Magnetic Moment of the Electron

P. Kusch

I must say, and with considerable regret, that I am not a theoretical physicist. A penetrating analysis of the part that the discovery and measurement of the anomalous magnetic moment of the electron has played in the development of certain aspects of contemporary theoretical physics must be left to the group of men who have in recent years devised the theoretical structure of quantum electrodynamics. My role has been that of an experimental physicist who, by observation and measurement of the properties and operation of the physical world, supplies the data that may lead to the formulation of conceptual structures. The consistency of the consequences of a conceptual structure with the data of physical experiment determines the validity of that structure as a description of the physical universe.

Our early predecessors observed nature as it was displayed to them. As knowledge of the world increased, however, it was not sufficient to observe only the most apparent aspects of nature to discover its more subtle properties; rather, it was necessary to interrogate nature and often to compel nature, by various devices, to yield an answer concerning its functioning. It is precisely the role of the experimental physicist to arrange devices and procedures that will provide information that will enable us to make quantitative statements concerning the properties and behavior of nature. It is in this spirit that I will discuss here my participation in a sequence of earlier experiments that made possible the precision determination of the magnetic moment of the electron. I will then discuss the experiments themselves—the experiments that have yielded our present knowledge of the magnetic properties of the electron.

Atomic and Molecular Beams

Research with atomic and molecular beams has had a long and fruitful record in the history of the growth of our present knowledge of matter. The experiments that I shall discuss are some in which the method of atomic and molecular beams is used essentially as a spectroscopic device for the observation of spectral lines in the range of frequencies within which power may be generated by electronic means. The general principles of radio-frequency spectroscopy by the method of molecular beams were first described by Rabi and a group of his coworkers (1) of which I was fortunate to be a member. It is here sufficient to say that a transition between energy levels may be observed through the circumstance that the magnetic moment of an atom or molecule may be changed in a transition. The method is characterized by a very high-potential resolution; in many observations of the frequency of a line, an accuracy of better than 1 part in 1 million has been achieved. It is of particular value as a tool in the investigation of the details of interactions within atoms and molecules because small interactions appear as first-order effects rather than as small superpositions on the relatively enormous energies that characterize optical spectra.

Electron Properties

The fact that the electron has a spin of $\frac{1}{2}$ and a magnetic moment at least approximately equal to 1 Bohr magneton has long been recognized. Uhlenbeck and Goudsmit first postulated these properties of the electron to explain the fine structure in atomic spectra and what has been called the anomalous Zeeman effect (2). An enormous body of evidence has given ever-increasing support to these postulates. The relativistic Dirac theory of the electron assumed a particle that was endowed with the properties of mass and charge. The spin and magnetic moment postulated by Uhlenbeck and Goudsmit were then found to be a consequence of the relativistic invariance of the Dirac equation. Indeed, one of the great triumphs of the Dirac electron theory was the prediction of these postulated electron properties. The spin and moment of the electron were thus removed from the realm of ad hoc assumptions, justified by experimental evidence, to the realm of an integral part of quantum theory. The Dirac electron theory did not, however, consider the interaction of the quantized electromagnetic field with the electron.

I shall talk of the measurement of the g value rather than the magnetic moment of the electron. The g value is, as usual, the negative ratio of the magnetic moment in terms of the Bohr magneton µ0 and the angular momentum in units of $h/2\pi$. Since, in all cases here under discussion, the angular momentum of the system is known, the moment can immediately be obtained from the g value. The most elementary of the g values, g_{L} , is that associated with the orbital motion of the electron. Its value is 1 within small and calculable corrections. The electron also has a magnetic moment by virtue of its angular momentum about a spin axis. The g value associated with the spin, g_{S} , is the quantity here under investigation; a value of 2 was obtained for it in the Dirac electron theory. Now the electrons in an atom have both spin and orbital angular momentum. To the total electronic angular momentum J, we assign the g value g_J . The atom contains a nucleus that may have a nuclear angular momentum and hence a nuclear magnetic moment. The nuclear g value, g_N , is designated as g_P in the special case when

SCIENCE

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the nucleus is a proton. To the total angular momentum of the atom we assign the g value g_F .

Early Measurements

The earliest measurements by the molecular-beams magnetic resonance method were undertaken by a group, of which I was a member, that worked in Rabi's laboratory and under his direction (1). The measurements consisted of the determination of nuclear g values by the observation in a molecule of the nuclear resonance frequency in a classically determined magnetic field. Even in the great national laboratories dedicated to the maintenance of physical standards, a precision of only about 1 part in 40,000 has been achieved in the measurement of a field (3). In a well-equipped laboratory that lacks the equipment and tradition of meticulous intercomparison of electric standards, a precision of perhaps ¹/₄ percent may be achieved in the determination of the magnitude of a field. While ratios of nuclear g values may be found without an explicit knowledge of the field, the accuracy of the determination of a nuclear moment in terms of the Bohr or nuclear magneton is limited by the uncertainty in the measurement of a field as well as by the uncertainties in a prior measurement of the Bohr or nuclear magneton. Thus, the desirability of a direct measurement of a nuclear g value in terms of the Bohr magneton is apparent.

The molecular-beam magnetic resonance method was originally applied to the determination of the nuclear g values in molecules that did not have a net electronic orbital or spin angular momentum. It is, however, possible to apply the same experimental techniques to an investigation of the hyperfine structure of atoms. If we observe transitions between the various F levels at zero or very low magnetic field, the hyperfine structure separation may readily be found. At a higher magnetic field, the observation of the frequency of transition between magnetic sublevels yields again the zero-field hyperfine structure splitting, the quantity $g_J \mu_0 H/h$, and the quantity g_N/g_J , although the latter quantity can be found with only limited precision. The group of which I was a member at Columbia made the first such studies on the commonly occurring isotopes of the alkali atoms and determined the magnetic hyperfine structure interaction constants of the alkali atoms (4). Extensive subsequent work in the observation of atomic hyperfine structure has, of course, been done in many laboratories with results of great interest in the study of higher order moments in nuclei and of the properties of radioactive nuclei. The alkali atoms were particularly adaptable to the

original experimental work first because the beams of the atoms may readily be produced and detected and second because they occur in ${}^{2}S_{1/2}$ states, almost wholly free of perturbation by other states.

The possibility of measuring the moment of a nucleus in terms of the Bohr magneton is a consequence of the possibility of observing both nuclear resonance in molecules with a frequency $g_N \mu_0 H/h$ and transitions among the magnetic components of hyperfine structure levels for which the dependence of frequency on field is of the order of $g_J \mu_0 H/h$. Millman and I then addressed ourselves to the problem of measuring the moment of the proton in terms of the electron spin moment (5). The experimental problems were considerable and arose from three factors.

The first of these was related to the fact that the effective moment of a molecule of zero electronic angular momentum is of the order of a nuclear magneton while that of an atom is of the order of a Bohr magneton. Deflecting fields that allow the observation of a change in trajectory of a molecule in which a hydrogen nucleus has undergone a transition will deflect an atom through unmanageably large excursions if the field is arbitrary. However, all atoms in which magnetic hyperfine structure occurs and in which the spin of the nucleus is greater than $\frac{1}{2}$ have, in certain states, zero magnetic moments at definite values of the magnetic field, which may be very high. Atoms in such states may thus traverse a carefully adjusted inhomogeneous field without catastrophic deflections; and a transition from such a state may at once be detected because the terminal state is generally characterized by a large magnetic moment. Thus, it is possible to choose the deflecting field in such a way that a change in the spin orientation of a nucleus in a molecule and a transition among the magnetic levels of the hyperfine structure may both be detected.

The second of our experimental problems was related to the production of a beam of molecules that contained hydrogen and, simultaneously, an alkali atom, which was requisite for the detection of beams by the techniques then available to us. Since atomic lines were to be observed at the same field as nuclear resonance lines, the simultaneous production of a beam of alkali atoms was necessary. We used beams of sodium and potassium hydroxides evaporated from silver ovens and noted that, at the temperatures required to generate a beam of the hydroxide, the reaction between an alkali halide and metallic calcium proceeded at such a rate that a convenient beam of atoms appeared.

The third experimental problem was associated with the need of applying to the same circuit, in succession, two frequencies that differed by as much as a factor of 70 and with the prewar difficulty of generating sufficient power at high radio frequencies.

Extensive intercomparison of the frequencies of the resonance line of the proton and of lines in the hyperfine structure spectra of sodium, rubidium, and cesium-for each of which a prior determination of the interaction constants had been made-led to a determination of the ratio of the proton moment and the spin magnetic moment of the electron in the calibrating atoms, which we, of course, assumed to be the Bohr magneton. The magnetic moment of the proton in terms of the nuclear magneton that was found on the basis of the assumption that the spin moment of the electron was indeed the Bohr magneton differed from the moment as determined from the measurement of a frequency in a classically determined field by about 0.1 percent. When, at a much later date, it was found that the spin magnetic moment of the electron deviates from the Bohr magneton by the order of 0.1 percent, the direction of the deviation in the older experiment was examined. It is perhaps a good commentary on the hazards of experimental physics that no significant effect had escaped us but that the error in the mutual inductance which we had used in the calibration of the magnetic field was of the order of 0.2 percent rather than 0.1 percent.

After World War II, Nafe and Nelson, working with Rabi, made the first of the measurements of the hyperfine structure splitting of hydrogen in the ground state (6). Now the hyperfine structure of hydrogen may be calculated explicitly in terms of the magnetic moment of the proton, the spin magnetic moment of the electron, and the electronic wave function at the nucleus. However, a discrepancy of about $\frac{1}{4}$ percent was noted between the observed and predicted magnitude of the hyperfine structure splitting when the value of the proton moment found by Millman and me was used. The assumption that the spin moment of the electron is 1 Bohr magneton enters into the calculation twice-first as an intrinsic property of the electron and second in the calculation of the proton moment from the observed ratio of the proton moment and the spin moment of the electron. The discrepancy led Breit to suggest (7) that the electron may possess an intrinsic magnetic moment greater than μ_0 by the order of $\alpha \mu_0$, where α is the usual fine structure constant.

Anomalous Magnetic Moment

The question of the existence of an anomalous magnetic moment was then investigated in detail by Foley and me

(8). In this inquiry, as in all others conducted in the atomic and molecular beams laboratory at Columbia University, we profited by Rabi's advice. The procedure that we employed made use of the fact that the g_J value associated with a state is a linear combination of the electronic orbital and spin g values, g_L and g_8 and that this combination is different for different states. That is, there is a contribution to the total electronic magnetic moment of an atom both from the orbital motion of the electrons and from the spin of the electrons; the contribution from each of these factors is dependent on the state of the atom. Since we considered only atoms with single electrons outside closed shells, Russell-Saunders coupling is a good approximation and the coefficients that relate the various g values are known. The ratio of the g_J values of two atoms that occur in different spectroscopic states yields g_S/g_L to an accuracy limited by the precision of observation and the precision with which the coefficients relating the various g values are known.

The intercomparison of g values to obtain a value of g_8 can only be made if atoms in several different spectroscopic states are available for observation. After our first investigation of the hyperfine structures of the alkali atoms, all in the ${}^{2}S_{1/2}$ state, Hardy and Millman (9) studied the hyperfine structure of indium in the ${}^{2}P_{1/2}$ state. Just after the war, Becker and I (10) determined the interaction constants that characterize the hyperfine structure of both the isotopes of gallium in the ${}^{2}P_{1/2}$ state. Gallium atoms in the excited and metastable ${}^{2}P_{3/2}$ state also occur in an atomic beam, and it was possible to determine the interaction constants for both isotopes in this state as well. We thus had available for study atoms in three different spectroscopic states.

In principle, the determination of the ratio of two g_J values is simple. Suppose we observe transitions for which F is constant and m_F changes by ± 1 for two different atoms or for the same atom in two different states. To the extent to which strictly low field conditions prevail, all lines in a given F state have the same frequency, and the ratio of the frequencies of such lines in two different states at a fixed field is simply the ratio of the g_F values. From this ratio, the ratio g_S/g_L for the electron may readily be derived with some additional knowledge of the properties of the nucleus in each atom. However, the hyperfine structure splitting of atomic states is generally small (from 200 to 20,000 megacycles per second) and the energies of the levels are far from linearly dependent on magnetic field at usefully high fields. Nevertheless, it is possible to obtain expressions for the energies of all levels in the hyperfine structure in terms of the zero field hyper-

Comparison	Nominal	Observed	g_s/g_L
$g_J({}^2P_{3/2}\text{Ga})/g_J({}^2P_{1/2}\text{Ga})$	2	$2(1.00172 \pm 0.00006)$	$2(1.00114 \pm 0.00004)$
$g_J(^2S_{1/2}\text{Na})/g_J(^2P_{1/2}\text{Ga})$	3	$3(1.00242 \pm 0.00006)$	$2(1.00121 \pm 0.00003)$
$g_J(^2S_{1/2}Na)/g_J(^2P_{1/2}In)$	3	$3(1.00243 \pm 0.00010)$	$2(1.00121 \pm 0.00005)$

fine structure splitting, the ratio g_N/g_J , and the quantity $g_J \mu_0 H/h$, or, where such expressions cannot be explicitly found, to determine the energies from the appropriate secular determinants. From the observed frequencies of appropriate lines and with a prior knowledge of the interaction constants which characterize an atom in the state in question, it is then possible to determine $g_J \mu_0 H/h$. Measurement of this quantity at the same field for atoms in two different states yields, at once, the important ratio g_8/g_L . The determination is independent of a knowledge of the magnetic field and of any fundamental constants.

Experimental Details

It is, perhaps, worthwhile to remark on some experimental details. The field in which the transition frequency was measured was so chosen that all the observed lines had a frequency of the order of 1 megacycle per second, per gauss. To avoid excessive distortion of the lines due to inhomogeneity of the field, a great deal of adjustment of the field was required before the lines approximated in width their theoretical value.

Special arrangements were made to allow the rapid interchange of ovens so that lines of different atoms could be measured in rapid succession. A considerable number of oscillators was required so that several frequencies which differed by large factors could be applied to the radio-frequency circuits that induced the transitions. Although the lines should, in principle, be measured at a fixed if unknown field, the actual measurements were made in a field that varied monotonically throughout a series of observations. The variation of field has the annoving effect of requiring a large body of data to establish the frequencies of two or more lines at a fixed field, but it also aids in avoiding repetitive errors that may occur when a reading of a fixed quantity is repeated.

Three intercomparisons of g_J values were made in these experiments. The results are given in Table 1.

It is to be noted that the ratio g_S/g_L which has been determined has been found from the ratio of the g_J values on the basis of the assumption that the coupling is Russell-Saunders coupling. Hence the deviation of the ratio g_S/g_L from its nominal value of 2 as determined from any pair of atoms or any pair of states does not constitute clear evidence that the spin moment of the electron is other than 1 Bohr magneton because of the possibility of occurrence of significant perturbations of the states. Theoretical arguments, however, indicate that such perturbations must be small. On experimental grounds, the agreement of the ratio obtained in three different ways from different atoms in different spectroscopic states offers overwhelming evidence that the spin momint of the electron does indeed differ from its nominal value by the indicated amount. The discrepancies between the three values of the ratio may, however, arise from perturbations of the indicated energy levels.

A later intercomparison of the g_J values of the alkali atoms and a comparison of the g_J value of potassium and hydrogen has demonstrated that the g_J values of the three alkali atoms of lowest atomic number are indeed equal to the spin gvalue of the electron to within 1 part in 40,000. A further intercomparison by Mann and me (11) of the g_J values of indium in the ${}^2P_{1/2}$ and ${}^2P_{3/2}$ states has given further confirmation to the interpretation of the discrepancy between a measured ratio of g_J values and the nominal value.

The experiments that have been described were performed at a field of about 400 gauss. In a wholly independent series of experiments, Taub and I (12) determined the ratio of the g_J value of indium in the ${}^2P_{1/2}$ state and that of sodium in the ${}^2S_{1/2}$ state by observations of lines in the hyperfine structure spectrum at fields that ranged from 3300 to 12,000 gauss. The method was to determine the nuclear g value of the proton in an alkali hydroxide in terms of the g_J values of indium and sodium. The result, insofar as it concerns the proton, is of no further interest here in view of the highly refined experiments which have been made in later years that allow the precise and direct determination of the nuclear g value of the proton in terms of both the nuclear and the Bohr magneton. The result, however, is of interest because it yields again the ratio of the g_J values in two different states on the basis of measurements at fields which differed from those in the earlier experiments by an order of magnitude. We found that

$g_s/g_L = 2(1 + 0.00119)$

We may therefore conclude on the basis of all evidence that the electron does indeed possess an "intrinsic" or "anomalous" magnetic moment over and above that deduced from the Dirac theory and whose magnitude is very close to 0.119 percent of the Bohr magneton.

Theoretical Interpretation

Perhaps it is well, at this point, to make a brief statement of the theoretical status of the spin magnetic moment of the electron. Soon after the publication of our first results, which gave substance to the assertion that the electron does have an anomalous moment, Schwinger (13) gave a result, based on new procedures in quantum electrodynamics, that

 $g_s/g_L = 2(1 + \alpha/2\pi) = 2(1.00116)$

The result is in excellent agreement with experimental measurements of the same quantity. The effect of the increased electron moment arises essentially as a consequence of the quantization of the electromagnetic field which always has a residual zero-point amplitude. Although the existence of this field had previously been recognized, it had not been possible to deal with the interaction prior to the formulation of contemporary quantum electrodynamics. The importance of the observation of the anomalous magnetic moment of the electron is in part in the demonstration that the procedures of quantum electrodynamics are, in fact, satisfactory in formulating a description of nature.

High-Precision Measurements

It is obvious that a more detailed study of the magnetic moment of the electron than that described thus far was desirable. The objective of a more extended investigation lies in the avoidance of theoretical difficulties in the interpretation, to a high precision, of the electronic g values of complex atoms. In the absence of substantial difficulties of interpretation, the very great precision of which spectroscopy by the method of atomic beams is capable may be used to obtain results of sufficient precision to test the validity of the calculations of quantum electrodynamics when they are made to a higher order than those originally made by Schwinger.

Barring only a measurement of the spin moment of the free electron itself, the best measurement that one may hope to make is on the electron in the hydrogen atom. In this atom in the ground state, the electron has no orbital angular momentum and hence there is no contribution to the electronic magnetic moment from the orbital motion. The entire electronic magnetic moment arises from the spin moment of the electron. Koenig, Prodell, and I (14) have determined the ratio of the electronic g value g_J of the hydrogen atom and the nuclear g value of the proton by experimental procedures to be described. To a very high order of accuracy, g_J is equal to g_S' , the spin g value of the electron bound in the hydrogen atom. The value of $g_{s'}$ differs from g_S of the free electron through a small relativistic effect of about 18 parts per million. Corrections due to a mixing of states and relativistic effects are well known and do not limit the accuracy with which the ratio g_S/g_P may be determined at the present time. Gardner and Purcell (15) have measured the ratio $2g_L/g_P$ of the cyclotron frequency of the electron and the precession frequency of the proton in a magnetic field to an accuracy of about 1 part in 80,000. Our result, when combined with that of Gardner and Purcell, yields g_S/g_L .

As a preliminary procedure, Prodell and I (16) determined the hyperfine structure separation of hydrogen with high precision. A subsequent investigation by both Wittke and Dicke (17) and by us (18) indicated an excessively optimistic estimate of the uncertainty. However, the value of the zero field hyperfine structure splitting of hydrogen that we used was sufficiently good to contribute no error to the value of g_J/g_P that was comparable to other uncertainties.

The apparatus designed for the purpose of these experiments had for the magnet that determined the transition frequencies one with a much better field homogeneity than that which usually characterizes the magnets used in atomic beams experiments. Ordinarily in an atomic beams apparatus, the magnet that determines the splitting of the levels is internal to the vacuum system. This arrangement permits small magnet gaps to be used and hence the production of large fields with electromagnets of moderate dimensions and power consumption. The use of a small gap, however, leads to a considerable hazard of field inhomogeneity. In the present case, the magnet was external to the vacuum envelope, the pole faces were of large diameter to reduce edge effects, and the magnet could be carefully shimmed after each change of externally imposed experimental parameters and from day to day to give good homogeneity in the volume within which transitions were observed. The deflecting magnets consisted of current-carrying conductors rather than the iron magnets that have become conventional in atomic beams experiments. This choice was made because of the smaller distortion of the transition field by current-carrying conductors than by massive blocks of iron.

Results

The experiment involved the measurement of the frequency of transition between the levels m = 0 and m = -1 in the state for which F = 1 alternately with the proton resonance frequency in the same magnetic field. The frequency of the first of these lines is of the order of 3600 megacycles per second at a field of 1500 gauss. The frequency of the proton resonance line is about 6.5 megacycles per second at the same field and is found by the methods of nuclear resonance in the same region of space as that traversed by the beam. An important component of the equipment is a device that can insert a cylindrical sample of water or mineral oil into a region as closely coincident as inherent limitations permit to that in which the atomic line has been observed. Various small corrections relating to the residual inhomogeneity of the field, bulk diamagnetism of the matter in the cylindrical sample that we employed, the presence of paramagnetic ions when we observed the resonance in water, and the differential internal diamagnetic shielding between oil and water must be applied.

We found that

$g_J/g_P = -658.2171 \pm 0.0006$

where g_P is the nuclear g value observed in a spherical sample of mineral oil. It is to be noted that this is only an apparent value of g_P because the externally applied field is modified by the internal diamagnetic shielding of the proton by the electrons in the molecules containing the proton. It is, nevertheless, of value to give the result in this form since the ratio $2g_L/g_P$ measured by Gardner and Purcell also refers to a spherical sample of mineral oil.

Application of a small relativistic term yields g_s , the spin g value of the electron in terms of g_p :

$g_{\rm S}/g_P = -658.2288 \pm 0.0006$

The combination of this result with that of Gardner and Purcell

$$2g_L/g_P = -657.475 \pm 0.008$$

yields

 $g_s/g_L = 2(1.001146 \pm 0.000012)$

where the principle uncertainty arises from the result of Gardner and Purcell. Since g_L equals 1, we can write

 $g_s = 2\mu_s = 2(1.001146 \pm 0.000012)$

where μ_s is the spin magnetic moment of the electron in terms of μ_0 .

The same result has subsequently been obtained by Beringer and Heald (19), who used a different experimental method involving, for atomic hydrogen, a microwave absorption technique and, for the observation of the proton resonance frequency, the usual nuclear resonance technique. The primary result obtained by them was

$$g_J/g_P = -653.2181 \pm 0.0003$$

In view of the stated uncertainties and the possibility of differences in the internal diamagnetic shielding in different samples of mineral oil, the agreement is good. Because of the limited accuracy for the result $2g_L/g_P$, the value of g_S/g_L is not affected, within the range of its uncertainty, by the discrepancy in the two results.

Conclusion

It is interesting to examine the ratio of g_{s}/g_{L} obtained by the sequence of experiments just described in light of the theoretical calculations of the electron moment. The result gives unambiguous evidence that the electron moment is anomalous and that the deviation of the moment from its nominal value is about $\alpha \mu_0/2\pi$. Karplus and Kroll (20) have calculated to a higher order the radiative correction to the spin moment of the electron and have found for the spin g value

$$g_s = 2(1 + \alpha/2\pi - 2.973 \ \alpha^2/\pi^2) = 2(1.0011454)$$

The result of the experiment is in remarkable agreement with the calculation, especially since the uncertainty in the experiment is much greater than the discrepancy between the experimental and calculated values. The agreement offers conclusive evidence of the validity of the calculation to the order α and very strong support to the validity of the calculations to the order α^2 . Thus the new procedures of quantum electrodynamicswhich have, perhaps, a questionable a priori validity-are demonstrated to be, in practice, valid for the interpretation of certain observed phenomena and, therefore, useful in the exploration of other aspects of the behavior of matter.

Airborne Radioactivity

Eizo Tajima and Tadayoshi Doke

Artificial radioactivity, in addition to the natural activity in the atmosphere, has been observed after atomic bomb tests. Other investigations in this field might be in progress, but few articles are available. Relatively thorough documents have been published by M. Eisenbud and J. H. Harley on radioactive fallout in the United States (1, 2). The present report describes the results of measurements of airborne radioactivity in Japan from 16 March to 4 May 1955 (3). We could detect not only the radon and thoron daughter products but also artificial radioactivity, including alpha emitters. The artificial radioactivity appears to have two or more components of different origins. The correlation of the daily variation with meteorological conditions is shown.

The air samplers used in this experiment were of the same type as the "coffee pot" sampler that was presented to the Japanese Science Council by the United States at the Japan-United States Radiobiological Conference that was held in the fall of 1954. A fan draws the air through a No. 41 Whatman filter paper 11 centimeters in diameter. The filter efficiency for normal dust is said to be high but it is not known for fine dust. Although a membrane filter has a higher efficiency for fine dust, it could not be used because the small air flow is not enough to cool the fan motor.

The change in flow rate of the air was checked by an anemometer because clogging of the filter paper might decrease it. The initial rate of the air flow was 27 cubic meters per hour; the rate did not change during the 24 hours of operation. On an average, the flow of the air for 24 hours of operation could be measured as 650 cubic meters with an

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error of less than \pm 10 percent. Little difference was observed among the three samplers that were used.

The efficiency of the filter paper was checked at high humidity. A pair of samplers was made to draw the air through chimneys that were each 3 meters in length and 11.5 centimeters in diameter. One of the chimneys was furnished with a long water-filled boat and a Nichrome winding. After 4 hours' simultaneous operation of both samplers, the natural radioactivity on the filter paper was measured. About 70 milliliters of water was evaporated during the period of operation. The cleaning efficiency of the filter paper in moist air was found to be only 4 percent greater than it was for ordinary air.

The filter papers were ashed in a crucible and the beta radioactivity of the ash was measured by a conventional Geiger-Müller counter with a mica window. The counting efficiency was estimated at 16.6 percent from a 75-disintegration-per-minute substandard of Ra-E. The activity of all samples was measured once a day for about 1 month to get the decay curve. Aluminum absorption curves of some samples of special interest were also run.

For alpha measurements, a scintillation counter with adjustable sample-to-phosphor distance was used. The counting efficiency for alpha particles of 5.3 Mev

The authors are on the faculty of St. Paul's University, Ikebukuro Toshima-ku, Tokyo, Japan.