RADIOCARBON – A UNIQUE TRACER OF GLOBAL CARBON CYCLE DYNAMICS

Ingeborg Levin • Vago Hesshaimer
Institut für Umwelphysik, University of Heidelberg, Im Neuenheimer Feld 229, D-69120 Heidelberg, Germany.
Email: Ingeborg.Levin@iup.uni-heidelberg.de.

INTRODUCTION

Climate on Earth strongly depends on the radiative balance of its atmosphere, and thus, on the abundance of the radiatively active greenhouse gases. Largely due to human activities since the Industrial Revolution, the atmospheric burden of many greenhouse gases has increased dramatically. Direct measurements during the last decades and analysis of ancient air trapped in ice from polar regions allow the quantification of the change in these trace gas concentrations in the atmosphere. From a presumably “undisturbed” preindustrial situation several hundred years ago until today, the CO₂ mixing ratio increased by almost 30% (Figure 1a) (Neftel et al. 1985; Conway et al. 1994; Etheridge et al. 1996). In the last decades this increase has been nearly exponential, leading to a global mean CO₂ mixing ratio of almost 370 ppm at the turn of the millennium (Keeling and Whorf 1999).

The atmospheric abundance of CO₂, the main greenhouse gas containing carbon, is strongly controlled by exchange with the organic and inorganic carbon reservoirs. The world oceans are definitively the most important carbon reservoir, with a buffering capacity for atmospheric CO₂ that is largest on time scales of centuries and longer. In contrast, the buffering capacity of the terrestrial biosphere is largest on shorter time scales from decades to centuries. Although equally important today, the role of the terrestrial biosphere as a sink of anthropogenic CO₂ emissions is still poorly understood. Any prediction of future climate strongly relies on an accurate knowledge of the greenhouse gas concentrations in the present-day atmosphere, and of their development in the future. This implies the need to quantitatively understand their natural geophysical and biochemical cycles including the important perturbations by human impact.

In attempting to disentangle the complexity of these cycles, radiocarbon observations have played a crucial role as an experimental tool enlightening the spatial and temporal variability of carbon sources and sinks. Studies of the “undisturbed” natural carbon cycle profit from the radioactive decay of ¹⁴C in using it as a dating tracer, e.g. to determine the turnover time of soil organic matter (SOM) or to study internal mixing rates of the global oceans. Moreover, the anthropogenic disturbance of ¹⁴C through atmospheric bomb tests has served as an invaluable tracer for gaining insight into the global carbon cycle on the decadal time scale.

Global Atmospheric Carbon Dioxide Cycle

The mean residence time of carbon dioxide in the atmosphere is in the order of only 5 yr. Accordingly, 20% of the atmospheric CO₂ inventory of today about 750 GtC (1 GtC = 10⁹ metric tons of Carbon = 10¹⁵ g C) are annually exchanged with the biosphere (ca. 60 GtC per year) and the ocean surface waters (ca. 90 GtC per year) (Schimel et al. 1995). Compared to this large gross exchange of carbon between reservoirs, the total yearly net perturbation fluxes are smaller by more than one order of magnitude: In the 1980s, the input of CO₂ into the atmosphere from burning of fossil fuels and cement production summed up to about 6 GtC yr⁻¹, and, in addition, about 1–2 GtC yr⁻¹ was released in the course of deforestation and land use change. Although the effect of these emissions on the atmospheric CO₂ concentration is clearly observed and well documented, it adds up to a mean increase of only 3 GtC yr⁻¹ in the atmospheric burden during the 1980s (Figure 1a). The remaining fraction, corresponding to about 50% of the fossil fuel emissions, is buffered away about equally by
the world oceans and the biosphere. This partitioning of anthropogenic CO₂ between buffer reservoirs is determined by the dynamics of their internal mixing as well as by the strength of their gross carbon exchange with the atmosphere and leads to a residence time of excess man-made CO₂ in the atmosphere as large as several hundred years. Despite intensive research in the last decades, the uncertainty of the gross exchange rates between carbon reservoirs is still in the order of ±20%, and it remains difficult to univocally quantify the repartition of net uptake between the biosphere and the oceans, and thus, to predict the fate of anthropogenic CO₂ emissions. The situation is further complicated by the fact that the share of excess CO₂ taken up by the biosphere and oceans may clearly change in the near future under possibly changing climatic conditions. This concerns in particular the terrestrial biospheric reservoir, which has only a small carbon inventory (about 2200 GtC) compared with the oceans (39,000 GtC). Moreover, this reservoir consists of living plants and dead SOM, which are highly vulnerable and already today strongly perturbed by human activity.

The terrestrial biosphere shows a large diversity, ranging from high-latitude tundra and boreal forests, to cultivated land and grassland, to equatorial evergreen forests. In the oceans, the repartition of carbon is less heterogeneous but a large temporal and spatial variability of dissolved inorganic carbon (DIC) is still observed. Therefore, reliable uptake rates of excess CO₂ resulting from net changes of the carbon inventory in the oceans and in the biosphere are difficult to obtain by direct long-term observations in these reservoirs. In contrast, the atmosphere, being well-mixed in comparison to the biosphere and the oceans, is presently the best-known of all carbon reservoirs. Substantial information about the carbon cycle has been derived from the interpretation of the observed spatial and temporal variability of atmospheric CO₂ in combination with model simulations of the biosphere and the oceans. Consequently, in the past, net anthropogenic carbon fluxes into the biosphere and oceans have been mainly estimated from the gross exchanges between the atmosphere and these reservoirs, also taking into account their internal dynamics. These models have been constrained by suitable (transient) tracers such as bomb ¹⁴C, tritium, and chlorofluorocarbons in the case of the oceans, and by natural and bomb ¹⁴C in the case of the biosphere. The most prominent examples of the application of ¹⁴C will be discussed in the following sections.

**Global ¹⁴C Suess Effect Derived from Tree-Ring Analyses**

It has been erroneously argued that the observed atmospheric CO₂ increase since the middle of the 19th century may be due to an ongoing natural perturbation of gross fluxes between the atmosphere, biosphere, and oceans. That the increase is in fact a predominantly anthropogenic disturbance, caused by accelerated release of CO₂ from burning of fossil fuels, has been elegantly demonstrated through ¹⁴C analyses of tree rings from the last two centuries (Stuiver and Quay 1981; Suess 1955; Tans et al. 1979). Figure 1c shows high-precision ¹⁴C data mostly from single tree rings of two Douglas-firs grown from 1815 to 1975 at the Olympic Peninsula (48°N, 124°W) about 15 km from the Pacific coast of the United States (Stuiver and Quay 1981). A strong decrease of about Δ¹⁴C = −20‰ from 1890 to 1950 is observed in these tree rings. Only a small fraction (Δ¹⁴C = −3‰) of this decrease is estimated by Stuiver and Quay to be due to natural processes (namely the changing natural ¹⁴C production rate through solar and geomagnetic variations), the remaining decrease being attributed to increasing dilution of ¹⁴C in atmospheric CO₂ by input of ¹⁴C-free fossil fuel CO₂ (the so-called ¹⁴C Suess effect). Taking into account the fossil fuel emission rates for the time span in question (Figure 1b) the Suess effect could be quantitatively reproduced by these authors (and independently by Siegenthaler [1983]) using a carbon reservoir box diffusion model. The overwhelming effect of bomb ¹⁴C on the atmospheric ¹⁴C level from 1950 onwards, which cancelled out the ¹⁴C Suess effect from 1955 onwards (shaded area in Figure 1c), is discussed in the next section.
The $\Delta^{14}C$ depletion in the first half of the 20th century has been shown to be larger in more polluted areas, such as Europe, than at the west coast of the United States (De Jong and Mook 1982). In particular, highly populated areas show regional Suess effects up to $\delta^{14}C = -100\%$, especially during winter (Levin et al. 1989). A discussion of the use of $^{14}C$ for regional studies in the context of verifying emission reductions (as agreed upon in the Kyoto Protocol), will be given below (Regional Suess Effect section).
Direct Long-Term Observations of Bomb $^{14}$CO$_2$ in the Global Carbon Reservoirs

During the atmospheric nuclear weapons tests in the 1950s and early 1960s, large amounts of $^{14}$C, in total $630 \times 10^{26}$ atoms of $^{14}$C or more (Hesshaimer et al. 1994), were produced in the atmosphere. This artificial $^{14}$C input caused a global increase in the $^{14}$C/$^{12}$C ratio of atmospheric CO$_2$, leading to a substantial disequilibrium of $^{14}$C between atmosphere, biosphere, and surface ocean water. In the decade following the testing, many laboratories from around the world conducted large observational programs to document stratospheric and tropospheric $^{14}$CO$_2$ changes as well as the penetration of bomb $^{14}$C in all coupled reservoirs of the atmospheric carbon cycle. These measurements enable us to use $^{14}$C as a unique tracer for carbon transfer processes between the reservoirs in question.

More than 30 years of direct tropospheric $^{14}$CO$_2$ observations from both hemispheres are now available since the nuclear test ban treaty in 1962 (Figure 2a) (Tans 1981; Nydal and Lövseth 1983; Levin et al. 1985, 1992; Manning et al. 1990; Meijer et al. 1995). Also, the stratosphere has been extensively sampled during and after the atmospheric bomb tests. The latter data were originally gathered by Telegadas (1971) who calculated bomb $^{14}$C inventories for eight major sub-compartments of the stratosphere. A reassessment of the same data set was performed recently by Hesshaimer and Levin (2000), providing revised stratospheric inventories suitable for global carbon cycle and transport modeling (Figure 2b). The bomb $^{14}$C decline observed in the troposphere mainly reflects the penetration of the $^{14}$CO$_2$ perturbation into the oceans and the biosphere, which is essentially driven by the internal circulation dynamics within each of these two reservoirs. But also, ongoing fossil fuel CO$_2$ emissions and $^{14}$C sources such as nuclear facilities (Otlet et al. 1992, and references therein), and possibly nuclear underground detonations, contribute to the observed trend.

The oceans have been extensively sampled for $^{14}$CO$_2$ analysis in DIC during several international surveys: the Geochemical Ocean Sections Study (GEOSECS, 1972–1978), the Transient Tracers in the Ocean (TTO, 1980–1982) study, and the South Atlantic Ventilation Experiment (SAVE) are prominent examples in this context. In addition, the time development of $^{14}$C in surface ocean water since preindustrial times, and including the bomb perturbation (Druffel and Suess 1983; Druffel 1995) was derived from corals that accrete annual density bands with $^{14}$C activities equal to those of DIC in the surface water of the oceans (see Figure 2a). These data have been evaluated by Broecker et al. (1985, 1995) to determine the distribution of the natural $^{14}$C component in DIC as well as the total bomb $^{14}$C inventories of the world oceans. Their best figure of the global oceanic bomb $^{14}$C inventory, normalized to 1 January 1974, amounts to 295 ± 30 × $10^{26}$ atoms of $^{14}$C (Broecker et al. 1995) (see Figure 2b).

Air–Sea Gas Exchange Rate and Internal Mixing of the Oceans

The combined analysis of the development of bomb $^{14}$C in the global atmosphere as well as in the surface and deep oceans has given us insight into the strength of CO$_2$ exchange between the atmosphere and the oceans, as well as into the mixing dynamics within the oceanic reservoir itself. Accurate knowledge of these processes is crucial to determine the oceanic uptake of anthropogenic CO$_2$. The transport of excess CO$_2$ from the atmosphere into the interior of the oceans is limited by two kinds of transport resistance: air–sea gas exchange which is controlled by a thin water film at the water surface, and oceanic “mixing” circulation responsible for the exchange between the surface and deeper layers of the oceans. In many areas of the oceans, i.e. in mid-to-low latitudes, the gas exchange at the air–sea interface is fast enough, respectively, surface water renewal through mixing e.g. with the deep ocean is slow enough to ensure surface water to be almost in equilibrium with atmospheric CO$_2$ (the global mean equilibration time of ocean surface water is about 1 yr). However, in regions of the ocean with large deepwater formation rates, and respective fast renewal of the
surface water (i.e. in polar regions), the rate of gas exchange becomes important for net transport of excess CO$_2$ into the deep ocean (Siegenthaler 1983).

Both natural and bomb $^{14}$C have been successfully used in the past to derive the mean atmosphere/ocean gas exchange rate for major ocean basins (Stuiver 1980; Stuiver et al. 1981). The net $^{14}$C influx from the atmosphere to the surface ocean at a given time is proportional to the $\Delta^{14}$C gradient...
at the interface multiplied by the gross CO\textsubscript{2} exchange flux. In the undisturbed natural case, the global mean difference of $\Delta^{14}\text{C}$ between the atmosphere and the surface ocean was about 40‰. At equilibrium, the corresponding net $^{14}\text{C}$ influx was just enough to compensate the $^{14}\text{C}$ decay in the oceans, immediately yielding the CO\textsubscript{2} gas exchange flux for the preindustrial situation (ca. 280 ppm). During the period after 1955, disturbed by nuclear tests, the CO\textsubscript{2} gas exchange rate could be determined in an independent manner from the, in this case, time dependent uptake of bomb $^{14}\text{C}$ in the oceans until the time of GEOSECS (see Stuiver 1980). In this way, Stuiver et al. (1981) derived mean area-weighted CO\textsubscript{2} exchange fluxes for the Pacific and for the Atlantic oceans between 50°N and 50°S for 1955–1974. The bomb $^{14}\text{C}$-derived preindustrial gas exchange rate turned out to be in very good agreement with the mean steady state value. However, the $^{14}\text{C}$-derived gas exchange rates are larger by more than 30% if compared to figures derived from direct measurements in wind tunnels, over lakes and open ocean (Liss and Merlivat 1986; Boutin and Etcheto 1997). Although this discrepancy has not been fully resolved yet (Broecker et al. 1986; Wannikhof 1992), the $^{14}\text{C}$-derived exchange rate between the atmosphere and the ocean, leading to a gross exchange flux of about 90 GtC yr$^{-1}$ for the period 1980–1990 is adopted for all carbon-cycle models.

The dynamic model parameters for the internal mixing of the ocean have also been frequently determined by the distribution of natural $^{14}\text{C}$. Unfortunately, this procedure gives quite different answers for these fluxes than calibration with the bomb-$^{14}\text{C}$ distribution (Oeschger et al. 1975; Broecker et al. 1980; Siegenthaler 1983). The explanation for this discrepancy is that the distribution of natural $^{14}\text{C}$ is more sensitive to mixing processes on time scales of hundreds or thousands of years, whereas the characteristic time scale for the bomb $^{14}\text{C}$ uptake perturbation is only several decades. Siegenthaler and Joos (1992) solved this problem with their High-Latitude Exchange/Interior Diffusion-Advection (HILDA) model. They introduced an eddy diffusivity in the interior of the ocean, which is decreasing with depth. (In the original Oeschger-type box diffusion models, the eddy diffusivity is assumed constant with depth.) With this modification, natural plus bomb $^{14}\text{C}$-calibration yields a reliable description for the whole range of mixing processes in the ocean on all relevant time scales from 10 to 1000 yr. In particular, the calibration now provides a realistic description of the time-dependent uptake rates for anthropogenic excess CO\textsubscript{2} for the time scale in question, namely several hundred years.

**Budgeting Bomb $^{14}\text{C}$ in the Global Carbon Cycle**

Natural and bomb $^{14}\text{C}$ have also been used in several studies to determine carbon turnover times within the terrestrial biosphere, in particular for SOM (e.g. Goudriaan 1992; Trumbore et al. 1993; Perruchoud et al. 1999). However, the heterogeneity of the terrestrial biosphere hampers the determination of globally valid bomb $^{14}\text{C}$ inventories for this reservoir based on observational data. Evidence on the exchange fluxes of bomb $^{14}\text{CO}_{2}$ with the biosphere is, however, also inherent to the atmospheric $^{14}\text{CO}_{2}$ record. It should, therefore, in principle be possible to gain insight into the cycling time of carbon within the biospheric reservoir by budgeting bomb $^{14}\text{C}$ in the global carbon system. This approach was used by Hesshaimer et al. (1994): The total decay-corrected bomb $^{14}\text{C}$ input into the carbon system at a given time must equal the sum of the corresponding bomb $^{14}\text{C}$ inventories of the stratosphere, the troposphere, the oceans, and the terrestrial biosphere. Besides the biospheric bomb $^{14}\text{C}$ inventory, the second unknown in this exercise is, however, the $^{14}\text{C}$ input from atmospheric bomb tests, i.e. the $^{14}\text{C}$ yield per Mt TNT. Theoretically, this $^{14}\text{C}$ yield can be derived from nuclear physics. However, those calculations have uncertainties in the order of nearly a factor of 2 (Bonka 1980; UNSCEAR 1982). The total bomb $^{14}\text{C}$ input into the atmosphere was, therefore, derived by Hesshaimer et al. (1994) on the basis of published bomb detonation strengths (Rath 1988) through adjusting the $^{14}\text{C}$ yield per Mt TNT to the tropospheric and stratospheric observations.
during the time period of the major $^{14}$CO$_2$ rises (i.e. 1950–1963). As long as we trust the atmospheric observations and the bomb detonation strengths the uncertainty of this adjustment is small (in the order of about ±10%) since the total bomb $^{14}$C uptake by the ocean and by the biosphere was not important before 1963.

A simple standard compartment model of the global carbon cycle (similar to the one by Oeschger et al. 1975) was then applied to infer the straightforward bomb $^{14}$C budgeting. In doing so, Hesshaimer et al. (1994) found a serious mismatch in the total bomb $^{14}$C inventory with an apparent $^{14}$C source missing. Possible solutions for this mismatch have been argued to be due to: 1) an over-estimate of the oceanic bomb $^{14}$C inventories, 2) residing bomb $^{14}$C hot spots in the very high stratosphere, or 3) much less bomb $^{14}$C uptake by the terrestrial biosphere than estimated from standard parameterizations of this reservoir (e.g. Goudriaan 1992). Other modeling studies, trying to globally match bomb $^{14}$CO$_2$ levels observed in the carbon cycle reservoirs (Enting et al. 1993; Broecker and Peng 1994; Lassey et al. 1996; Jain et al. 1997) solved inconsistencies in their budgets by generally mistrusting the stratospheric inventories of Telegadas (1971). These inventories have turned out to be largely confirmed by the revised calculations of Hesshaimer and Levin (2000) so that further investigations are necessary to solve the problem of budgeting bomb $^{14}$C in the global carbon cycle. Solving this inconsistency, and in particular, independently confirming the penetration rate of bomb $^{14}$C into the oceans, is crucial for the prediction of future atmospheric CO$_2$ concentrations because the oceanic uptake of excess CO$_2$ is generally normalized with bomb $^{14}$C (see the Air–Sea Gas Exchange section above). As mentioned earlier, it is important to know explicitly which fraction of the anthropogenic emissions enters the biosphere and which fraction enters the oceans, because carbon storage in these two reservoirs behaves very differently. If the anthropogenic CO$_2$ is eventually stored in the deep oceans it is unlikely that it will re-enter the atmosphere within the next centuries. In contrast, CO$_2$ stored in the biosphere may be released back to the atmosphere within a few decades. Moreover, the capability of the terrestrial biosphere for uptake of excess-CO$_2$ is very sensitive to changes through human activities, in supply of nutrients and water, but also due to climatic feedback.

**Global Distribution of Atmospheric $^{14}$CO$_2$ in the 1990s and Beyond**

The most prominent applications of the bomb $^{14}$C perturbation as a transient tracer for carbon cycle studies took place in the 1970s and 1980s when the signal was largest in all reservoirs. After 1990, many of the extensive observational programs phased out as $^{14}$CO$_2$ approached a new quasi-natural equilibrium state. From 1982 onwards the globally decreasing $\Delta^{14}$CO$_2$ trend in fact almost exactly follows an exponential curve with a time constant of 18.70 ± 0.15 yr, thus changing from a decrease of about $\Delta^{14}$CO$_2 = -13\%\ \mathrm{yr}^{-1}$ in 1982 to about $-4\%\ \mathrm{yr}^{-1}$ in 1998 (see Figure 2a). With the last atmospheric bomb test being conducted in 1980, and although much weaker underground tests are still performed, bomb $^{14}$C homogenized in the troposphere and additional input from the stratosphere became negligible. In the 1990s and today $^{14}$CO$_2$ in the troposphere, therefore, again reflects the distribution of natural and anthropogenic CO$_2$ sources and sinks. This is nicely illustrated by our modern high-precision measurements at seven globally distributed background stations located between $82^\circ$N (Alert, Nunavut, Canada) and $71^\circ$S (Neumayer, Antarctica) (Figure 3b). Although the north-south gradients are in the order of only a few per mil, a distinct structure is observed in the yearly mean $^{14}$CO$_2$ profile.

Figure 3b shows a relative $\Delta^{14}$CO$_2$ minimum in mid-northern latitudes. It is caused by the maximum effect of fossil fuel CO$_2$ emissions in this latitude belt. These anthropogenic emissions are also responsible for the north-south gradient of the CO$_2$ mixing ratio shown in Figure 3a, with about 1% higher concentrations in the northern than in the southern hemisphere. A second relative $\Delta^{14}$CO$_2$ minimum is observed in mid-to-high latitudes of the southern hemisphere and is associated with gas
exchange with the circum Antarctic ocean surface: upwelling of intermediate water around Antarctica, which is depleted in $^{14}$C (Figure 3c), causes a strong disequilibrium between atmospheric $^{14}$CO$_2$ and $^{14}$C in DIC of the surface ocean water. The respective CO$_2$ flux from the ocean to the atmosphere is, therefore, depleted in $^{14}$C. Driven by the large gross exchange rate of CO$_2$ at these latitudes due to high wind speeds, a strong $^{14}$C draw down is observed in the atmosphere at Macquarie Island and Neumayer. The two $\Delta^{14}$C depleting effects in mid to high latitudes of both hemispheres cause a relative $\Delta^{14}$CO$_2$ maximum to appear in tropical and
subtropical regions (Figure 3b). This maximum is probably further enhanced by a net flux of bomb
$^{14}$C from the terrestrial biosphere to the atmosphere; two-thirds of the yearly terrestrial biospheric
$\text{CO}_2$ exchange flux is located between $30^\circ$N and $30^\circ$S, the mean carbon turnover time of these
ecosystems (i.e. tropical rain forests) being in the order of 30 yr. Bomb $^{14}$C that was taken up in the
early 1960s is now re-emitted to the atmosphere in the 1990s (see Figure 2b). This $^{14}$C enriched
biospheric $\text{CO}_2$ probably contributes to the $\Delta^{14}$CO$_2$ maximum observed at our tropical and
subtropical sites. Although the $\Delta^{14}$C signals in recent global background atmospheric $\text{CO}_2$ are rather
small, they provide independent information on global $\text{CO}_2$ sources and sinks. This information has
not yet been retrieved from the data sets by carbon-cycle modeling exercises.

Figure 4  (a) Monthly mean $\Delta^{14}$C observations in the upper Rhine valley at the polluted site Heidelberg together with
the maritime background level, derived from observations at the baseline station Izaña, Tenerife (28°N) (Levin et
al. 1989 and unpublished Heidelberg data). The maritime background $^{14}$C level before 1988 is calculated from
observations at Vermunt, Austria (47°N) (Levin et al. 1985), which were corrected by $\Delta^{14}$C = +2‰, therewith
accounting for a general European fossil fuel offset. (b) Mean fossil fuel contribution at Heidelberg calculated with
a two-component mixing model from the $\Delta^{14}$C difference between Heidelberg and Izaña, the latter representing the
maritime background level.

Regional $^{14}$C Suess Effect

As has been discussed above, the most important man-made perturbation of the atmospheric $\text{CO}_2$
cycle is the burning of fossil fuels, with about 95% of the $\text{CO}_2$ being released in the northern hemi-
sphere. The fossil emissions today contribute about 50% to the decreasing $\Delta^{14}$CO$_2$ trend observed in
the global atmosphere (Figure 2a). In polluted areas, e.g. over the European continent, an additional
regional surplus of $\text{CO}_2$ from the burning of fossil fuels is clearly detectable as $\Delta^{14}$C depletion relative
to maritime levels. Levin et al. (1995) used $\Delta^{14}$C observations to estimate the fossil fuel contribu-
tion to the continental $\text{CO}_2$ offset at Schauinsland station (Black Forest, Germany) to be around
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1–3 ppm, depending on season. In more polluted areas, such as the upper Rhine valley in Heidelberg, the monthly mean fossil fuel contribution can be as large as several 10 ppm (Levin et al. 1989).

Figure 4a shows the updated Δ14CO2 record for the Heidelberg site. Particularly during the winter months, mean Δ14C depletions in the order of 50‰ are frequently observed. These translate into mean fossil fuel contributions of 20 ppm during this time of the year (Figure 4b). In fact, during summer, the fossil fuel contributions are generally expected to be much lower due to enhanced vertical mixing and dilution of ground-level pollutants but also due to reduced emissions (no domestic heating). The interannual changes of the yearly mean regional fossil fuel contributions are small, however, and a general trend, i.e. towards decreasing offsets, cannot be detected in the last 20 years. In fact, from statistical emissions data of southern Germany no significant trend should be expected from 1987 to 1996. The mean values for the five-year periods from 1987 to 1991 (8.98 ± 0.31 ppm) and from 1992 to 1996 (8.90 ± 0.58 ppm) compare extremely well within ±1%. This opens the possibility of using regional Δ14C observations to validate fossil fuel emissions reductions, e.g. in the frame of the Kyoto Protocol; here, Germany and the European Union as a whole are obliged to reduce their emissions of CO2 and other greenhouse gases by 8% by 2008–2012, a reduction that will be clearly visible and quantifiable using our 14C-based regional observations in the time frame of a five-year period.

CONCLUSION

The radioactive lifetime of 14C is perfectly suited for dating of carbon pools interacting with the atmospheric CO2 reservoir on the time scale of several hundred to several thousand years. The above examples clearly show that our quantitative knowledge of the present gross and net fluxes of CO2 between the dominant carbon reservoirs is significantly based on respective information provided through radiocarbon observations. In particular the use of bomb 14C as a transient tracer in the carbon system provides invaluable insight into processes on the time scale where the largest man made CO2 perturbations took place, namely the last 50–100 yr. The most prominent example is the air–sea gas exchange and the penetration of human CO2 disturbances from the surface into deeper layers of the oceans. Finally, the unique characteristics of fossil fuel derived CO2 being 14C-free allows the tracking of respective emission changes in the past and also in coming years when reliable tools are needed to validate national emission claims of greenhouse gases in the frame of the Kyoto negotiations.

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