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Atmospheric Depositional Fluxes of ⁷Be and ²¹⁰Pb at Galveston and College Station, Texas

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The bulk depositional fluxes of ^{21 o}Pb and ⁷Be were measured at a cosstal (Galveston) and an inland (College
Station) station for about 3 years, between 1989 and 1991 The annual depositional fluxes of ^{7B}e and ^{21 o} The volume weighted 210Pb concentrations, when normalized to the amount of precipitation, seem to be constant over the time period of this study. Four to six heavy rain events (> 5 cm) in a single day account for 20-30% of the annual deposition of 7 Be and 2 lip. Such events account, however, for only about 4-6% of the total number of
rainy days in a year. The dry depositional fluxes of these nuclides appear to be a significant fraction of the b depositional flux only during the months when there is very little rain. The fraction of dry to total depositional flux of 21 OPb appears to be higher than that of ⁷Be. The strong positive correlation between ⁷Be a correlation also indicates that a major portion of the air masses that brings precipitation to Galveston and College
Station is of continental origin. Our data therefore suggest that ⁷Be and ²¹⁰Pb cannot be used as ind atmospheric tracers in our coastal station. This observation is consistent with those observed at many other continental and coastal stations.

INTRODUCTION

Beryllium 7 (half-life $= 53.3$ days) and lead 210 (half-life $= 22.1$ years) are two radionuclides which have been widely used as tracers and chronometers in aquatic and atmospheric systems [Wogman et al., 1968; Perkins et al., 1970; Martell, 1970; Poet et al., 1972; Craig et al., 1973; Somayajulu and Craig, 1976; Young and Silver, 1974, 1980; Bacon et al., 1976; Benninger et al., 1979; Krishnaswami et al., 1980; Aaboe et al., 1981; Turekian et al., 1983; Olsen et al., 1985, 1986; Casey et al., 1986; Dominik et al., 1989; Santschi and Honeyman, 1989; Todd et al., 1989; Brost et al., 1991; Feichter et al., 1991; Kritz et al., 1991; Schuler et al., 1991; Wieland et al., 1991]. Even though the sources of these two nuclides to the atmosphere are distinctly different, both are highly particle reactive and thus get attached to aerosols in the atmosphere soon after production.

Beryllium 7 is produced throughout the atmosphere as a product of the spallation of oxygen and nitrogen nuclei by energetic cosmic rays [Lal et al., 1958]. Owing to its short mean life (76.9 days) and the longer residence time of stratospheric aerosols (about 1 year [Kuroda et al., 1962]), most of the $7B$ e nuclei that are produced in the stratosphere do not readily reach the troposphere except during spring when seasonal thinning of the tropopause takes place at midlatitudes, resulting in air exchange between stratosphere and troposphere. Since $7Be$ is of cosmogenic origin, its flux to the Earth's surface has a latitudinal dependence [Lal and Peters, 1967]. Its concentration in the air increases with increasing altitude from the surface of the Earth, and its atmospheric flux to the Earth's surface should be independent of geography at any particular latitude [Turekian et al., 1983]. In other words, the standing crop of ⁷Be in the atmosphere at a particular latitude should be the same irrespective of whether it is located over the ocean or a continent.

In contrast, 210Pb is produced by radioactive decay from its progenitor, ²²²Rn. This Rn nuclide, which is a noble gas isotope in the ²³⁸U decay chain, emanates primarily from land surfaces. The concentration of 210Pb in the air over the continents decreases with

its elevation from the ground due to a decrease in the concentration of 222Rn [Moore et al., 1973]. As a consequence of its continental source, the ²²²Rn flux from soils is about 100 times higher than the oceanic ²²²Rn flux [Wilkening et al., 1975]. Therefore the standing crop of $210Pb$ in the atmosphere strongly depends on the longitude, depending on whether it is above the ocean or a continent.

These two radionuclides with their different source functions are therefore useful in studies designed to better understand the mechanisms and rates of removal of aerosols. The activity ratio in the air, as well as in the precipitation, is expected to vary with location and time. The $7Be/210Pb$ activity ratios in the precipitation are useful for two purposes: (1) the variations in the ratio can be used to predict the physical behavior of some of the chemical species injected into the troposphere and (2) they could be used, in principle, to predict the elevation at which the cloud condensation takes place. For example, aerosols from the lower part of the troposphere will be enriched in ²¹⁰Pb over ⁷Be, while those from the upper troposphere will be enriched in ⁷Be. Thus qualitative information on the elevation at which the precipitation originated can, in principle, be obtained from this ratio. Precise determination of the value of the activity ratio is required to obtain this information.

Our investigation is designed to shed light on the following questions: (1) How do the annual ⁷Be and ²¹⁰Pb depositional fluxes vary with varying amounts of precipitation? (2) How do the $7Be/210Pb$ activity ratios from individual rain events vary within the same month? (3) Do these ratios vary from season to season? (4) What are the relationships between activity ratios and amounts of precipitation? (5) Can ⁷Be and ²¹⁰Pb be used as independent atmospheric tracers for oceanic and continental air? To answer these questions, we have collected bulk atmospheric precipitation samples for ⁷Be and ²¹⁰Pb analysis at two sites, one in Galveston and the other in College Station, Texas. In addition, a compilation of all earlier data has been made and is compared with our data. This is one of the few studies in which the bulk deposition samples were collected and analyzed for ⁷Be and ²¹⁰Pb at very close time intervals.

MATERIALS AND METHODS

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Two sampling sites were chosen for this study, the roof of the Kirkham Hall (11 m above ground level) at the Mitchell campus of

Texas A&M University, Pelican Island, Galveston (29°18'N, 94°48'W) and the roof of the Oceanography and Meteorology Building (58 m above ground level) in College Station (30"35'N, 96'22'W), Texas. Polyethylene drums of 20O-L capacity and a surface area of 2800 cm^2 were used as rain collectors. Bulk deposition samples were obtained from December 1988 through February 1992 for Galveston and from June 1989 through February 1992 for College Station. Since evaporation rates in the containers were high in some of the samples, the precipitation record for Galveston was obtained from a meteorological station located about 3 km southeast of the sampling site, and for College Station it was obtained from the airport about 3 km away from the sampling site. Although during periods of individual convective storms the amounts of precipitation could be different, the integrated amounts of precipitation over the sampling period for most of our samples are likely to remain constant within a spatial distance of 3 km. A comparison of the meteorological record from the Galveston site with the record obtained from a rain gauge installed with our rain collector during the second half of the collection period indicates that most of the time the amounts of precipitation in individual rain events are essentially the same for the two sites, which are only 3 km apart. Since the rain collector was exposed to the atmosphere continuously, the sum of wet and dry fallout (equal to bulk fallout) was collected. To determine the dry fallout alone, the 200-L drum's lid (2800 cm², height of 3.5 cm) was deployed 3 times continuously for a finite period of time (7-14 days) when there was no rain. Since we primarily use these radionuclides as tracers in aquatic systems where deposition rates occur onto a water surface, the lid was filled with distilled water up to 1.5 cm $(-1/2)$ of the total height).

The drums were cleaned by repeated 6 M HCI rinsings. Prior to each deployment, 1 mg of stable Be and Pb in 100 mL of 6 M HCl were added to the drum or the lid. The acid prevents adsorption of Be and Pb isotopcs onto the polyethylene drum surfaces. The collectors were retrieved at monthly intervals in College Station and more frequently in Galveston. The more frequent retrieval enabled us to determine the fallout of these nuclides during thunder storms and other short-term rain events. After the collectors were emptied, the walls of the drum were rinsed with 2 L of 6 M HCl, which was then added to the sample. Following this thorough acid cleaning and a distilled water rinsing, the collectors were redeployed after the addition of dilute acid and stable Pb and Be carriers. In the first set of Galveston samples (during the first year), the samples were evaporated to about 5O mL, and the reduced volumc was subsequently filtered through a 0.4-um Nuclepore filter paper. Ferric chloride was added to this filtrate, which had a $pH < 1$, and then, after 1 hour of equilibration, $Fe(OH)$ ₃ was precipitated by adding NH₄OH. The precipitate was then filtered through a Whatman 42 filter. In all College Station samples and in the Galveston samples after the first year, known amounts of FeCl3 were added and Fe(OH)3 was precipitated directly from the original rainwater samples, without an evaporation step, by adding NH4OH. Thc precipitates were separated by filtering through a Whatman 42 filter paper. The residues retained on the filter were washcd with 50 mL deionized water and redissolved in a minimum amount of 6 M HCl. The solution was subsequcntly dried on a hot plate.

The dried residues were packed into 10-mL gamma counting vials and counted on a high-purity Ge well detector coupled to a Canberra 5-100 multichannel analyzer. Most of the recently published reasearch has used gamma counting methods to determine the ²¹⁰Pb concentrations in soils and rainwater (soil profile [Graustein and Turekian, 1986]) [Olsen et al., 1985; Todd et al., 1989; Monaghan, 1989; Dibb, 1989; Schuler et al., 1991]. This gamma counting method eliminates the problems of processing each sample twice (first, in situ ²¹⁰Po is separated from ²¹⁰Pb and then, after waiting for about 3-6 months, the $210p_0$ grown in from 210Pb is measured). Furthermore, it involves relatively little manual labor and analytical time.

Typically, the samples were counted for about 6-24 hours, depending on the activity of 2l0Pb in the sample, since all the

samples had relatively high concentrations of $7Be$. The peak analysis of ⁷Be (I = 10.3%, 477.7 keV) and ²¹⁰Pb (I = 4.05%, 46.5) keV) was done using SPECTRAN-AT peak analysis soffware (CANBERRA Company). There is no peak background in the 210Pb energy range, 44.5-48.5 keV. The count rates of rainwater samples were typically between 2 and 1O counts per minute (cpm). In selected samples, the net counts obtained from the peak analysis software were compared with the peak analysis carried out manually in order to convince ourselves that both methods essentially gave the same net counts. In a suite of samples from a sediment core, 210Pb concentrations were measured by alpha and gamma spectrometry. The $210Pb$ concentrations obtained by these two different methods were in good agreement, always within \pm 5% (M. Baskaran, unpublished data, 1992). The peak/Compton ratio for 60 Co (1332 keV) was 45.0:1.

One of the 7Be standard solutions was obtained from Swiss Federal Institute of Water Resources and Water Pollution Control (EAWAG), Switzerland, but originated from the Laboratoire de Meteorologie des Rayonnements Ionisants, Gif-sur-lvette. France, and the other from Brookhaven National Laboratory, New York. Our gamma counting equipment was calibrated with both of these $7Be$ standards. The $210Pb$ standard solution was obtained from Amersham Company and was calibrated with respect to a working standard which in tum was calibrated with respect to a National Institute of Standards and Technology standard. Using these standard solutions, the disintegrations per minute (dpm)/cpm conversion factors for various vial geonretries were determined. This conversion factor was then used to calculate all 7Be and 2l0Pb activities. The error associated with the dpm/cpm conversion factor was always less than 1% . The final $7Be$ concentrations and deposition fluxes were corrected for radioactive decay from the eud of sample collection to midtime of counting, as well as for the ingrowth and decay during the deployment period. The yields of chemical extraction were determined using the Pb and Be recoveries in the final $Fe(OH)$ ³ precipitate measured by atomic absorption spectrometry. The overall propagated error $(\pm 1 \text{ sigma})$ in the value of thc final concentration and flux is estimated to be less than 5% for $7Be$ and less than 10% for $210Pb$.

RESULTS AND DIscUssIoN

The amount of precipitation, periods of rain collector deployment, specific concentrations, and depositional fluxes of 7Be and ²¹⁰Pb results are given in Tables 1 and 2 for Galveston and College Station, respectively. Between December 16, 1988, and February 24, 1992, 67 bulk deposition samples and three dry-only deposition samples werg collected and measurcd at Galveston. In College Station, between June 1989 and May 1992, 22 bulk deposition samples were collccted. One bulk deposition sample was lost from both Galveston (May 17 to June 14) and College Station (March 22 to April 22, 1991) during sample collection and handling. The annual amount of precipitation in 1989, 1990, and 1991 was 103, 97, and 150 cm, respectively, at Galveston and 80, 98, and 146 cm, respectively, at College Station. The rainfall in 1991 for both Galveston and College Station was thus about 50% higher than during the other two years.

The monthly precipitation data for Galveston are plotted in Figure I for the years 1989-1991. Generally, the amount of prccipitation is higher in the summer months. In 1991 (and also in 1992), January and February months also had relatively high rainfall. The monthly precipitation data for College Sration are plotted in Figure 2. In general, the precipitation records for the two stations are similar.

Monthly and Seasonal Variations of 7Be and 210Pb Fluxes

At Galveston. the samples were collected more frequently between May 1991 and February 1992. In those samples which were collected in between two calendar months, the monthly flux was calculated by assuming the specific concentration during the

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Fig. 1. Total monthly precipitation at Galveston, Texas,

whole deployment period was constant and by using the amount of precipitation during the deployment period. College Station rainwater collectors were sometimes deployed for periods longer than I month, and thus reliable estimation of monthly fluxes was not attempted under those circumstances, The frequency distributions of the depositional fluxes of $7Be$ and $210Pb$ for Galveston in individual bulk depositional samples are plotted in Figures 3 and 4, respectively. These figures indicate that in $>80\%$ of these individual samples, the variations of the depositional fluxes are within a factor of 10. The temporal trend of the monthly bulk depositional fluxes of ⁷Be and ^{210p}b at the Galveston station is shown in Figure 5. There are no clear consistent seasonal trends in the distribution of monthly fluxes during the study period. However, the amounts of precipitation seem to be controlling the magnitude of the depositional fluxes. For example, at Galveston, the 38 months for which the depositional fluxes are reported in

Table 1, the highest monthly fluxes were during winter of 1991, coinciding with relatively high rainfall during those months. During January and February 1991, the ⁷Re and ²¹⁰Pb fallout was equivalent to about 29.0% of the total annual fallout, which is similar to the percentage of the amount of precipitation of 43 cm, which was also 29% of the annual rainfall. During May and June, the $7Be$ and $210Pb$ fallout was equivalent to 35 and 32%, respcctively, of the total annual fallout, slightly higher than the percentage of the amount of precipitation of 39 cm, which was -26% of the annual rainfall.

The monthly ⁷Be depositional fluxes at Galveston varied between 1.2 and 74.5 dpm cm⁻² yr⁻¹ with a mean of 13.9 dpm cm⁻² yr¹ over the 38 months for which the depositional fluxes were measured. The corresponding monthly fluxes of ²¹⁰Pb varied between 0.072 and 4.75 dpm cm⁻² yr⁻¹ with a mean of 0.97 dpm cm^{-2} yr⁻¹. There was a spring maximum of ⁷Be or ²¹⁰Pb fallout in

Fig. 3. Frequency distribution of the depositional fluxes of ⁷Be in the 67 samples collected in Galveston, Texas.

only one of the three years (1991) for which we have a complete data set. The ⁷Be flux maximum during spring is generally attributed to a significant injection of stratospheric ⁷Be into the troposphere due to midlatitude (38°-51°N) folding of the tropopause which enhances stratospheric-tropospheric exchange [Young et al., 1970; Rangarajan and Gopalakrishnan, 1970; Dutkiewicz and Husain, 1979, 1985; Olsen et al., 1985; Todd et al., 1989]. The 210Pb flux maximum is likely related to the spring increase in the amount and frequency of precipitation, but in other places, it could be due to warming of frozen soils, or the drying out of saturated soils during spring or summer [Olsen et al., 1985; Schuler et al., 1991]. These differences in the depositional fluxes of ⁷Be and 210Pb during spring can also be due to the differences in the relative proportions of scavenging from stratiform and convective clouds. Frequent passage of strong fronts (associated with high rainfall) during all times of the year could also be a reason why we do not see a consistent ⁷Be fallout maximum during spring.

During the spring months March, April and May, the fractions of $7B$ e annual fluxes deposited in Galveston were 25, 29, and 37%, respectively, for the years 1989, 1990, and 1991. The corresponding values for ²¹⁰Pb are 19, 26, and 26% for the years 1989, 1990, and 1991, respectively. While we do not observe a markedly consistent seasonality in the ⁷Be and ²¹⁰Pb fluxes at Galveston, possibly due to more complicated atmospheric conditions, such as differences in the relative proportion of the stratiform and convective cloud scavenging, seasonal variations on the depositional fluxes have been reported for other continental and coastal stations. About 40-45% of the total annual deposition is reported to occur in three spring months (March, April, and May) at several places, such as Oak Ridge, Tennessee [Olsen et al., 1985], and Norfolk, Virginia [Todd et al., 1989]. Dibb [1989] and Canuel et al. [1990] found higher fluxes during the months May and June over the entire 1-year study period, and Schuler et al. [1991] reported consistent flux maxima during the summer months. Tsunogai et al. [1985] reported that deposition rates of 210Pb were higher in the winter at stations along the Japan Sea coasts, in contrast to the higher rates observed in the summer at stations along the Pacific coast.

Variations of the Specific Concentrations of 7Be

Specific concentrations of ⁷Be in all the samples collected are given in Tables 1 (Galveston) and 2 (College Station). More than 95% of both Galveston and College Station samples have activity concentrations of less than 250 dpm L⁻¹. The ⁷Be specific concentration in bulk deposition samples from Galveston ranged between 5.5 dpm L^{-1} and 1243 dpm L^{-1} , with a geometric mean of 130 dpm L⁻¹ (Table 1; since two of the 67 samples have an order of magnitude higher concentrations than the mean value, the geometric mean is reported). In College Station, 7Be concentrations varied between 15.9 and 298 dpm L^{-1} (Table 2), with a geometric mean of 112 dpm L⁻¹. This difference between sites, somewhat greater nuclide deposition at Galveston, may be related to its coastal meteorology and availability of condensation nuclei such as marine sulfate aerosols. The rain collectors were deployed in College Station for relatively longer time intervals, and therefore, the spread in radionuclide concentrations in the collected rainwater was considerably smaller. This range is comparable to the range of 29 to 191 dpm L^{-1} for the southeastern Virginia coast [Todd et al., 1989]. The higher end of the range in Galveston is due to very close interval sampling. Whenever there was a light shower (or drizzle) after a long dry spell, the samples contained very high specific concentrations of ⁷Be. During drizzles, smaller-size cloud drops likely have higher radionuclide concentrations. Growth of smaller water droplets (the smaller the size, the higher the affinity to aerosols) through coalescence will tend to keep this concentration constant, while further water moisture condensation would dilute the radionuclide concentration. Furthermore, smaller amounts of rainfall are always associated with higher evaporation rates of droplets owing to the lower humidity below cloud cover. This is true both during their in-cloud cycling and during their fall to the surface of the Earth. In such cases, these droplets will be enriched in radionuclide concentration [Wogman et al., 1968].

Relationship Between 7Be Specific Concentration and Precipitation

The ⁷Be specific concentration (in disintegrations per minute per liter) measured for Galveston is plotted against the amount of precipitation in Figure 6. The linear correlation between 7Be concentration and rainfall is poor $(r = 0.27, P < 0.05)$. However, if the parameters are plotted semi-logarithmically, an inverse dependency of the concentration on the amount of precipitation, x , is observed $(^{7}Be (dpm L^{-1}) = 334 - 288.8 \log(x); r = 0.70; n = 67, P$ < 0.001; Figure 6). For the rainwater samples from College Station, there is a significant correlation neither between ⁷Be concentration and rainfall ($r = 0.23$, $n = 21$, Figure 7) nor between ⁷Be flux and rainfall ($r = 0.072$, $n = 21$). However, other workers reported significant correlations between ⁷Be specific concentration and the amount of precipitation (Connecticut, $r = 0.74$, $P < 0.001$; Bermuda, $r = 0.66$, $\bar{P} < 0.05$ [Turekian et al., 1983]; Norfolk, Virginia, r =

Fig. 4. Frequency distribution of the depositional fluxes of ²¹⁰Pb in the 67 samples collected in Galveston, Texas.

-0.73 P < 0.001; Oak Ridge, Tennessee, r = -Q.79, P < 0.0O1 [Olsan et al., 1985]). It has been shown that the concentration of $7Be$ in rain decreasedby a factor of 3 during a storm event which deposited 6 cm of rain [Olsen et al., 1985]. Similarly, Dibb [1989] found that the specific concentration in samples taken serially during a single precipitation event decreased over time. Canuel et al. [1990] suggested that the concentration of $7Be$ in rain may be higher during drier months and periods characterized by short-duration precipitation events. A very weak correlation or lack thereof in the Texas data set suggests that the $7B$ e concentration is not mainly controlled by the amount of precipitation. A poor correlation has also been reported by Brown et al. [1989] and Todd et al. [1989], who similarly suggested that dilution is not the only process that controls the concentration of 7Be in rainwater.

Volume-Weighted Concentration of 7Be

Volume-weighted annual average ?Be concentrations were calculated as total amount of 7Be deposited in a ycar in the rain collector divided by total volumc of water in the rain collector, takcn either from the precipitation record (annual precipitation times surface area of the collector) or the actual volume of the water sample. At Galveston during 1989, 1990, and 1991, the average 7Be concentration was 87, 124, and 155 dpm L⁻¹, respectively. In 1990, this average 7Be concentration is idcntical at Galveston and College Station, whereas the 1991 year values are distinctly diffcrent (104 dpm L-l at College Station versus 155 dpm L-l at Galveston) even though the amounts of precipitation are comparable (Table 3). In Table 4, the volume-weighted $7Be$

Fig. 6. Specific concentrations of ⁷Be (disintegrations per minute per liter) versus amount of precipitation at Galveston, Texas.

Fig. 7. Specific concentrations of 7 Be (disintegrations per minute per liter) versus amount of precipitation at College Station, Texas.

concentrations are compared with values reported in the literature. The value at Galveston in 1991 is much higher than the values reported in many other places; for example, for the years 1983 and 1984, at Norfolk, Virginia (36'53'N), Todd et al. [1989] reported 90 and 100 dpm L⁻¹ respectively. Earlier results from Bombay (19°N, 33 dpm L^{-1} [Lal et al., 1979]) and Quillayute, Washington (49°N, 30 dpm L^{-1} [Crecelius, 1981]) are much lower than the values we are rcporting herc. and this differencc can be partly attributed to experimental artifacts in samplc collection (the water in the rain collectors was not acidified, as reported by Turekian et al. [1983], and thus a significant portion of $7\overline{B}e$ could have been adsorbed onto the container walls).

When these volume-weighted concentrations are normalized to a constant precipitation of 100 cm (by correcting the volumeweighted concentration by the ratios of 100 cm yr⁻¹ to the actual amount of rainfall), one obtains $84-127$ dpm L^{-1} for Galveston and 71-126 dpm L⁻¹ for College Station. This variation from year to year in the average ⁷Be concentration can be attributed to differences in frequencies of atmospheric turbulence and to sequences and altitude of washout events.

Variations of the Specific Concentration of 210Pb

The specific concentrations of $210Pb$ in bulk deposition samples from Galveston varied befween 2.03 and 218.1 dpm L-l (geometric mean of 10.8 dpm L⁻¹; Table 1). In College Station, they ranged between 2.13 and 25.7 dpm L^{-1} (geometric mean of 8.6 dpm L^{-1} ; Table 2). Most of the $210Pb$ specific concentrations fall within a narow range of 2-20 (only three samples have more than 100 dpm L-l) for Galveston, whereas in College Station, they fall within an even narrower range of 2-18 dpm L^{-1} (with the exception of one sample). These variations in the specific ²¹⁰Pb concentrations can be attributed to factors such as scavenging efficiency, the altitude from which the precipitation occurred, and the sources of the air masses (as the continental and oceanic air masses will have very different atmospheric inventories of ²¹⁰Pb [Turekian et al., 1977]). Since the deployment times of the rain collectors are longer for College Station samples, the specific concentration there is a result of integration of many rain events; thus the range of valucs beconres narrower than for the Galveston samples.

Relationship Between 210Pb Concentrations and Anounts of Precipitation

The ²¹⁰Pb specific concentrations in individual samples are plotted against amount of precipitation in Figure 8 for Galveston and in Figure 9 for College Station. There is only a very weak correlation between the specific concentration and amount of precipitation (for Galveston, $r = 0.29$ and $P < 0.05$; for College Station, $r = 0.31$ and $n = 21$). Benninger [1976, 1978] found a correlation of 0.67 (P < O.OOI) for the four years when precipitation samples were collected monthly. A relatively weak positive or negative correlation between ²¹⁰Pb concentration and amount of precipitation has been observed by other workers in Oak Ridge. Tennessee $(r = 0.65, P < 0.001$ [Olsen et al., 1985]), Norfolk, Virginia ($r = -0.49$, $P < 0.05$ [Todd et al., 1989]), and Texel, De Bilt, and Groningen, in the Netherlands ($r = 0.74-0.94$, $P < 0.001$ [Zuo, 1992]). Nevissi [1985] observed a positive correlation between monthly rainfall and the ²¹⁰Pb deposition at Seattle, Washington; however, for the same amount of rainfall the ²¹⁰Pb deposition rate also varied from year to year. Hussain et al. [1990] found no significant correlation between monthly atmospheric 210Pb fallout and rainfall. This lack of correlation can be attributed to several factors that control the depositional fluxes of $210Pb$ such as sources, pathways, altitude of the clouds, intensity of fronts,

GVS is Galveston, and CLL is College Station. NM means not measured for the whole year; samples collected from June 1989 onward.

* One rain sample from College Station was lost during collection between March 22 and April 22 (14.5 cm rain).

+ Total concentrations collected in container with 2800 cm^2 area.

These are the values from rainwater samples collected during 333 days in 1991 (except March 22 to April 22).

 $$$ The fluxes (dpm cm⁻² yr⁻¹) and specific concentrations (dpm L⁻¹) were calculated for 333 days and then prorated for 365 days.

T Concentration (dpm L⁻¹) = annual flux (dpm cm⁻² yr⁻¹)* 1000 / annual precipitation in cm = total amount (disintegration per minute)/total volume (L).

TABLE 4. Comparison of Volume-Weighted Concentrations of ⁷Be and ²¹⁰Pb Measured in Galveston and College Station, Texas in 1989 through 1991 With Those Measured From Other
Regions of the World after 1980

Volume-weighted average concentrations were calculated as in Table 3. $*$ Based on 8 months of precipitation data

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Fig. 8. Specific concentrations of 210Pb (disintegrations per minute per liter) versus amount of precipitation at Galveston, Texas.

relative humidity, and the transit time of aerosols that contain 210Pb from their source to the sink in rain collectors.

Volume-Weighted Concentration of 210Pb

The volume-weighted ²¹⁰Pb concentrations for Galveston and College Station, calculated as the total amount of 210Pb deposited divided by total rain volume, are given in Table 3. During the years 1989-1991, the concentrations were 6.75, 6.95, and 11.4 dpm L^{-1} , respectively, for Galveston and 7.94 dpm L⁻¹ (1990) and 9.23 dpm L-1 (1991) for College Station. When the 210Pb specific concentrations were normalized to a constant amount of precipitation, by taking the product of average concentration and the ratio of 100 cm yr^{-1} to the actual amount of rainfall, a relatively constant value resulted for both Galveston $(7.10 \text{ dpm } L^{-1})$ and College Station $(7.21 \text{ dpm } L^{-1})$. Thus it seems that the annual atmospheric depositional flux of ²¹⁰Pb is controlled by the amount of precipitation for the years we have studied, as may well be the case in other places.

In Table 4, our volume-weighted 210Pb concentrations are compared with values reported in literature from other places. The value at Galveston in 1991 is much higher than the values reported in many other places; for example, for the years 1983 and 1984, at Norfolk, Virginia (36°53'N) Todd et al. [1989] reported only 5.9 and 6.6 dpm L⁻¹ respectively. For New Haven, Connecticut (41°N) and Bermuda (33°N) Turekian et al.. [1983] reported 8.0 dpm L-1 and 4.1 dpm L^{-1} , respectively.

Variations of Specific Concentrations and Fluxes 7Be of and 210Pb Sampled During Individual Rain Events Within One Month

In 1991, some of the Galveston rainwater samples were collected right after individual rain events. Information that can be obtained from these samples will be useful in studying the variations of specific concentrations and fluxes between one rainout event and another, as well in determining the variability for a single month from year to year. For example, depositional fluxes in September 10-13, 1990 (all 4 days were rainy days) can be compared to the depositional flux during September 5-7, 1991 (all 3 days were rainy). The depositional flux of $7Be$ differed by a factor of almost 2 between the two collection periods (27.2 and 17.9 dpm cm⁻² yr⁻¹ for the first and second periods, respectively) whereas the 210Pb fluxes (1.59 and 1.42 dpm cm⁻² yr⁻¹) remained essentially the same.

Another example is the four rain events that were collected continuously between September 2 and 8, 1991. If all these rain events were derived from the same water vapor, then one would expect a dilution of the concentration with time. However, the specific concentrations of ⁷Be decreased from 67.4 dpm L^{-1} in the first rain to 31.7 dpm L^{-1} in the third rain and then increased again to 47.7 dpm L^{-1} in the fourth rain event. Similarly, the specific concentration of ²¹⁰Pb decreased from 3.16 dpm L⁻¹ in the first rain to 2.52 dpm L⁻¹ in the third rain and again increased to 5.73 dpm L⁻ ¹ in the fourth rain event. The nuclide fluxes varied by a factor of 3 for 210Pb and a factor of 5 for $7Be$. The $7Be/210Pb$ activity ratios varied by a factor of 2.6 within these 8 days. Thus it appears that a significant amount of 210Pb-rich aerosols in the fourth rain is derived from entrained continental air. A record of the wind directions during these 8 days indicates that the fourth rain event was brought by southern winds, different from the others.

Fig. 9. Specific concentrations of 210Pb (disintegrations per minute per liter) versus amount of precipitation at College Station, Texas.

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Relationship Between Wind Direction and Depositional Fluxes of $7Be$ and $210Pb$

The depositional fluxes of $7Be$ and $210Pb$ are plotted against wind direction in Figures 10 and 11, respectively. The common wind direction is from the southeast. On only one day (January 15, 1991) did the wind originate from the northwest, and this wind appears to have brought significant amounts of 2l0Pb and ?Be, resulting in the highest ⁷Be and ²¹⁰Pb fluxes. These higher levels could be due to longer transit time of aerosols which brought large amounts of ⁷Be and ²¹⁰Pb nuclides or stronger interactions between higher-altitude jet stream and near-surface air masses. However, there is no significant relationship between the depositional fluxes of these nuclides and wind direction. AIso. there is no correlation between wind speed and depositional fluxes of $7Be$ or $210Pb$ at any of the stations. Thus it appears that strong fronts passing through Texas entrain variable amounts of oceanic and continent-derived acrosols.

Fraction of the Annual Depositional Fuxes of $7Be$ and $210Pb$ From Pulse Rain Events

Even though the rainy days are spread over the whole year, during a few days, the rainfall was heavy due to thunderstoms, hurricanes, or tomadoes. The contribution of ⁷Be and ²¹⁰Pb to the annual bulk depositional fluxes during these heavy rainout events is not very well known. Since the input of these nuclides during heavy rains into the coastal water is useful for studying the particle dynamics in coastal and bay waters [Olsen et al., 1989; Baskaran and Santschi, 1993], it is important to know the relative contribution during these events when using the nuclides as tracers for particle cycling.

In Galveston, there were 82, 88, and 119 rainy days during 1989, 1990, and 1991, respectively (Table 5). Of these, 4, 6, and 5 rainy days had greater than 5 cm rainfall during the years 1989, 1990, and 1991, respectively. Even though the fractional amount of precipitation and the depositional fluxes of radionuclides are comparable in these pulse events, a higher amount of depositional flux into the coastal waters can be used as a tracer to investigate the pollutant transport and particle cycling. For example, in Galveston Bay, with a mean depth of 2 m, one of these individual rainout events (January $6-8$, 1992, Table 1) typically would have resulted in total concentrations (dissolved and particulate) of 2.2 dpm L^{-1} for

Fig. 10. The depositional flux of $7Be$ at Galveston, Texas is plotted against wind direction.

Fig. 11. The depositional flux of ²¹⁰Pb at Galveston, Texas is plotted against wind direction.

7Be and 0.15 dpm L -1 for 210Pb in the bay water. These concentration levels are easily measurable, and thus the fate of particles as well as the dissolved nuclides can be studied [Baskaran and Santschi, 1993].

In 1990, six heavy rainfalls with more than 5 cm precipitation deposited 22% of the total annual ⁷Be flux and 21% of the annual flux of $210Pb$ (Table 5). During 1991, the ⁷Be and $210Pb$ deposited during these 5 days of heavy rainfall amounted to 26 and 20%, respectively, of the annual flux (Table 5). The fraction of the depositional flux of 7Be and 2l0Pb for 1989 was not calculated as one of the four rain samples with more than 5 cm rain was lost. It appears that 4-6 heavy rain events per year (typically 4-6% of the total number of rainy days) account for about $20-30\%$ of the total annual deposition rate of $7Be$ and $210Pb$. This is likely to be true for other atmospherically delivered stable elements as well.

Dry Depositional Fluxes of ⁷Be and ²¹⁰Pb

The time during which the dry depositional flux study was carricd out was limited to about a weck cach in October and November 1990 and to 2 wccks in October 1991. The dry depositional fluxes of radionuclides are presented in Table 6. These dry depositional fluxes were obtained by exposing the rain collector's lid (surface area of 2800 cm^2 , depth of 3.5 cm) for 7-14 days with 1.5 cm distilled water. Such a method was advisable, sincc we were primarily interested in estimating the depositional fluxes of ⁷Be and ²¹⁰Pb into coastal waters [Dasch, 1985]. Another method of determining the dry fallout is from the measurement of bulk (wet and dry) fallout of these nuclides and extrapolation to zero precipitation. Soil profiles have been mentioned too, as they provide bulk atmospheric depositional fluxes over the mean life of the radionuclide [Graustein and Turekian, 1986]. However, it will be difficult to get information on the annual dry fallout for $210Pb$ and ⁷Be from such studies.

The dry depositional flux for ⁷Be varied between 0.249 and 0.394 dpm cm⁻² yr⁻¹. This dry depositional flux was 3.0 to 7.4% of. the bulk depositional flux during October 1990, but it amounted to about 877o during October 1991. This high value is likely due to the fact that the bulk deposition rate during that period was low, as there was no rainfall for a period of about a month (September 25 to October 27), thus giving rise to high concentrations of 7Be and

Collection Interval	Total amount of precipitation, cm.	Number of Days With > 5 cm Rain	Be Flux	Percentage of Annual Percentage of Annual 210_{Pb} Flux
1989	103	4	IC	IС
1990	97	6	22.2(2.66)	20.5(0.137)
1991	150	5	25.8(5.99)	20.0(0.342)

TABLE 5. Percentage of Bulk ⁷Be and ²¹⁰Pb Depositional Fluxes During Heavy Rain at Galveston, Texas, During 1990 and 1991

Numbers in parentheses denote depositional fluxes (disintegrations cm^{-2} yr⁻¹) during the heavy rain events. IC means ⁷Be and 210 Pb annual flux for the year 1989 is incomplete and hence was not calculated.

210Pb in surface air aerosols. Todd et al. [1989] observed a weak positive correlation between dry ⁷Be depositional flux and the amount of monthly precipitation and attributed this to higher relative humidity during dry fallout periods. However, our present data do not allow us to address this question to any great extent. It is sufficient to say that humidity is high in Galveston year around. Furthermore, Santschi et al. [1988] reported higher dry fallout values during humid, foggy days for Chernobyl-derived radionuclides. Also, traditionally, the linear relationship between amount of precipitation and bulk depositional flux can enable one to estimate the dry fallout at zero precipitation. Since we do not find any relationship between amount of precipitation and depositional fluxes of ⁷Be and ²¹⁰Pb it is not possible to estimate dry

depositional fluxes from such an approach.
The dry depositional flux for ²¹⁰Pb varied between 0.032 and 0.092 dpm cm⁻² yr¹ during 7- to 14-day periods, which appears to be a higher fraction of the total fallout than was measured for ⁷Be for the same period (6 and 22% in October/November 1990 and 41% in October 1991; Table 6). This higher fraction of dry fallout for $210Pb$ relative to ⁷Be could likely be caused by $210Pb$ leaching from fine resuspended dust from ground in the acidified dry fallout collector. Alternatively, dry deposition could be more important for 210Pb than for ⁷Be. A similar observation has been reported by Olsen et al. [1985], based on the relatively weaker correlation between total monthly 210Pb deposition and the amount of precipitation. The average fraction of dry deposition flux of ²¹⁰Pb, 14% in 1990, appears to be closer to the value reported by other workers. For example, Brown et al. [1989] measured dry 7Be depositional fluxes and estimated that less than 10% of the bulk fallout is dry fallout. The $210Pb$ dry depositional flux in 1991 is higher than in 1990, as was also the case for 7Be.

Annual Bulk Depositional Fluxes of 7Be and 210Pb

The annual bulk depositional fluxes of 7Be and 210Pb at Galveston and College Station are given in Table 3. During the years 1989 to 1991, depositional fluxes of ⁷Be in Galveston varied between 8.9 and 23.2 dpm cm⁻² yr⁻¹, with an average of 14.7 dpm $cm⁻²$ yr⁻¹. The highest radionuclide fallout in 1991 corresponds to a maximum annual rainfall. Even though the annual depositional flux of $7Be$ varied by a factor of 2.6, the $7Be$ flux normalized to 100 cm yr^{-1} rainfall varies only by a factor of 1.8. The College Station annual depositional values of ⁷Be in 1990 and 1991 were 12.7 and 15.0 dpm cm⁻² yr^{-1} . This range of values can be compared to the values reported from other places, including New Haven,
Connecticut (22.7 dpm cm⁻² yr⁻¹ [*Turekian et al.*, 1983]), Bermuda (17.1 dpm cm⁻² yr¹ [Turekian et al., 1983]), Oak Ridge, Tennessee (12.0 dpm cm⁻² yr¹ [Olsen et al., 1985]), Norfolk, Virginia (12.0-12.9 dpm cm⁻² yr⁻¹ [Todd et al., 1989]), Chesapeake Bay, Maryland (13.6 dpm dpm cm⁻² yr⁻¹ [Dibb, 1989]), and Dubendorf, Switzerland (16.0-16.5 dpm cm⁻² yr⁻¹ [Schuler et al., 1991]).

The annual ²¹⁰Pb depositional fluxes in Galveston range between 0.67 and 1.71 dpm cm⁻² yr⁻¹, with a mean of 1.03 dpm cm⁻ ² yr¹. Depositional fluxes in College Station during 1990 were 0.78 dpm cm⁻² yr¹ and in 1991 1.33 dpm cm⁻² yr¹. The ²¹⁰Pb fluxes normalized to 100 cm yr⁻¹ rainfall for College Station are, however, about the same in both years, 0.80 dpm cm⁻² yr^{-1} in 1990 and 0.77 dpm cm⁻² yr⁻¹ in 1991. The corresponding values for Galveston range between 0.68 and 1.14 dpm cm⁻² yr⁻¹, with a mean of 0.84 dpm cm⁻² yr⁻¹ for 100 cm yr⁻¹ rainfall. These values can be compared to the values reported from other regions. Benninger [1976, 1978] reported for the 4-year period between 1973 and 1977, annual ²¹⁰Pb fluxes in New Haven, Connecticut, of 0.86 to 1.04

Collection Interval	Time for Which Sample Was Collected days	7 Be Flux dpm cm ⁻² yr ⁻¹	$\frac{210 \text{pb} \text{ Flux}}{4 \text{pm cm}^{-2} \text{ yr}^{-1}}$	$7_{Be}/210_{Pb}$ Activity Ratio
Oct. 26 to Nov. 2, 1990	7	0.249(8.40)	0.031(0.486)	8.11(17.3)
Nov. 12 to Nov. 19, 1990	7	0.394(5.35)	0.092(0.420)	4.28(12.7)
Oct. 7 to Oct. 21, 1991	14	0.277(0.318)	0.059(0.144)	4.72(2.21)

 $\mathbf{R} \mathbf{I} \mathbf{E}$ 6. Dry Depositional Fluxes of 7 Be and 210 Ph Measured in Galveston. Te

Numbers in parentheses denote bulk depositional fluxes (or activity ratio) during this sampling period.

dpm cm⁻² yr⁻¹. Nevissi [1985] measured the monthly depositional fluxes of ²¹⁰Pb in Seattle, Washington, over a period of 7 years, and the annual depositional flux varied between 0.18 and 0.81 dpm cm⁻² yr⁻¹ Average fluxes were 0.44 dpm cm⁻² yr⁻¹ for Seattle, Washington [Nevissi, 1985], 1.2 dpm cm⁻² yr⁻¹ for New Haven,
Connecticut [*Turekian et al.*, 1983], 0.69 dpm cm⁻² yr⁻¹ for Bermuda [Turekian et al., 1983], 1.04 dpm cm-2 yr-1 for Oak Ridge, Tennessee [Olsen et al., 1985], 0.79-0.85 dpm cm⁻² yr¹ for Norfolk, Virginia [Todd et al., 1989], 0.83-0.91 dpm cm⁻² yr⁻¹ for Dubendorf and Geneva, Switzerland [Dominik et al., 1987; Schuler et al., 1991], and 0.32-0.43 dpm cm⁻² yr⁻¹ for Texel, De Bilt, and Groningen, the Netherlands [Zuo, 1992]. Moore and Poet [1976] estimated the local ^{210}Pb flux from a correlation of ^{210}Pb concentration profiles versus latitude. For the westem central United States, they reported an average value of 0.8 dpm cm⁻² yr¹.

Our measured 210Pb fallout of 0.67-1.71 dpm cm⁻² yr⁻¹ corresponds to a $222Rn$ emanation rate from continental soils of 0.36-0.91 atoms cm⁻² s⁻¹, with a mean of 0.55 atoms cm⁻² s⁻¹ for Galveston and College Station. This is in reasonable agreement with 0.40 atoms cm^{-2} s⁻¹ for the central and coastal Texas and 0.7 atoms cm^{-2} s⁻¹ for Texas plains [Wilkening et al., 1975]. However, the 210Pb containing aerosols may have been derived from continental sites, which were farther away and had different rndon exhalation rates.

Can 7Be and 210Pb Depositional Fluxes be Used as Two Independent Atmospheric Tracers?

Since their modes of production are different, $7Be$ and $210Pb$ have two distinct sources; howcver, they are removed from the atrnosphere by very similar mechanisms. If the depositional fluxes of these two nuclidcs were found to be linearly related to each other, then they could not be used as two independent atmospheric tracers for continental and occanic air.

The ⁷Be/²¹⁰Pb ratios in all the rainwater samples, from both Galveston and College Station, are given in Tables L and 2. The $7Be/210Pb$ activity ratio histograms are plotted in Figures 12 (Galveston) and 13 (College Station). The ratios varied over an order of magnitude, between 2.21 and 32.6. The annual $7Be/210Pb$ ratios varied between 11.3 and 17.8, with a mean of 14.4 (Table 3). Over the 7 consecutive days during which rainfall was collected (four rains, September 2-8, 1991), this ratio varied between 8.33 and 21.4, Thus, when it was raining evcry day for several days, thc ratio gradually decreased and this decrease can be attributed to the

Fig. 13. Frequency distribution of $7Be/210Pb$ activity ratio in College Station, Texas samples.

variable mixing of continental aerosols that are enriched in 210Pb compared to oceanic aerosols. For a given latitude, however, aerosols are cxpected to have morc or less the same 7Be concentration.

The bulk depositional fluxes of $7Be$ are plotted against $210Pb$ depositionat fluxcs in Figure 14 for Galveston and in Figure 15 for College Station. The strong correlation between the two nuclides, for both Galveston ($r = 0.94$, significant at $P < 0.001$) and College Station ($r = 0.91$, significant at $P < 0.001$), indicates dependence rather than independence of these two tracers.

There is another reason for exploring the possibility of using $7Be$ and 2l0Pb nuclides as two independent tracers. In the years 1989 and 1990, with about the same rainfall, the depositional fluxes of 210Pb remained the same at Galveston (0.70 and 0.67 dpm cm⁻² yr 1, respectively), while the annual $7Be$ flux in 1990 was 35% higher than that of 1989. However, the good correlation between the two (as seen in Figures 14 and 15) suggests that they were generally removed by the same atmospheric process(cs) with only minor variations in the annual flux ratios. Theoretically, one would expect a good correlation between the two mainly for continental stations,

Fig. 12. Frequency distribution of $7Be/210Pb$ activity ratio in Galveston, Texas samples.

Fig. 14. Bulk depositional fluxes of $7Be$ are plotted against $210Pb$ fluxes for Galveston, Texas.

Fig. 15. Bulk depositional fluxes of ⁷Be are plotted against ²¹⁰Pb fluxes for Collcgc Station, Tcxas.

such as Dubendorf, Switzerland, since most of the ^{"210}Pb-tagged" aerosols are land derived and are primarily removed by precipitation. In a purely oceanic station, like Bcrmuda, parts of the air mass are likely derived from continents and so are enriched in 210Pb, whereas other parts are ocean derived, and so are depleted in 210Pb. Thus we do not expect any correlation between the two nuclides in an oceanic station even if they are removed by the same kinds of atmospheric processcs. In coastal stations, we might expect a mixhrre of the two end-member cases and thus a relatively weak correlation.

The depositional flux of ²¹⁰Pb is plotted against depositional flux of ⁷Be in Figure 16 for continental stations Dubendorf, Switzerland ($r = 0.90$, $P < 0.001$) and Oak Ridge, Tennessee, ($r =$ 0.86, $P < 0.001$), coastal stations Galveston, Texas ($r = 0.94$, $P <$ 0.001], New Haven, Connecticut $(r = 0.59, P < 0.05)$, and Norfolk, Virginia ($r = 0.77$, $P < 0.001$), and oceanic station Bermuda ($r =$ 0.43 , $n = 8$). The least squares best fit line for the oceanic station is distinctly different from all stations, even from most coastal stations. The regression line for New Haven, Connecticut, appears to be that expected for a coastal station, Galveston, Texas, and Norfolk, Virginia ($r = 0.65$, $P < 0.001$) appear to behave more like continental stations in terms of the relationship between ⁷Be and 210Pb depositional fluxes (data taken from Olsen et al. [1985], Todd et al. [1989] and Schuler et al. [1991]). It appears that in most of the continental and coastal stations, 7Be and ²¹⁰Pb cannot be used as two independent atrnospheric tracers. Only in oceanic and a few coastal stations do $7\hat{B}e$ and $210Pb$ fluxes seem to vary independently, and in such places these nuclides can be used as independent air mass tracers.

Inplications of theVariations of Annual 21OPb Depositional Fluxes for the ²¹⁰Pb-Based Sediment Chronologies in Lakes and Coastal **Waters**

The ²¹⁰Pb sediment chronology in aquatic systems is based on the assumption that the annual flux of ²¹⁰Pb to the sediments remains constant [*Robbins*, 1978; *Robbins and Edgington*, 1975]. When the residence time of ²¹⁰Pb in the water column is much shorter than the half-life of $210Pb$, that is, 22 years, then any variability in the annual atmospheric delivery of ²¹⁰Pb will cause a similar variability in thc annual delivery to the sediments. For example, in coastal and estuarine waters, the residence time of dissolved ²¹⁰Pb is of the order of a month or less [Baskaran and Santschi, 1993, and references therein]. Under such circumstances, all of the atmospheric ²¹⁰Pb will be delivered to the sediments, and therefore the annual variability of the depositional flux of ²¹⁰Pb will result in variable flux to the sediment-water interface. Annual depositionat fluxes of 2l0Pb in Galveston varied between 0.67 and $1.\overline{7}1$ dpm cm⁻² yr⁻¹ over the 3 years this study was carried out. Similar depositional flux variations havc been previously reported by Neviosi [1985] for Seattle, Washington (a 4.4-fold range in the 210Pb annual flux during the periods 1973 and 1980). Thus it is likely that a vertical profile of $210Pb$ concentrations in coastal (as well as lake) sediments will be affected not only by sediment mixing (physical and biological) but also by thc variabiliry of thc

Fig. 16. Bulk depositional fluxes of ⁷Be are plotted against ²¹⁰Pb fluxes for data collected from continental, coastal, and oceanic stations.

atmospheric depositional flux of 2l0Pb. In open ocean waters, on the other hand. such as in the North Atlantic and North Pacific oceans, the dissolved ^{210}Pb residence time is of the order of 10-100 years [Craig et al., 1973; Somayajulu and Craig, 1976]; thus the annual variability of the atmospheric 2l0Pb flux will be smoothed out, and the variations of the annual depositional flux of $210Pb$ to the sediments will be much more smaller when sediment delivery rates axe constant.

CONCLUSIONS

Simultaneous measurements of bulk atmospheric depositional fluxes of $7Be$ and $210Pb$ were made at a coastal station (Galveston, Texas) and an inland site (College Station, Texas), for about 3 years. From this extensive database, we draw the following conclusions.

1. The annual depositional flux of 7Be, measured at Galveston during 1989 to 1991, does not remain constant but varies between 8.9 and 23.2 dpm cm⁻² yr⁻¹, with a mean of 14.7 dpm cm⁻² yr⁻¹. The flux varies with the amount of precipitation. The precipitationnormalized ?Be flux is not constant either but varies within a factor of 1.8. The ²¹⁰Pb depositional fluxes range between 0.67 and 1.71 dpm cm⁻² yr¹, with a mean of 1.03 dpm cm⁻² yr¹. The highest values are in the year with the maximum rainfall. The precipitationnormalized ²¹⁰Pb fluxes are constant for College Station, while they vary by a factor of 1.7 for Galveston,

2. The bulk depositional fluxes of $7Be$ and $210Pb$ do not show any consistent seasonal trends. In one year there was a spring maximum, and in another year there were flux maxima in both winter and summer. However, the annual depositional fluxes are most likely controlled by the amount of precipitation during that vear.

3. It appears that 4-6 heavy rain events in a year (typically 4-6% of the total number of rainy days) account for about 20-30% of the total annual deposition rate of 7Be and 210p6. This is likely to be true for other atmospherically deliverrd stable elements also.

4. The dry depositional fluxes were generally less than 10% for /Be. However, in the months when there was scanty rain. dry depositional fluxes of ⁷Be and ²¹⁰Pb became a major portion of the bulk depositional fluxes. The fraction of dry depositional flux to total flux appears to be higher for $210Pb$ than ⁷Be. This higher $210Pb$ dry depositional flux relative to ⁷Be could also be due to leaching of resuspended dust by acid solution in the dry fallout collector,

5. On the basis of comparison with other measurements reported in thc literature, it appears that in most of the continental and coastal stations, ⁷Be and ^{210p}b fallout pattern is tightly correlated, as is observed in Galveston and College Station, and thus the two nuclides cannot be used as two independent atmospheric air mass tracers; only in oceanic and a few coastal stations do $7Be$ and $210Pb$ behave as two independent atmospheric tracers.

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