COSMOGENIC $^{14}$CO FOR ASSESSING THE OH-BASED SELF-CLEANING CAPACITY OF THE TROPOSPHERE†

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ABSTRACT. An application of radiocarbon ($^{14}$C) in atmospheric chemistry is reviewed. $^{14}$C produced by cosmic neutrons immediately forms $^{14}$CO, which reacts with hydroxyl radicals (OH) to $^{14}$CO$_2$. By this the distribution and seasonality (the lifetime of $^{14}$CO is $\sim$1 month) of the pivotal atmospheric oxidant OH can be established. $^{14}$CO measurement is a complex but unique application which benefitted enormously from the realization of AMS, bearing in mind that $^{14}$CO abundance is of the order of merely 10 molecules per cm$^3$ not only provides $^{14}$CO an independent measure for the OH based self-cleansing capacity of the troposphere, but also enabled detection of $^{14}$C production due to high energy solar protons in 1989. Although its production takes place throughout the atmosphere and does not have the character of a point source, transport processes in the atmosphere affect the distribution of $^{14}$CO. Vertical mixing in the troposphere renders gradients in its production rate less critical, but considerable meridional gradients exist. One question has remained open, namely confirmation of calculated $^{14}$C production by direct measurement. A new sampling method is proposed. The conclusions are a guide to future work on $^{14}$CO in relation to OH and atmospheric transport.

KEYWORDS: atmospheric $^{14}$CO, cosmogenic, hydroxyl radical, radiocarbon AMS dating, solar flares.

INTRODUCTION

Before a fresh cosmogenic $^{14}$C atom (produced by $^{14}$N(n,p)$^{14}$C) enters the main carbon cycle, it almost instantaneously forms $^{14}$CO and a such partakes in atmospheric chemistry for a period of a month or so. This brief intermezzo of not yet being fully oxidized in an otherwise fairly long life is of interest to atmospheric chemists. Bernard Weinstock had been a student of Harold Urey and worked on the Manhattan project. Subsequently he was for 20 years research scientist at Ford Motor Company, Dearborn, Michigan, a company with a brilliant scientist researching air pollution and the impact of combustion engine exhaust gases. In 1969, Weinstock published a brief paper in Science in which he estimated the atmospheric residence time of carbon monoxide. Pandow et al. (1960) already had focused on the fate of $^{14}$C recoil atoms ($\sim$95% forms $^{14}$CO). Subsequently, MacKay et al. (1963) achieved the amazing first $^{14}$CO measurements (by Minze Stuiver) using CO samples obtained from an air liquification plant (huge air volumes needed to be processed) and Mackay suggested $^{14}$CO to be a useful atmospheric tracer. But Weinstock (1969) managed to use these $^{14}$CO data to estimate the lifetime of $^{14}$CO and thus of CO itself ($^{12}$CO and $^{14}$CO for this purpose behave identically). When one of us, as a Dutch student (Groningen Radiocarbon Lab under W. G. Mook and P. M. Grootes) in New Zealand (Institute of Nuclear Science under A. Rafter) in 1976, had written Bernard Weinstock about his approach, proposing to do more $^{14}$CO measurements, an encouraging answer arrived within two weeks by airmail. This was an exposure to the generous attitude of American scientists.

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Why was this estimate of the lifetime of CO so important? Concern about air pollution (e.g., Los Angeles) was certainly a factor, but moreover; when the number of automobiles increases strongly and each produces about 100 g CO (or more) per gallon gas the question is, “Will this toxic gas accumulate in the global atmosphere?” It was early days for atmospheric chemistry. Christian Junge (1959) had listed a mere 9 atmospheric trace gases in his 1959 “Air Chemistry” paper (he noted: “It is very likely that only a few of the trace gases are known”). Furthermore, consensus was that CO had only an anthropogenic source and that soils and the stratosphere were its only sinks. Having a guess of the input (it was thought that CO was mostly of fossil fuel origin) and of its atmospheric inventory did give a lifetime estimate. A totally independent lifetime calculation however, was worth a lot. Weinstock estimated using a rate of production of $^{14}$C at $1.3 \times 10^{19}$ atomsec$^{-1}$, a lower limit of 0.1 year. An excellent number. The big problem in those days was, however, that it was becoming clear that neither the stratosphere nor soils (the known sinks) could take up tropospheric CO (and thus $^{14}$CO) at the rate necessary to obtain such a short lifetime.

The fast developments in early atmospheric chemistry led to the landmark paper by Levy (1971) in which he showed that high concentrations of the hydroxyl radical OH do occur in the troposphere. High levels of formaldehyde were predicted too, which would be a source of CO, a partly natural source. Prior to this paper by Levy, it was assumed that OH abundance was negligible. But suddenly, the troposphere turned out to be a true chemical reactor, alike a slowly burning cold flame. OH is the troposphere’s pivotal oxidant. For instance, nearly the entire annual input of CH$_4$ is removed by this radical, in fact >500 Tg/year. Likewise, CO and most Volatile Organic Carbons are removed by OH. It is hard to imagine how a biological-life-friendly atmosphere could function without OH.

The most compelling case for understanding the cycle and tropospheric distribution of OH is the budget of the important greenhouse gas methane. CH$_4$ has been monitored on a large scale for decades and changes in its natural and anthropogenic sources cause variations that are difficult to trace back. Time and again growth rate changes of the global atmospheric burden CH$_4$ have surprised us. The main factor in the budget of CH$_4$ is of course its sink, namely OH. Numerous detailed studies address the role of changes in OH (e.g., Zhao et al. 2019), which—to complicate matters—are partly attributable to CH$_4$ emission changes and their global pattern.

The UV photolysis of tropospheric O$_3$ leads to O$_2$ molecules and extremely short-lived excited O$^{(1)}D$ radicals. These can react with H$_2$O to form 2 OH radicals. These stable but highly reactive OH radicals live for about a second before reacting with either CH$_4$, CO, organics and so on. One can by measurement of $^{14}$CO calculate the abundance of OH radicals, provided one knows the reaction rate constant for $^{14}$CO + OH (The rate is pressure dependent, at 200 hPa it is $\sim$30% lower than at sea level). This is the “$^{14}$CO method” for determining OH, the subject of this article.

What makes $^{14}$CO relevant for OH determination? First of all, OH has a lifetime of seconds only. Its formation by daylight produces a strong diurnal cycle. Its concentration is much higher in summer than winter leading to a seasonal cycle. It is much more abundant in the tropics than at high latitudes. Its abundance varies with altitude due to mainly water vapour and UV flux gradients. In short, OH is extremely variable in space and time at different scales. Peak values are below $10^7$ radicals per cm$^3$, sub parts per trillion levels. This rarity combined with its high reactivity made it initially impossible to detect OH, but
successful detailed experiments to use tracer $^{14}$CO in a flow reactor with ambient air were done (Felton et al. 1992). This radiochemical method proved to be problematic. To date, in-situ measurement of OH still pose an experimental challenge in atmospheric chemistry (Cho et al. 2021). And even though presently one can obtain in-situ OH data, no large-scale distribution of OH can be reconstructed this way as too many detection systems globally distributed would be needed. Indirect methods for gauging OH are therefore called upon. Instead of trying to measure OH, one observes the removal of certain trace gases that react specifically with OH. The indirect methods have the advantage of mapping the effect of OH on reactive gases, all of which have different reaction times, sources and distributions. The in-situ OH detection is on the other hand essential in achieving a detailed understanding of radical chemistry. Atmospheric chemistry models allow us to construct OH fields, but for verification, measurements are needed. The indirect methods use a trace gas with known input into the atmosphere that predominantly reacts with OH. Initially (Singh 1977; Prinn et al. 1983) this was methylchloroform (CH$_3$CCl$_3$). However, methylchloroform has been almost eliminated from the atmosphere thanks to the Montreal Protocol measures. A recent paper by Patra et al. (2021) notes: “Estimations of tropospheric OH trends and variability based on budget analysis of methyl chloroform (CH$_3$CCl$_3$) and process-based chemistry transport models often produce conflicting results.” In their paper they show that no major change in the chemical removal is likely.

Apart from this problem of phasing out, an inherent problem remains that uncertainties about the rate of release exist, which also applies to other industrial trace gases that potentially may be applied in constraining OH estimates. The conclusions about OH from the use of tracers like methylchloroform, often in painstaking work and finest analytical work and modeling support over decades, are not always in agreement.

Alternatively, $^{14}$CO is a geophysical tracer that will be around for a long time. Work is even in progress to use $^{14}$CO measurements in air extracted from firn and ice to reconstruct OH concentrations in the past. We review the application of $^{14}$CO in atmospheric science with some explanation for a broader readership. We also mention a few less successful studies to help to avoid pitfalls in future research. We do not deal with in-situ production of $^{14}$Ci in solids (e.g., ice).

**APPLICATION**

**Distribution of Tropospheric OH Concentration**

Figure 1 shows the zonally averaged distribution of OH over one year derived with an atmospheric chemistry model. These ever more sophisticated models incorporate hundreds of chemical reactions that take place in the atmosphere. They are in essence atmospheric transport/climate models with embedded chemistry modules dealing with trace gases, particles, their sources, clouds, precipitation chemistry and so on. The tantalizing aspect is that these models show us a distribution of OH that we cannot measure, in contrast for instance to the model derived distribution of O$_3$, which is the most widely measured reactive trace gas. Different models have different OH distributions. Nicely et al. (2020) show that there are still large differences between OH distributions generated in current models. The main feature of model results is the high concentration of OH in the tropospheric tropics (the tropics are the atmosphere’s cooking pot). Basically, the amount of UV light, ozone and water vapor determine OH. With most anthropogenic emissions (which are considerable compared to natural emissions in this period of the Anthropocene;
Crutzen 2021), including the key catalyst NO that sustains the oxidating cycle, being in the northern hemisphere, most models produced a higher OH abundance in the Northern Hemisphere. The publication by Spivakovsky et al. (1991) on OH has been frequently cited, underscoring the key role of OH. Work on using trace gases for assessing OH distribution is of course ongoing. For instance, Wolfe et al. (2019) mapped the variability of OH throughout the global remote troposphere via the synthesis of airborne and satellite formaldehyde observations.

Over the years, much has been learned about OH, in particular its degree of recycling (Lelieveld 2004). For instance, when an OH radical reacts with CH4, it is lost. However, some of the follow-on products are again radicals and with NO these can lead to the re-formation of OH. To “definitely” remove one OH radical takes up to 20 CO molecules (Gromov 2013).

**Distribution of $^{14}$CO Production**

Figure 2 shows the calculated production rate distribution of $^{14}$CO based on Lingenfelter (1963). The mismatch with the OH distribution is large. We see high $^{14}$CO production rates at high latitudes and higher altitudes determined by the interaction of cosmic rays with the atmosphere and the nudging by the Earth’s magnetic field. Lowest production is in the tropics. High production combined with the longer lifetime of $^{14}$CO in much of the stratosphere (by mass) implies that its rate of transport into the troposphere (overall variable with season and stronger in the NH) is a key factor. In the troposphere $^{14}$CO is transported as well, be it to a degree that depends on its regional lifetime, ranging from one (tropics) to several months (at high latitudes).
The result of the contrasting distributions of the rate of $^{14}$C production (Figure 2) and of OH (removal) is shown in Figure 3 as the resulting distribution of $^{14}$CO itself. A model takes transport and the pressure dependence of the reaction rate constant into account. $^{14}$CO mixing ratios are high in the maximum production region (Figure 2) and almost exactly there, OH bottoms out. Moreover, in the tropics where $^{14}$CO is rarer, annual variations are small. High concentrations occur at higher latitudes, accompanied by large annual variations reflecting OH seasonality. Figure 4 shows a high latitude annual cycle of $^{14}$CO measured at Spitsbergen and Alert (Ellesmere Island, Canada), as a result of the ongoing production of $^{14}$CO, its transport and mixing of air masses, the annual averaged level of OH and its strong local seasonal cycle. $^{14}$CO accumulates in the high latitude winter limited only by transport and mixing with air masses at mid latitudes (lower $^{14}$CO). In summer local photochemical removal by OH becomes important.

Not all atmospheric $^{14}$CO is of direct cosmogenic origin. When we consider an air mass its CO content is highly variable because of its relatively short lifetime and the occurrence of concentrated sources, for instance industry (fossil fuel) and forest fires. Forest fires

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**Figure 2** Zonal-annual-mean galactic cosmic ray $^{14}$CO production rate (Lingenfelter 1963). The unit is $10^{-3}$ molec g$^{-1}$ s$^{-1}$ normalized to a global average production rate of 1 molec cm$^{-3}$ s$^{-1}$. The yield of $^{14}$CO from $^{14}$C oxidation is assumed to be 95%. The contour lines refer to $^{14}$C production by solar protons which is discussed later in the text. (Figure from Jöckel 2000.)
(biomass burning in general) releases copious amounts of CO into the troposphere with a $^{14}$C content determined by that of the burned matter. Likewise, the ongoing global removal of CH$_4$ by mainly OH leads to $^{14}$CO production in the atmosphere itself. This "recycled $^{14}$C" in the form of biogenic $^{14}$CO forms a complication in the use of cosmogenic $^{14}$CO for reconstructing OH. Besides this biogenic source, there are poorly quantified sources of $^{14}$CO from the nuclear power industry and possibly from the use of $^{14}$C containing radio-chemicals/pharmaceuticals. There are no indications so far that these two types of sources play a significant role for the total atmospheric $^{14}$CO inventory, but no detailed research has been done so far. To correct for biogenic $^{14}$CO ($\sim$20%) one has to subtract its contribution from measured $^{14}$CO values. Knowledge about the cycle of CO is necessary
and measuring stable isotopic composition of CO namely $^{13}$C, $^{18}$O, and $^{17}$O (a mass independent effect exists (Röckmann et al. 1998)), is helpful. The contribution of recycled $^{14}$CO is higher in the NH and more use could be made of $^{13}$CO observations to improve the corrections. Detailed experimental work on CO isotopologues is available (Röckmann 2002) complemented by use of modeling (Gromov 2010, 2013). A rule of thumb is that 30 nmole/mole biogenic CO is equivalent to 1 molecule $^{14}$CO per cm$^3$ (Brenninkmeijer 1993).

A 10% uncertainty in the estimate of biogenic $^{14}$CO results in a $\sim$2.5% uncertainty of the inferred amount of cosmogenic $^{14}$CO.

RESULTS

After Mackay’s pioneering data, Volz et al. (1981) carried out systematic $^{14}$CO measurements. They removed CO$_2$ from 200 m$^3$ sample air at the sampling site, oxidized its CO content to CO$_2$ (20–40 mL) which was collected for proportional counting. They obtained the first seasonal cycles of $^{14}$CO and measured at different locations, including shipboard measurements. Using a global 2D model they derived a tropospheric OH concentration below 10$^6$ per cm$^3$.

Subsequently $^{14}$CO measurements were set up at the Institute of Nuclear Sciences in Lower Hutt, New Zealand. Roger Sparks and colleagues had turned a decommissioned Pelletron accelerator from Australia into a $^{14}$C AMS system (Sparks et al. 1984). Using AMS the required sample size for $^{14}$CO dropped from 200 to less than 1 m$^3$. AMS was a breakthrough for $^{14}$CO applications, especially because air samples of that smaller size could be collected in pressurized cylinders anywhere for processing in the laboratory. The required air sample size can be estimated as follows. When a conservative estimate of the efficiency of $^{14}$C detection of 1% is used in relation to the initial amount of $^{14}$C in the CO$_2$ sample and a 1% precision is required, the presence of 10 $^{14}$CO molecules per cm$^3$ air STP...
implies a minimum air sample size of 100 L. Besides $^{14}$C determination, the CO derived CO$_2$ samples are analysed by stable isotope mass spectrometry to determine the $^{13}$C/$^{12}$C and $^{18}$O/$^{16}$O ratio. These numbers are useful for reconstructing the local budget of atmospheric CO which helps to estimate the mentioned biogenic $^{14}$CO fraction.

The CO extraction method is described by Brenninkmeijer (1993). Sample air is cryogenically purified by means of “Russian doll” traps submerged in liquid nitrogen. These traps consisting of nested borosilicate glass fibre thimbles remove all condensable gases and achieve the reduction of the CO$_2$ content of the air to below 0.1 nmole/mole. Depending on the sample CO content, the specific activity of CO is up to 1000 pMC (lowermost stratosphere) at CO mixing ratios down to 50 nmole/mole or less. Residual traces of CO$_2$ would dilute the $^{14}$CO samples in terms of pMC. After this purification step, the CO content is specifically oxidized to CO$_2$ using an iodine pentoxide-based reactant. The resulting micromole CO$_2$ samples are measured manometrically which gives the value of the CO mixing ratio. After off-line stable isotope mass spectrometry, the small sample is diluted with “dead” CO$_2$ for obtaining a CO$_2$ sample of sufficient size for target preparation. The advantage of the oxidant used (I$_2$O$_5$) is that it does not scramble the original oxygen isotopic composition of the CO because it merely adds an oxygen atom of known isotopic composition to a CO molecule. Insofar air samples are collected in cylinders the CO content can increase in time. This spurious CO would generally affect the $^{14}$C result little because of a low or zero $^{14}$C content. Sampling and storage systems limit spurious CO “growth” to less than the percent level. Irrespective of this, delay between sampling and processing can lead to significant in situ produced additional $^{14}$CO, for which a correction must be applied (Lowe et al. 2002).

The accuracy and precision of a $^{14}$CO determination is composed of the AMS error, the manometric error in dilution of the CO sample (dilution up to a factor 10), and the error in determining the CO content of the air sample and can be kept at about 1%. Adding the larger uncertainty of the biogenic fraction estimation of 2%, gives a combined uncertainty that is below the variability of $^{14}$CO in most air masses (remote SH air showed the least variability) during a typical 1-hr air sample collection period using a high-pressure compressor. Recently, Petrenko et al. (2021) reported an air sample collection and processing method for much smaller air samples, namely less than 100 L. This avoids the use of high-pressure sample air cylinders, which are more expensive and time consuming to transport from the point of collection to the laboratory.

Using this method (with on-site CO extraction, however) Brenninkmeijer et al. (1992) had measured $^{14}$CO data at the clean air station Baring Head, New Zealand and compared the results for the seasonal cycle with those of Volz et al. (1981) for Schauinsland, Germany. Lowest summertime values at Baring Head were about 5 molec cm$^{-3}$, compared to 10 at Schauinsland. These are two totally different sites from an atmospheric chemistry point of view, namely clean mid-latitude maritime SH air against continental NH air in the Black Forest. After correcting for the effect of the solar cycle and the difference in latitude and accounting for the biogenic $^{14}$CO fraction, Brenninkmeijer et al. (1993) concluded that OH levels in the SH were higher.

Using a large set of $^{14}$CO data Jöckel and Brenninkmeijer (2002) developed a climatology by which Jöckel et al. (2002) could evaluate global atmospheric transport (including stratosphere-troposphere exchange) and the hydroxyl radical (OH)-based oxidation in three-dimensional (3-D) atmospheric chemistry transport models (CTMs) and general circulation models.
(GCMs). They showed that the correct representation of phase (annual cycle) and strength of the STE, including its asymmetry between NH and SH, is essential to reproduce the observation based seasonal cycle of $^{14}$CO. Moreover, their results did not support the previously inferred interhemispheric asymmetry in the OH abundance with an on average higher concentration in the SH.

A question that had to be addressed was the response time of the $^{14}$CO concentration to changes in total $^{14}$C production caused by modulation through solar activity (Jöckel et al. 2000) which is related to, but not the same as the lifetime. Figure 5 shows model lifetimes of $^{14}$CO throughout the atmosphere. At the surface lifetimes are below 1 month for parts of the NH tropics.

Figure 5  Lifetime of $^{14}$CO in days (color scale) simulated with the EMAC chemistry climate model (year 2000, Jöckel et al. 2016). The troposphere shows a considerable asymmetry with higher OH abundance in the NH. (Please see electronic version for color figures.)
Quay et al. (2000) applied a 2D transport model to their measured and several other $^{14}$CO datasets. They found good agreement and emphasize that in their model an increase in advection and slower stratosphere-troposphere exchange in the SH would best explain the lower $^{14}$CO levels of Baring Head.

The properties of $^{14}$CO in relation to OH and transport have been characterized in detail using the TM5 global chemistry transport model and inverse modeling (Krol et al. 2008). It was established that in the tropics $^{14}$CO is sensitive to regional OH. Therefore, it is a useful indicator for the local “speed” of the rate of removal of reduced gases. At the hemispheric scale however, transport of $^{14}$CO plays a bigger role. Variability is introduced by synoptic scale processes. The model research also compares $^{14}$CO with CH$_3$CCl$_3$, which appears to be more sensitive to bulk tropical OH. The model results closely follow the annual cycle of $^{14}$CO at baring Head, NZ, with an offset probably due to the assumed production rate. Recommendation about optimizing locations for measurements can be gleaned from this work.

The success of the $^{14}$CO method for inferring changes in OH is best illustrated by Figure 6 from Manning et al. (2005). It shows the longest available record, namely that of Baring Head including data from Scott Base, Antarctica. Cylinders with sample air from Antarctica often were stored on site for months before shipment was possible. Lowe et al. (2002) corrected the $^{14}$CO content for in-situ production. One clearly sees the solar cycle induced change of about 20% for the period 1989 until 2003. The seasonal cycle due to OH is about $\pm$ 40%. When taking the $^{14}$CO production variations into account and applying an atmospheric model the residuals relative to a fitted curve are within 10%, except for two
periods. The first one is in 1991, after the eruption of Mount Pinatubo. Reduced UV radiation in the troposphere did lead to a reduced production of OH (Dlugokencky et al. 1996) and thus an increase in $^{14}$CO. The second period is 1997/1998. During this period a strong El Niño led to intensive biomass burning (Southeast Asia, particularly Indonesia) thus burdening the troposphere with reduced gases apparently leading to a reduction in OH in this part of the SH. Also, radiative effects of a large atmospheric load with soot can have influenced OH levels. There are no other observations of this kind that document such changes in OH.

Several experiments have addressed free tropospheric and lowermost stratospheric $^{14}$CO measurements using air samples collected using aircraft. Initially modified scuba diving compressors were used (Mak and Brenninkmeijer 1994) and later membrane compressors (Brenninkmeijer et al. 2007; Brenninkmeijer and Roberts 1994) with the low ambient pressure being a challenge considering the large amount of air required.

Detection of 1989 Solar Proton Events

Figure 6 shows the calculated $^{14}$C production rate during solar cycle 22. The effect of the modulation of the cosmic ray flux by solar activity is considerable. Apart from this 11-yr gradual modulation, short-term fluctuations in $^{14}$C production do occur. For calculating the actual variation in the production of $^{14}$C, neutron count rates were used (Lowe et al. 2002). Both these longer-term and short-term variations have to be taken into account when reconstructing the actual $^{14}$C production rate for application to OH.

Besides this, independent, sporadic bursts (lasting days only) of atmospheric $^{14}$C production are due to high energy solar protons from coronal mass ejections. Such events can occur throughout a solar cycle and lead to elevated neutron count rates at ground level (Ground Level Enhancements, GLE). Lingenfelter and Flamm (1964) estimated the average $^{14}$C production due to solar protons to be at most 5% of the cosmogenic production. For events to be of significance for $^{14}$C production the proton energy has to exceed $\sim 30\text{MeV}$ and the flux has to be sufficiently high. During solar cycle 22 several high energy proton events occurred. Three of these events between August and December 1989 carried sufficient high energy protons to produce—via additional neutrons—$^{14}$C that was detectable above the background. The GLE on 29 September 1989 was the largest recorded since the giant event of 23 February 1956 (Hessel de Vries submitted a paper about its effects on background count rates in low level $^{14}$C proportional counters a mere 2 weeks later; de Vries 1956). Figure 2 (contour lines) shows that the solar protons in 1989, despite their high energy, produced $^{14}$C higher in the polar stratosphere than the typical maximum of the production by galactic cosmic rays. By analysing the Baring Head $^{14}$CO record, using modeling and satellite data for the proton flux, Jöckel et al. (2003) established that between 3 and 30% additional $^{14}$C production due to solar protons had occurred in 1989. Such an application of $^{14}$CO is a unique case. To extract the solar flare signal (Figure 7) the period June 1990–June 1991 was used a reference (no significant solar flares). The model calculated increase overlaps with the measured $^{14}$C enhancement.

Calculated Production of $^{14}$CO in the Atmosphere

The application of $^{14}$CO requires accurate information about the $^{14}$C production distribution (on average about 7 kg $^{14}$C is produced annually). Its subsequent fate by oxidation to $^{14}$CO$_2$ is then determined by transport and OH chemistry, with a difficult aspect being its transport from the stratosphere ($^{14}$C production but low OH) into the troposphere (also production but high
The calculation of $^{14}$C production by Lingenfelter (1963) (Figure 2) has been used most. Obviously, the position of the tropopause (the dynamic interface between the two spheres) is of no relevance to the $^{14}$C production field. All neutrons generated by nuclear reactions in the earth atmosphere lead to the production of $^{14}$C via $^{14}$N(n,p)$^{14}$C. Lingenfelter used neutron count rates to derive 3D neutron production rates for solar maximum and minimum conditions. O’Brien et al (O’Brien et al. 1979) carried out calculations based on hadronic cascades in the atmosphere. Masarik and Beer (1999) developed a pure physical model. All three approaches give a few % more $^{14}$C production in the SH due to the Earth’s magnetic field distribution. In the given order of principal author 52 (Lingenfelter), 57 (O’Brien) and 64% (Masarik), respectively, is produced in the stratosphere (Jöckel 2000). Whereas SH-NH differences are small, the vertical distribution needs attention in view of the role of stratosphere-troposphere exchange. Jöckel et al. (Jöckel et al. 1999) showed by means of a model that tropospheric $^{14}$CO concentrations are relatively insensitive to these source differences and therefore not critical for the $^{14}$CO method. More extensive and refined work on cosmogenic isotope production was carried out by Usoskin and coworkers (Poluianov et al. 2016). Not only did this work at long last close the gap between calculated global $^{14}$C production and observations in the carbon cycle but also revealed a larger solar modulation of $^{14}$C production than in previous works.

Figure 8 shows vertical profiles of production rates from the four approaches. There are significant differences of a factor 5 at the surface and in the free troposphere. The steep production rate gradient in the troposphere is a lesser concern because of the relatively rapid vertical exchange of air masses and thus mixing of $^{14}$CO in the vertical. For local $^{14}$CO production the gradient matters.

In-Situ $^{14}$CO Production Experiments

From an experimentalist’s point of view, it is comforting to know that (in principle) it is straightforward to measure the actual production of $^{14}$C (as $^{14}$CO) by exposing a quantity
of air to cosmic radiation. Mak et al. (1999) reported on such experiments based on exposing cylinders with compressed air at different altitudes and latitudes (South Pole Station and Mount Cook, NZ) and at different heights on a tall tower (Boulder Atmospheric Observatory, USA). The fundamental problem is that production rates at the surface are low (production peaks at 100 hPa), which renders the result as a measure for the column integrated production intrinsically less informative. An additional complication is the enhancement of the neutron flux close to the surface. The purpose of the tall tower experiment was to deal with this. The results showed that measured production rates were consistently lower by $\sim 50\%$ than values derived from Lingenfelter’s calculations.

Further experiments were made using cylinders with compressed air stowed in passenger aircraft (Ramsey et al. 2007). Such experiments are ideal from the point of view that civil aircraft happen to cruise at altitudes where $^{14}$C production peaks. By simulating the aircraft flights over weeks between cylinder exchanges in a model of the $^{14}$C production field expected accumulated production could be calculated. Also, calculations were carried out to correct for local neutron enhancement due to aircraft structure and payload. But there also was a big obstacle. Namely, only cylinders that were certified to fly on passenger aircraft could be used. These aviation oxygen supply (mild steel) cylinders were not the treated inert aluminum cylinders normally used in $^{14}$CO air sampling and storage. During the experiment, the aviation cylinders had to undergo hydraulic pressure tests. Altogether 90 experiments could be carried out. Sets of cylinders were kept in the laboratory for background testing. Measured was not only $^{14}$CO but also $^{14}$CO$_2$. The results showed not a $^{14}$CO$_2$ to $^{14}$CO ratio of about 5%, as expected on the basis of the early radiochemistry experiment, but about 20%. No explanation could be given. Furthermore, production rates were well above those calculated using the Masarik and Beer distribution. Despite a considerable effort the results were less conclusive.


14CO as the Precursor of 14CO2

Because soils and oceans take up negligible amounts of 14CO and the chemical fate of CO is its oxidation, nearly all primary cosmogenic 14CO molecules will form 14CO2. In equilibrium both 14CO and 14CO2 have exactly the same flux (some deviations occur because of 14C from the nuclear weapons tests and variations in the rate of uptake of 14CO2 by the biosphere and oceans). As a result of their vastly different residence times however, the atmospheric inventory of 14C as 14CO is more than 1000 times smaller than 14C as 14CO2. Also, their distributions in the atmosphere bear no resemblance. Kanu et al. (2016) calculated an independent cosmogenic production rate of 14C using data from air samples collected in the stratosphere up to above 30 km. Their 14CO2 data show a compact correlation with decreasing N2O mixing ratios, a gas that is destroyed in the stratosphere. Δ14CO2 reached about 140 per mil. Brenninkmeijer et al. (1996) report 14CO values in the lowermost stratosphere over Antarctica of up to 150 molec cm⁻³, 10 times tropospheric values. To our knowledge, no model analysis for 14CO2 production from 14CO in the stratosphere have been published so far. Our preliminary model calculations show that ~30% of 14CO produced in the stratosphere is exported to the troposphere. About 10% of the tropospheric production (which is less than 50% of the total) ends up in the stratosphere. Therefore, the net 14CO loss for the stratosphere is only 20%. Consequently, 80% of 14CO in the stratosphere is oxidized there to 14CO2.

Firn Air Measurements of 14CO

The analysis of air extracted in-situ from firn allows reconstruction of past atmospheric composition of lesser reactive trace gases. In the case of 14CO hopes were high to be able to shed some light on changes in OH over past decades. Indeed, CO itself appeared to be stable in firn air. Although the in-situ production of 14C of the air retained in the firn layers is small, in situ production in the solid phase appeared to complicate matters. As can be seen in the paper by Assonov et al. (2005) 14CO levels in firn air (Dronning Maud Land, Dome Concordia and Berkner Island) were above atmospheric levels and increased considerably near pore close-off depth. This points to a very complicated situation when one wishes to reconstruct past OH concentrations. In-situ production of 14C (Jull et al. 1994) complicates the use of 14CO in firn air. A sufficiently accurate correction for in situ production in the 14CO firn air application, will be difficult, if possible, at all. Nevertheless, work on this is in progress (Petrenko et al. 2020).

Suggested Improvements for the 14CO Application

Given the lack of alternatives for OH and the improvements in modeling including the recent 14C production calculation by Usoskin’s group, 14CO work should continue with improved methods. The optimal number and location of sampling sites can be better determined now (Krol et al. 2008). From an air sampling method point of view, a degree of time integrated collection optimizes the results. Short term variations are average by this. Up to now it has been a major effort to collect, compress and ship a dry air sample, but why this fast sampling? There is a lot of air for sampling, in contrast to many other 14C AMS applications where smallest amounts need to be assayed. Volz et al. (1981) and Brenninkmeijer et al. (1992) used on-site extraction, which was cumbersome. Thereafter, Mak and Brenninkmeijer (1994) used the experience obtained in Boulder (NOAA) for using high-pressure air (diving air) compressors. This facilitated getting samples from remote locations enormously and many 14CO data were collected (e.g., Rom et al. 2000;
Gros et al. 2001). An excellent example of a sampling technique is the pioneering work by Östlund (1970), who measured the Tritium content of atmospheric H₂ already in 1970 using automated collection systems (HT is rarer in the atmosphere than ¹⁴CO) at remote locations, including Baring Head.

For the ¹⁴CO application one should ironically not think about small air samples as we did so far. We propose here that sample air, say 5000 L (flowrate of 1 L/min) is dried using Nafion® and subsequently cryogenically purified using Russian Doll traps. The next step is oxidation of the CO content at ambient temperature using a CO selective catalyst (e.g., Sofnocat®). The resulting CO₂ is collected on a molecular sieve cartridge for shipment to the AMS lab. Alternatively, molecular sieve cartridges are used for the thorough pre-purification after sample air drying. A large volume cartridge with molecular sieve would lead to a CO gas chromatography separation effect. Smaller cartridges in parallel, that are sequentially heated and backflushed for re-generation, avoid such a problem. After this set of traps one final molecular sieve cartridge acts as a safety barrier before the air flow reaches the catalyst. Another advantage of using molecular sieve (instead of the Russian doll traps) is that cooling systems are not required. Moreover, the air processing occurs at above atmospheric pressure. Cryogenic purification requires sub-ambient pressures.

The advantages compared to the method developed by Brenninkmeijer et al. (1999) are (1) large, “robust” CO-derived CO₂ samples, (2) time-integrated air sampling, (3) samples can be shipped without restrictions, (4) storage will most likely not lead to in situ ¹⁴C formation, and (5) the difficult I₂O₅ oxidant need not to be used (this “Schütze reagent” is very sensitive to moisture and difficult to prepare). The disadvantage is that no δ¹⁸O(CO) information is obtained in this way. CO mixing ratios are already obtained at many sampling sites in the framework of Global Atmosphere Watch. What remains difficult is the collection of air samples using civil aircraft. The CARIBIC Boeing 737 used for ¹⁴CO work (Brenninkmeijer et al. 2007) had a typical cruise speed of well over 800 km per hour, which meant that with the given air sample collection time of about ½ hour, sampling in the tropopause region where civil aircraft most frequently fly at mid-latitudes an averaging between ¹⁴CO in tropospheric and stratospheric air masses was inevitable.

CONCLUSIONS

The distribution and possible changes of OH in the troposphere have been a bone of contention for years (one can find several lively discussions in the literature, e.g., Spivakovsky 1991). It was (and still is) not known if OH does change over large parts of the troposphere or even globally on a scale of several years and if OH remained stable at all over the current decades of the Anthropocene. The clear detection of reductions in OH due to the 1991 Mount Pinatubo eruption and the 1998 El Nino by Manning et al. (2005) is an important demonstration of the use of ¹⁴CO. Over many years emphasis has been on using methyl chloroform records, but this substance disappears from the atmosphere after having been banned by the Montreal Protocol. Its mixing ratio dropped from over 100 to a few pmole/mole (lifetime about 5 years). Clarissa Spivakovsky wrote about CH₃CCl₃ based estimates of tropospheric OH “Last Offerings of a Departing Friend” (Spivakovsky et al. 2000). A main concern is uncertainty about its source strength, the role of the oceans with a dwindling atmospheric abundance. Nevertheless, a recent paper seems to have given methyl chloroform back a brief lease of life (Patra et al. 2021). Some other industrial trace gases also help to “constrain” OH, but there remain inherent uncertainties. It is difficult.
The initial interest in $^{14}$CO from an atmospheric application point of view was definitely the determination of OH. In the application of atmospheric models to support this use, $^{14}$CO also proved to be a tough test for the simulated transport processes (Jöckel et al. 2002). A difficult aspect in this is the cross-tropopause transport. At civil aircraft cruise-altitude we frequently measured changes from 10 to almost 100 $^{14}$CO molecules per cm$^3$ STP when crossing tropopause folds (CARIBIC unpublished results). $^{14}$CO correlates closely with O$_3$ in the lowermost stratosphere. This strong tropopause gradient means that a cut-off low releases $^{14}$CO enriched air into the troposphere. Such information—detailed as it is—is difficult to model. Today’s models can have a higher vertical degree of resolution in the tropopause region, which is helpful.

Given the improved $^{14}$C production calculations and continued $^{14}$CO applications it is worthwhile to carry out improved in-situ production experiments by exposing air samples. It seems to be a trivial undertaking, but so far, no really useful conclusions have been reached.

$^{14}$CO has the great advantage that it tracks closely the seasonal cycle of OH. Corresponding variations in CH$_3$CCl$_3$ are below 3% only. $^{14}$CO can help to derive OH levels on a regional scale. $^{14}$CO has a source that is probably not affected noticeably by industrial emissions, it is a geophysical tracer. From this point of view is stands to reason to continue $^{14}$CO observations, and indeed recently further improvements were just reported (Petrenko et al. 2021). Current atmospheric chemistry models are of great complexity and the reconstructed rate of removal of trace gases by OH remains an essential piece of output. The discussion about OH is ongoing. In this, measurement of $^{14}$CO provides a definite test option, in absence of other suitable trace gases. Support for long-term times series, namely Baring Head (the $^{14}$CO “Mauna Loa”), will be most helpful.

In summary, given a need to gauge tropospheric OH in a changing atmosphere and the lack of suitable trace gases, and given the improvements in cosmogenic $^{14}$C production knowledge, and given improvements in atmospheric modeling, and knowing an optimized sampling strategy and using an improved $^{14}$CO measurement methodology as suggested, continued progress is possible.

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