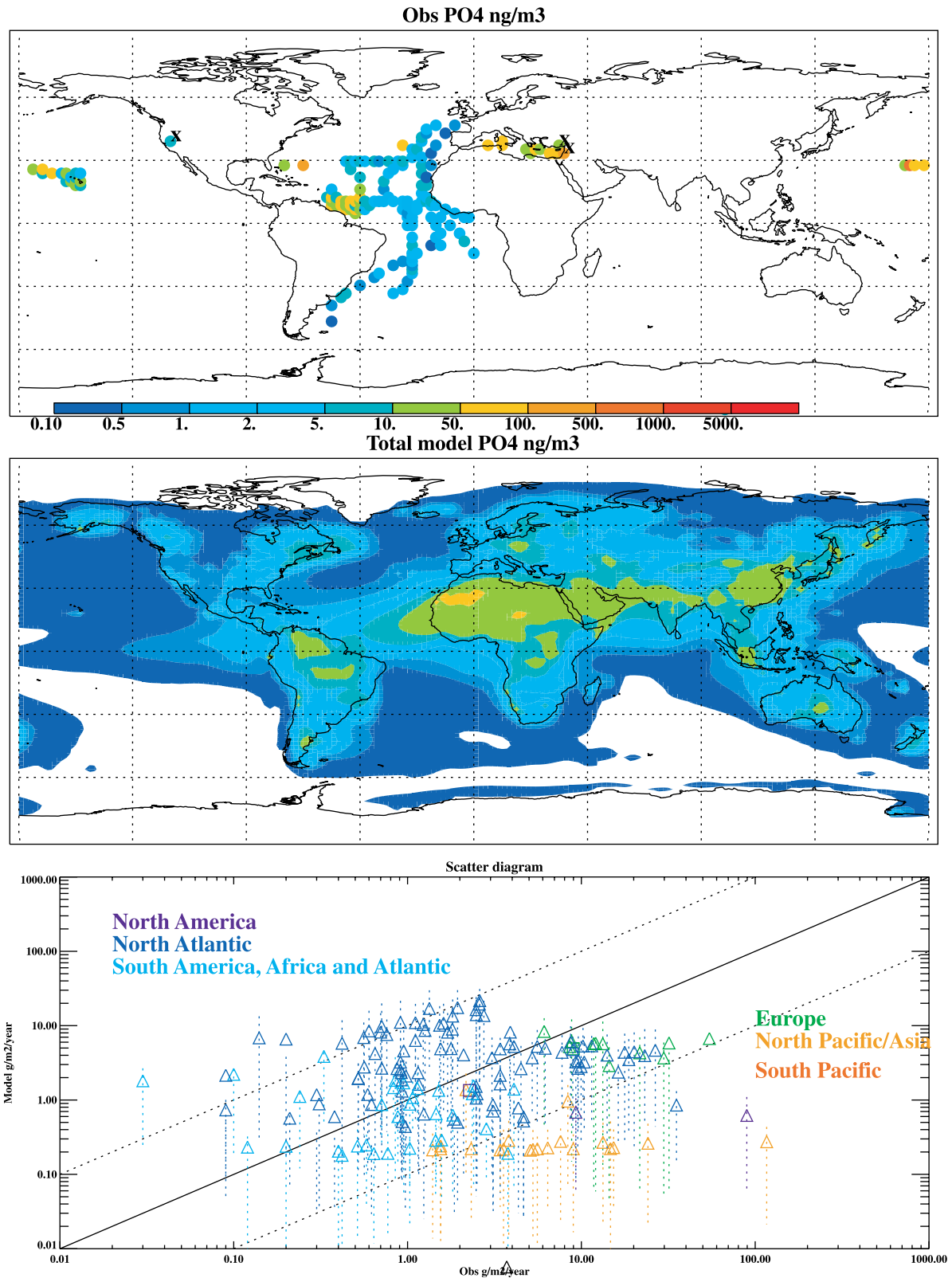


Figure 4. (top) Observational and (middle) modeled PO₄ concentrations (ng m⁻³). The observations are from Table S2a. The scale is the same for both panels. (bottom) A scatterplot shows the comparison between the model and observations. In the scatterplot, the colored dotted lines are estimates of the error in the model-data comparison when the model represents an annual mean, while the data is taken on 1–2 days (triangles). The methodology is discussed in more detail in the text.

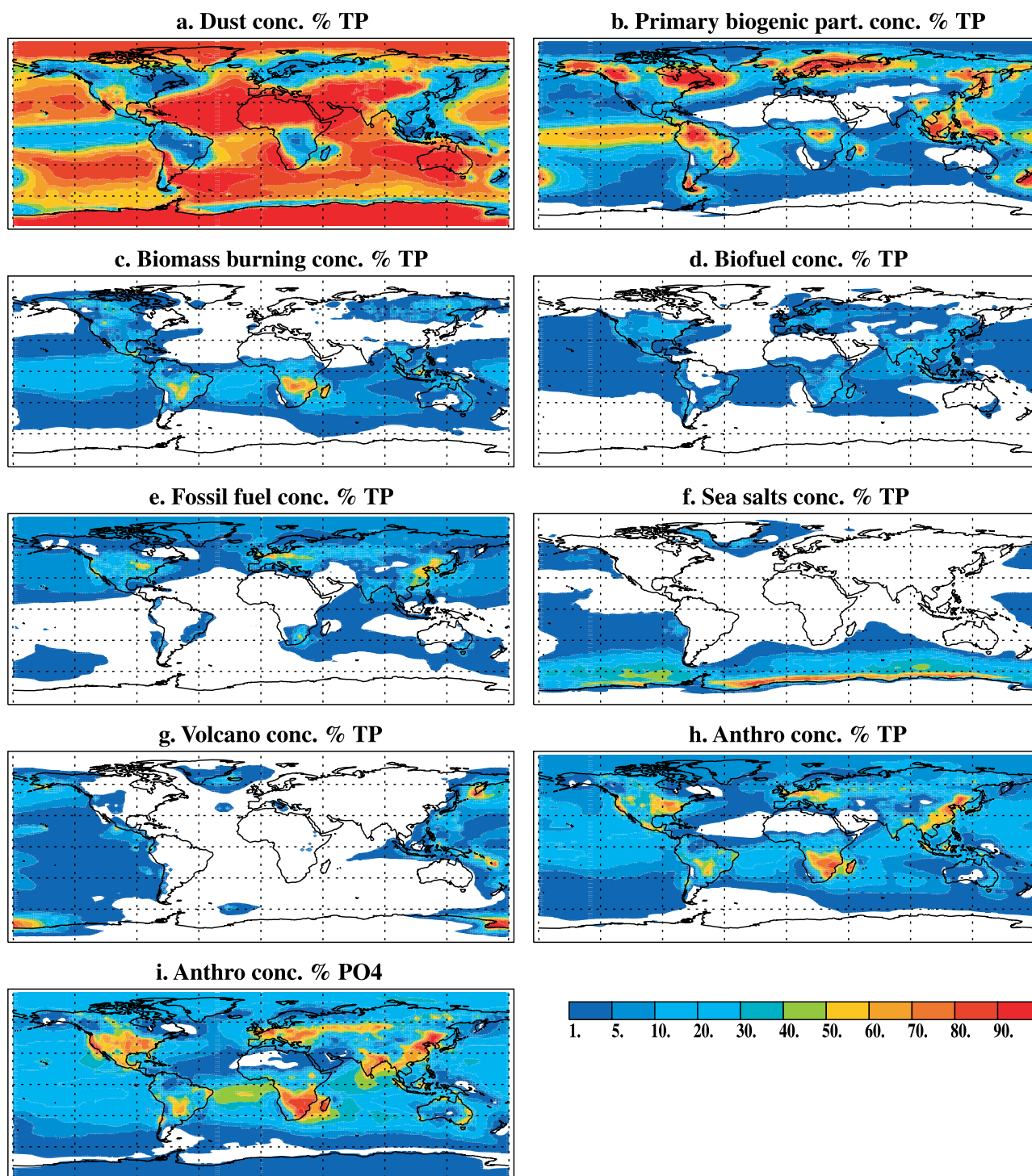


Figure 5. Percentage contribution of different modeled sources to annual average concentration of phosphorus for (a) mineral aerosols (dust), (b) primary biogenic particles, (c) biomass burning, (d) biofuels, (e) fossil fuels, (f) sea salts, and (g) volcanoes. Also shown is the relative importance of anthropogenic disturbance to the (h) atmospheric TP and (i) PO₄ concentration as modeled (assuming all biofuels and fossil fuels and 90% of biomass burning emissions are anthropogenic).

regarding the sources at Tel-Shikmona, Israel, but the model suggests that the site is dominated by dust derived TP.

[32] The model underestimates the fraction of TP from coarse primary biogenic particles at Waasmunster, Belgium

and Erdemli, Turkey and overestimates it at Skreadalen, Finland, Hong Kong, and Balbina and Rondonia, Brazil (Figure 3). In the fine mode, the combustion sources of TP tend to be more important than in the coarse mode in both

the observations and the model. Combustion sources of TP, including biomass burning, appear to be too low in the model in the coarse mode in Gent, and in the fine mode in the Amazon (Balbina, Santarem, Alta Floresta and Rondonia) and Zimbabwe. There is some evidence for higher TP from combustion at several stations where size fractionation is not available (fine + coarse mode). The source apportionment analysis on the deposition measurements from Ashiu, Japan suggest that the model over predicts combustion sources, but model results may still be consistent with natural and locally produced large particles (in sizes $> 10 \mu\text{m}$) contributing to the TP budget.

[33] Because of data limitations, we know very little about the sources of TP in Hawaii, although the literature suggests a combination of dust from Asia [Chadwick *et al.*, 1999] and volcanic activity [Sansone *et al.*, 2002]. Our study suggests TP concentrations (Figure 3) and deposition at Hawaii are dominated by mineral aerosols (85% of deposition), with 5–10% contributions from combustion and primary biogenic particles, and only 2% from volcanic activity.

[34] The model further allows us to estimate the fraction of TP in the coarse mode (Table S3a). In the observations, most stations are dominated by the coarse fraction, and this general feature is simulated by the model. However, at the Asian stations (Cheju, Korea and Hong Kong) and at Cuiaba, Brazil, the fine fraction dominates, perhaps due to specific anthropogenic activities close to these sites. Data collected over the Mediterranean suggests that the median fraction of $\text{P} < 1 \mu\text{m}$ is 55%, roughly consistent with our model values in that region in Table S3b (R. Losno, personal communication, 2008). The model is also within the very broad range of estimates of combustion sources to total aerosols in at the observational sites (fine plus coarse) (Table S3b).

[35] The model predicts the distribution of atmospheric PO_4 using simple assumptions regarding the fraction of TP that is PO_4 (see section 2.2.6, Figure 4). Unfortunately, the model does not simulate the observations well, with a much smaller PO_4 dynamic range in the model relative to the measurements (the correlation coefficient is not statistically significant) (Figure 4). Modeled concentrations tend to be too low, especially in the Pacific Ocean. These discrepancies are likely due in part to our assumptions regarding the PO_4 concentrations from the various sources, but may also be due to inconsistencies in the measurement techniques or the fact that we are comparing modeled annual averages to site specific daily observations. Also shown in Figure 4 is the error bar estimated from the model for comparison of an annual mean to a daily averaged value (similar to Figure 1). Baker *et al.* [2006a] suggest that there is atmospheric processing of P to form PO_4 in the atmosphere, because of an aerosol P solubility gradient away from dust regions in a transect across the North Atlantic. The inclusion of a mix of sources with different solubilities in our model cannot match the gradient in solubility seen by Baker *et al.* [2006a] (Figure S2), suggesting that we should include some processing of P, or stronger gradients in P solubility in the sources.

[36] At several stations we have PO_4 size distributions and estimates of TP solubility (PO_4/P) (Table S3b). The observations suggest that the PO_4 aerosol is more evenly distributed between the fine and coarse fraction than TP, with slightly more in the fine fraction (Table S3a). The model overestimates PO_4 in the coarse fraction which is likely related to why the model underestimates the PO_4 contribution from combustion (we expect more combustion particles in the fine mode). Nonetheless, the observations are consistent with combustions sources accounting for 30–100% of the PO_4 . The model also predicts lower values over the Mediterranean (perhaps due to higher solubilities occurring here than being assumed in our study [e.g., Bergametti *et al.*, 1992b]), and larger values over the Pacific. The observed P solubility in the Mediterranean is between 30 and 70%, while the model only predicts $\sim 14\%$. These comparisons highlight how little we understand or can simulate the observed distributions of PO_4 .

[37] Given the model's ability to capture mean distributions of atmospheric TP (Figure 1), we more closely examine the model results across different regions of the world (Figure 5). While mineral aerosol TP dominates close to dusty regions, other sources of TP dominate elsewhere, consistent with the source apportionment analysis of observations in Figure 3. Sea salts are predicted by the model to contribute less than 5% of the P, except in remote parts of the Southern Ocean, where it can reach as much as 70%. Volcanoes are predicted to contribute less than 5% over most of the globe, but can be substantial in regions of the Southern Ocean, and close to volcanic activity in the Pacific, where it can reach as much as 70% (Figure 5). Assuming that fossil fuel and biofuel combustion are solely anthropogenic and biomass burning is 90% anthropogenic, Figure 5h shows the relative contribution of humans to TP concentrations across the world. Figure 5h suggests that over much of the globe, humans contribute 10–50% of atmospheric TP concentrations, and can exceed 50% in heavily perturbed areas (industrialized and biomass burning regions). Given our assumptions in section 2.2.6, anthropogenic sources of TP are thought to be more soluble. Thus PO_4 concentrations are much more affected by anthropogenic sources (Figure 5i).

[38] Modeled deposition estimates for TP and PO_4 suggest maximum depositions close to the desert regions, and near combustion sources (Figures 6a and 6b). Using the model simulations we can speculate on where atmospheric TP has been the most perturbed. These simulations include many uncertainties, so they should be considered preliminary estimates. The ratio of current to preindustrial total TP and PO_4 deposition (Figures 6c and 6d) suggests that the most impacted areas are experiencing more than 30% increases in TP deposition, with about one third of the oceans experiencing more than 10% anthropogenic TP deposition. The anthropogenic impact on PO_4 is much larger than on TP, although the uncertainties are also larger (see section 3.2).

[39] Globally averaged P sources and deposition derived from the model simulations are shown in Table 4. Similar to previous studies [Graham and Duce, 1979], mineral aerosols dominate the atmospheric TP global cycle by 1 order of

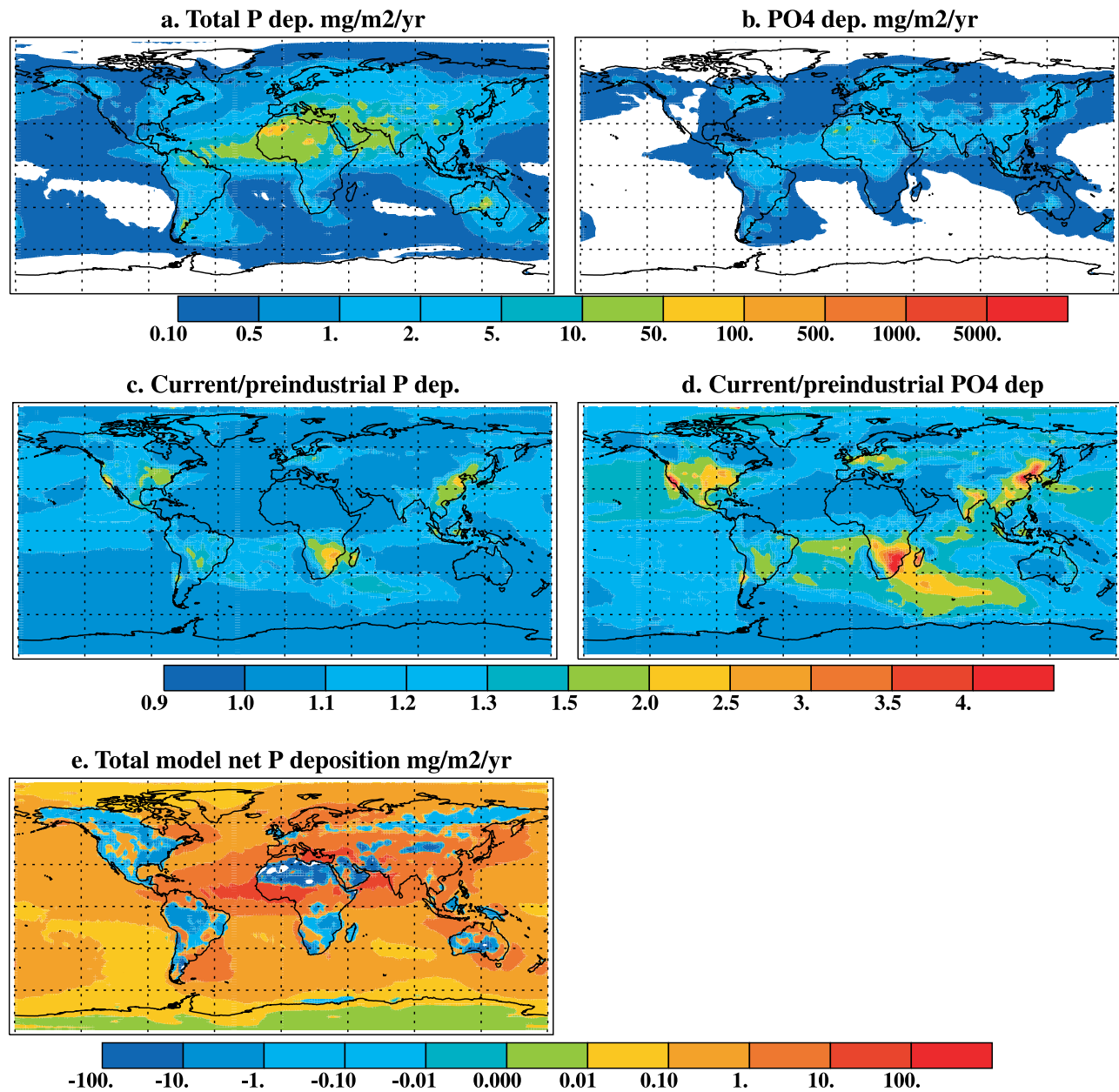


Figure 6. Model estimated deposition fluxes of (a) TP and (b) PO_4 in $\text{mg m}^{-2} \text{a}^{-1}$. Ratio of current to preindustrial (c) TP and (d) PO_4 deposition fluxes, assuming no change in atmospheric dust, and the same assumptions as Figures 5h and 5i. Net flux of TP PO_4 from land and ocean ecosystems ($\text{ng m}^{-2} \text{a}^{-1}$), where downward is a positive flux. Net flux is calculated as the deposition from the atmosphere, minus the source to the atmosphere from dust, biomass burning, biofuels (assuming that these biofuels were generated within one grid box: 180 km), primary biogenic particles, and sea salts.

magnitude. For PO_4 , the assumptions used here imply that mineral aerosols are only responsible for about 50% of the total atmospheric PO_4 . Globally averaged anthropogenic sources (assuming biomass burning is 90% anthropogenic) are $\sim 5\%$ of total TP and $\sim 15\%$ of PO_4 . We use P in dust of 720 ppm, typical for North African dust transported to the North Atlantic, while observations suggest that there can be up to 30% more TP in dust in other locations. If we include the larger estimates of P in dust, our results suggest a 3.8

and 12.5% contribution of anthropogenic sources to TP and PO_4 , respectively. The uncertainties in other sources of TP and PO_4 are even larger, perhaps an order of magnitude.

[40] Using our model simulations, we can further create a global P mass balance and examine which regions are gaining and losing TP through the atmosphere. The net flux is calculated as the deposition flux minus the source flux from the land surface (excluding fossil fuels and volcanoes). Figure 6e shows that regions with large dust

Table 4. Global Sources of Atmospheric Phosphorus^a

	Total P (Tg P a ⁻¹)	PO ₄ (Tg P a ⁻¹)
Dust	1.150	0.115
Primary biogenic particles	0.164	0.082
Biomass burning	0.025	0.012
Fossil Fuels	0.024	0.012
Biofuels	0.021	0.010
Volcanoes	0.006	0.003
Sea salts	0.0049	0.0049
Total	1.39	0.24
Percent anthropogenic	4.8	14.3

^aLess than 10 μm.

entrainment into the atmosphere (North Africa, Asia, Australia), and forested regions (tropical forests, boreal forests) with both primary biogenic emissions and biomass burning tend to be regions that lose TP to the atmosphere. Generally speaking, many (but not all) land regions are sources of atmospheric TP, and all oceans are sinks of atmospheric TP.

[41] Different oceans receive different amounts of TP and PO₄, as estimated here by the model (Table 5). As expected, most of the TP deposition occurs in the ocean basins just downwind of the desert regions (North Atlantic and North Pacific). The soluble fraction of the modeled P deposited ranges from 10% in the Mediterranean, to 34% in the North Pacific. The anthropogenic P fraction ranges between 0.8 and 16%. For PO₄ the fraction from the anthropogenic sources ranges between 4 and 24%.

4. Summary and Conclusions

[42] A combination of model and observations suggests that the global source of atmospheric phosphorus (<10 μm) is 1.39 Tg P a⁻¹, of which 4.8% is anthropogenic. In some heavily perturbed regions influenced by industrialization or biomass burning, the anthropogenic component as estimated by both models and observations may be as much as 50% of TP deposition. We have ignored anthropogenic perturbations to mineral aerosols, which dominate the TP budget close to desert source areas. Mineral aerosols may be significantly influenced (±50%) by human land use, climate or carbon dioxide fertilization [e.g., Mahowald and Luo, 2003].

[43] We estimate a globally averaged soluble phosphorus or phosphate source of 0.24 Tg P a⁻¹ with 14% being of anthropogenic origin. For ocean regions, the phosphate

derived from anthropogenic activities is likely to be even more important, since manmade emissions of phosphorus tend to be more soluble. Unfortunately, there are large uncertainties in the phosphate budget due to the lack of observations and the poor match of the model to observations.

[44] The observations and model results suggest that mineral aerosols are globally the most important source of atmospheric TP, consistent with most of the literature [e.g., Graham and Duce, 1979]. In nondesert areas, other sources dominate, such as primary biogenic aerosols and combustion sources (fossil fuel, biomass burning, biofuels) (Table 4). Sea salt and volcanic sources of TP are estimated to be small, except over very remote regions, or adjacent to volcanic activity. This is the first global model to include primary biogenic particles. We estimate a source of primary biogenic particles of 1.64 Tg a⁻¹, much smaller than previous estimates [Jaenicke, 2005], but large enough that their contribution to aerosol mass should not be ignored.

[45] There are large gaps in our knowledge of atmospheric P and PO₄. First there is a paucity of observations relative to the other major nutrients, partly due to current analytical detection limits. We have only limited measurements globally, and this especially true over North America, Asia and Africa, and oceanic regions, where phosphorus deposition is likely to be important to biological cycling [e.g., Wu *et al.*, 2000; Mills *et al.*, 2004]. We have even less knowledge of the soluble P, or PO₄. The modeling of P is limited both by the availability of observations of P and PO₄, as well as limited knowledge of the source distribution over both space and time, and a lack of knowledge of the aerosol source size distribution. One potentially important source of P that we were unable to include in our model, because of a lack of information, is P emissions from fertilizer factories. In urban areas, fugitive dust sources (e.g., from construction or road dust) also may be important, but data sets describing global sources of mineral aerosols from fugitive sources are not currently available. Because mineral aerosols dominate the TP budget, the uncertainties in the amount of TP in soils (between 700 and 1030 ppm in natural dust source areas, and 1200 ppm in aerosols from fertilized fields) represent a large source of uncertainty of the downwind deposited TP.

[46] Our analysis suggests that atmospheric transport serves as a mechanism for the loss of P from some ecosystems (deserts, forested regions), and a source of new P to other ecosystems, including all of the open ocean.

Table 5. Estimated TP and PO₄ Deposition to Different Ocean Basins

Ocean Basin	P Deposition (Gg a ⁻¹)	PO ₄ Deposition (Gg P a ⁻¹)	Percent Solubility	Percent Anthro P	Percent Anthro PO ₄
Global	558.2	96.5	17.3	4.8	13.9
North Atlantic	189.8	27.9	14.7	2.2	7.3
South Atlantic	41.5	7.9	19.1	7.7	20.2
North Pacific	63.0	22.0	34.9	15.7	22.5
South Pacific	22.2	6.0	26.8	5.2	9.7
North Indian	115.9	13.5	11.7	2.9	12.3
South Indian	33.0	6.8	20.5	9.7	23.6
Southern (<50S)	12.1	1.7	14.3	2.9	10.2
Arctic	6.9	2.2	32.3	6.0	9.3
Mediterranean	53.9	5.7	10.6	0.8	4.0

As shown before [Mahowald *et al.*, 2005a], there is a net loss of P from the phosphorus-limited Amazon because export to the ocean of P in locally generated primary biogenic and biomass burning aerosols exceeds P inputs to the region from North African dust. The anthropogenic impact of biomass burning in the Amazon is estimated to increase the loss of phosphorus from this phosphorus-limited region by 25% [Mahowald *et al.*, 2005a], although nonburned regions may be fertilized by biomass burning phosphorus. This is also true for many land ecosystems, and would ultimately increase the phosphorus that reaches the oceans. Furthermore, while weathering of phosphorus from soils or transport downriver are likely to be the major source of phosphorus to the oceans, human perturbations to biomass burning regimes may significantly perturb phosphorus availability in terrestrial systems.

[47] We estimate that the oceans are receiving a global average of 558 and 96.5 Gg P a⁻¹ of atmospheric phosphorus and phosphate, respectively, 5 and 15% of which are estimated to be anthropogenic. Assuming that the open ocean is phosphorus-limited, and using a Redfield ratio of 106C:P, we calculate a potential new primary production of 0.5 TgC a⁻¹ just from anthropogenic phosphate alone (compared with the 6.4 PgC a⁻¹ emitted by fossil fuel burning). Thus, the total phosphorus deposition to the oceans (558 Gg P a⁻¹ or $\sim 2 \times 10^{10}$ moles P a⁻¹) could be responsible for 22 TgC a⁻¹ of CO₂ uptake, ignoring ocean recycling of P. In some ecosystems, recycling of phosphorus takes place on timescales of days [e.g., Benitez-Nelson, 2000; Paytan and McLaughlin, 2007], suggesting this P could be much more important for primary productivity in phosphorus-limited regions than these simple calculations suggest. On a globally averaged basis, the total phosphorus exported by rivers to the coastal oceans (11 Tg a⁻¹) [Seitzinger *et al.*, 2005], much larger than the atmospheric inputs. However, the riverine inputs may be trapped in coastal zones, making atmospheric P source more important in the open ocean [e.g., Benitez-Nelson, 2000; Paytan and McLaughlin, 2007].

[48] Humans are perturbing the atmospheric nitrogen and P cycles at differing rates. The well established large increases in nitrogen inputs to ocean regions [Galloway *et al.*, 2003] may tend to move ecosystems toward phosphorus limitation, making human perturbations to atmospheric phosphorus more important to understand in the future.

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