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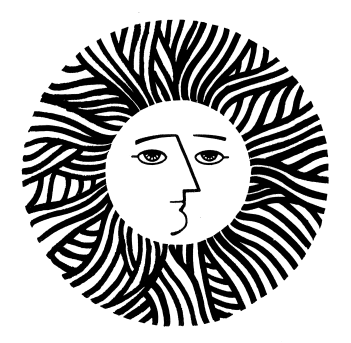
## ENERGY & ENVIRONMENT DIVISION

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# AN INTERCOMPARISON OF TRACER GASES USED FOR AIR INFILTRATION MEASUREMENTS

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#### ABSTRACT

A direct intercomparison has been made between common tracer gases used to measure air infiltration rates in buildings. The results indicate that air exchange rates measured using sulfur hexaflouride,  $\mathrm{SF}_6$ , are slightly larger than those measured using methane,  $\mathrm{CH}_4$ , or nitrous oxide,  $\mathrm{N}_2\mathrm{O}$ . The ratio of air change rates measured using  $\mathrm{SF}_6$  to air change rates measured concurrently using a lighter tracer gas was found to be 1.10  $\pm$  0.10.

#### INTRODUCTION

The energy cost of excessive air infiltration in buildings has spurred a sharp increase in the number of research projects investigating the magnitude of this phenomenon. The most common technique used to monitor air infiltration rates requires measurements of the concentration of a tracer gas. The tracer gas, a material easily monitored which normally is not present in the atmosphere, is injected into the space to be tested. When the injection ends, the concentration of tracer is measured as a function of time. Outside air, leaking into the test space, replaces the tracer - indoor air mixture which leaks out at the same rate. The rate of change of the concentration of tracer in the indoor air is therefore proportional to the concentration of tracer in the test space; i.e. the concentration decreases exponentially.

This verbal description of the measurement process contains several assumptions about the nature of air infiltration. For example, it assumes that the rate of air infiltration remains constant during the measurement period. In addition, it assumes that the outside infiltrating air mixes uniformly with the indoor air during measurement. In this investigation we examine the assumption that measured air change rates are independent of the type of tracer gas used if adequate mixing of the tracer in the test space occurs.

An ideal tracer gas should meet the following criteria [1]:

- (1) be inexpensive,
- (2) be easily measurable at low concentrations,
- (3) be non-toxic and non-allergenic,
- (4) be non-flammable,
- (5) have approximately the same molecular weight as air,
- (6) not be absorbed on any surfaces within the space under test,
- (7) not be a normal constituent of the air in the test space.

No tracer gas meets all these requirements. Within recent years improvements in instrumentation have permitted measurements of concentration of sulfur hexaflouride,  $SF_6$ , at levels of

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parts per billion. This is three orders of magnitude smaller than previous techniques which yield measurements in the range of parts per million. The ability to observe such low concentrations makes  ${\rm SF}_6$  an attractive possibility for use as a tracer gas. However, several researchers have expressed the concern that the large molecular weight of  ${\rm SF}_6$  (146) will cause stratification of the tracer gas after injection. Hunt  $^{[1]}$  has discussed the errors which will result if a tracer gas is mixed poorly within the test space. Stratification, which will yield poor mixing, will lead to erroneous infiltration results unless multiple sampling and properly weighted averaging of the tracer from several points occurs.

The experiment described in this paper was designed to make concurrent direct comparisons of air exchange rates measured with different tracer gases in the same test space.

#### LITERATURE REVIEW

The paper of Hitchin and Wilson [2] is an excellent review of the experimental techniques used in measuring air infiltration. This work has been extended to include work through 1978 by Hunt's careful review of current techniques [1]. Both authors cite previous direct intercomparison results. These are shown in Table 1.

Warner  $^{[3]}$  reported comparisons between coal gas (a mixture of  $\mathrm{H}_2$  and CO obtained by passing steam over hot carbon) detected using a Katharometer and  $\mathrm{CO}_2$  whose concentration was measured by Haldane gas analysis.

Collins and Smith  $^{[4]}$  used the radioactive argon isotope  $^{41}$ A as a tracer; its concentration was measured with a geiger counter and a ratemeter. A direct comparison was made of the infiltration rate detected using  $H_2$  detected with a katharometer and  $^{41}$ A; agreement within 8% was seen in 2 trials.

Howland, Kimber and Littlejohn  $^{[5]}$  reported comparisons between air changes measured with the radioactive isotope  $^{85}$ Kr using a geiger counter and a ratemeter. The decay rates were compared with measurements which used  ${\rm CO}_2$  as the tracer. Its concentration was determined by drawing samples of air periodically and using chemical analysis (the Haldame apparatus ) to find the amount of tracer remaining in the test space. Results of 3 tests varied by about 9%.

Lidwell  $^{[6]}$  compared results obtained when nitrous oxide, N<sub>2</sub>0, is compared with acetone, C<sub>3</sub>H<sub>6</sub>0, as a tracer gas. Infrared absorption was used to measure the concentration of N<sub>2</sub>0; acetone concentration was determined by measuring the change in pH which occurred when air containing acetone is absorbed into solutions of hydroxylamine hydrochloride. A single measurement (judged to be accurate to within 10%) produced 3% agreement.

Howard  $^{[7]}$  compared  $^{N}_20$  with both  $^{H}_2$  and  $^{O}_2$ .  $^{N}_20$  concentrations were determined with an infrared analyzer,  $^{H}_2$  concentrations with a katharometer and  $^{O}_2$  by absorption in aqueous chromous chloride. Specific results are not quoted in the paper. The author states that close agreement between decay rates using  $^{N}_20$  and  $^{O}_2$  were seen over wide ranges of wind speeds. On the other hand,  $^{H}_2$  decay rates were substantially higher than  $^{N}_20$ . The evidence suggested that diffusion of  $^{H}_2$  through the walls of the the unpainted gypsum of the test space was the source of the discrepancy. This hypothesis was tested by repeating the tests after the walls were sealed with 2 coats of latex paint and also repeating the tests in a laboratory with masonry walls. The discrepancy was not present in the latter 2 sets of tests.

Hunt and Burch  $^{[8]}$  compared air change rates using He and SF $_6$  as tracer gases to examine the influence of molecular diffusion on the infiltration process. Their test space was a 4-bedroom townhouse constructed within an environmental test chamber. If molecular diffusion were important in the infiltration process, the air change rate measured with He would be significantly larger than that measured with SF $_6$ . In fact, slightly larger air change rates were seen when SF $_6$  was used as the tracer rather than He. Six trials were made. The ratio of the air change rate measured with SF $_6$  to that measured with He was 1.17 with a standard deviation of 0.14.

#### TEST SPACE

The test reported in this paper were conducted in a 1-story unoccupied residence in Walnut Creek, Calif. This house is leased by the Lawrence Berkeley Laboratory as a research facility for the Energy Efficient Building program. A plan view of the house and its surroundings is shown in Fig. 1. The floor plan of the house is shown in Fig. 2. The volume of the living space of the house is 230 m $^3$ , its floor area is 100 m $^2$  and the area of the 6 surfaces bounding the living space is 300 m $^2$ .

The windows of the house are single-glazed aluminum sliding windows with rubber weather-stripping. The house has 2 wooden exterior doors and a single sliding glass patio door. Prior to these tests, one of the doors had been removed and replaced by a plywood panel upon which a fan was mounted for air leakage tests.

The house is heated with a forced air gas-fired furnace system. A central air-conditioning system uses the same duct system. The return duct opening located in the ceiling of the hallway leading to the bedrooms; and the supply duct diffusers located on the floor throughout the house, are both shown in Fig. 2.

#### TEST PROCEDURE

All tests in this intercomparison used a tracer gas concentration decay technique to measure the air exchange rate of the test space.

During all tests except test 8 and 9 (cf. Table 3,) the doors and windows were closed and the furnace was off. During tests 8 and 9 windows were open 1 cm. The furnace blower, however, ran continuously to provide mixing for the injection of the tracer gas and to provide a suitable location for sampling the air-tracer gas mixture. The blower capacity of the furnace fan was measured and was found to be 0.4 m<sup>3</sup>/s; this is equivalent to a volume flow rate of 6 house volumes/hr.

Indoor temperature and relative humidity were measured using a hygrothermograph located in the living room. Outdoor weather conditions, dry-bulb temperature, wind speed and direction, were measured using a weather tower mounted on the roof of the garage (cf. Fig. 1).

The fireplace chimney and the stove vent in the kitchen were both covered with plastic and taped closed.

The groups conducting the tests, and the tracer gases used are shown in Table 2.

#### ANALYSIS PROCEDURE

After injection, the rate of change of tracer gas concentration is the product of the air change rate, A, and the concentration in the test space.

$$\frac{dC}{dt} = -AC \tag{1}$$

If the air change rate is constant the solution of Eq 1 is simply

$$C(t) = C_0 e - At$$
 (2)

where  $C_0$  is the concentration of tracer at time t = o.

Several procedures can be used to analyze data relating concentration and time. In these investigations, concentration was plotted as a function of time using semilog graph paper.

Three  $SF_6$  decays, trials 6, ll and l4 were rejected after examining the graphs. Either non-uniform mixing, non-constant decay rates or instrumentation malfunctions caused this behavior.

#### EXPERIMENTAL RESULTS

Table 4 shows the results of the measurements.

The mean value of the ratio for all the tests was 1.10  $\pm$  0.10. The mean value for the comparisons of SF<sub>6</sub> with N<sub>2</sub>O was 1.09  $\pm$  0.09 while the value for the comparisons with CH<sub>4</sub> was 1.16  $\pm$  0.09.

The uncertainties listed with each of the ratios is the standard deviation of a single measurement. The t distribution with 9 deg. of freedom was used to calculate the expected range of the ratio. This result predicts that the actual range of the ratio, r, lies within the range

$$1.01 \le r \le 1.20$$

at the 99% level of confidence <sup>[9]</sup>. The t distribution assumes sampling from an infinite sample in which the scatter of results is only due to random effects. Our results, therefore, suggest that:

- (a) Systematic errors exist in the measurement procedure which results in SF $_6$  concentration decays that are too large, or N $_2$ O and CH $_4$  decays which are too small; or
- (b) Sulfur hexaflouride overestimates the "true" air infiltration rate when used as a tracer gas.

We remind the reader that the results above refer to 2 independent sets of measurements of the concentration decay of  $SF_6$  that were made using equipment at Princeton and LBL.

A source of additional information to help resolve this problem is contained in the paper of Hunt and Burch  $^{[8]}$ . As discussed above, these authors compared tracer gas measurements with the measurements we have described, we obtain a ratio  $^{A}_{SF}$ /A lighter gas of 1.13 + 0.12. Again, if we use the t- distribution with 15 deg. of freedon we predict that the actual ratio, r bar, will lie within the range

$$1.04 \le \frac{1}{r} \le 1.22$$

with 99% confidence.

Adding an additional set of independent measurements to the sample of measurements described in this report did not increase the likelihood that the discrepancy seen is the result of systematic errors in the measurements. Rather it supports hypothesis (b).

The results show that a difference exists between air exchange rates measured using  $SF_6$  and air exchange rates measured using lighter tracer gases. The difference, however, is small and represents the range of uncertainty which we estimate is present in any tracer gas measurement (5% to 10%). An example of this is test 12 in the measurements reported above. The air exchange rates measured using  $C_2H_6$ ,  $SF_6$  and  $N_2O$  were 0.68 hr<sup>-1</sup>, 0.66 hr<sup>-1</sup>, and 0.61 hr<sup>-1</sup>. These values represent the range of values seen whenever air exchange rates are measured; consequently the differences seen in this intercomparison are scarcely large enough to be significant.

We have examined the measurement process for physical effects which would bias the data in the direction observed. On this basis two effects, molecular diffusion and absorption of  $N_2O$  by water vapor, can be eliminated immediately.

If molecular diffusion were important in air infiltration, air change rates measured with light gases which have larger thermal speeds would be larger than those measured with heavy gases. This is the opposite of what we have observed.

If a significant amount of  $\rm N_20$  were absorbed by water vapor in the test space, air exchange rates measured with  $\rm N_20$  would tend to be larger than those using  $\rm SF_6$ . Again the opposite result was actually seen.

Another possibility to consider is settling of the tracer gas in the test space. Since the tracer is sampled at the return duct of the furnace, which is located on the ceiling in the test space, settling of the heavy gases during the course of the measurement would appear to increase the air exchange rate measured using a heavy tracer gas such as  $SF_6$ .

However, a simple calculation will show that this is quite unlikely. The tracer gases were injected into the return duct of a forced air heating system and are well mixed after a short time. Other results  $^{\left[10\right]}$  allow us to quote a mixing time of the order of 5 min. for this house. Therefore, after 5 min. the tracer gas is well mixed throughout the test space — and this mixing continues throughout the concentration decay measurement.

Buoyancy effects occur on the macroscopic, not microscopic, level. The fractional difference in density between a macroscopic volume of gas containing air and one containing 1 ppb (part per billion)  $SF_6$  is  $4 \times 10^{-9}$ . Since the acceleration due to buoyant forces is (P)g, the effective acceleration of our volume element containing  $SF_6$  is  $4 \times 10^{-10}$  g. It would take about 3 hr for such an element to settle 2m in still air under an acceleration of that magnitude. However, since the furnace blower moves 6 volumes of house air through it each hour, forced mixing and convective mixing certainly dominate buoyancy effects. We conclude that stratification due to the heavy  $SF_6$  molecule is unlikely after the gas is initially mixed with room air.

Other possibilities exist. Rather than comparing tracer gases we may, in fact, be comparing instrumentation system. The  ${\rm SF}_6$  is detected with electron capture gas chromatographic techniques while the concentrations of the lighter gases are measured using infrared absorption techniques or the change in thermal conductivity of helium-air mixtures.

Another possibility may be absorption rate could have a significant effect on measurements in the parts per billion range while they would not be noticed in measurements of concentrations of parts per million.

We must be careful to emphasize, after this extensive discussion, that we are searching for an explanation to a discrepancy that is typical of the uncertainties seen in infiltration measurements. While the difference may be real, it should not preclude use of one gas in preference for another when chosing a tracer gas.

#### CONCLUSIONS

A direct comparison of air change rates measured using  $SF_6$  and either  $N_2O$  or  $CH_4$  shows that  $SF_6$  gives a slightly larger value than the lighter gases. The difference cannot be explained using ideas of molecular diffusion. Stratification due to the settling of the heavier tracer gas seems to be an unlikely cause of the effect. While the difference appears to be real, it is small enough so that it is likely unnoticed in the uncertainty associated with a single tracer gas measurement.

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#### ACKNOWLEDGEMENTS

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TABLE 1
Published Comparisons between Tracer Gases

Reference	<u>Tracer</u> <u>Gases</u>	Number of Tests	Results*
Warner [3]	Coal Gas, CO <sub>2</sub>	3	1.05 <u>+</u> 0.18
Collins and Smith [4]	$H_2$ , $^{41}A$	2	0.93 <u>+</u> 0.01
Howland, et al. [5]	co <sub>2</sub> , <sup>85</sup> Kr	3	1.00 <u>+</u> 0.09
Lidwell [6]	N <sub>2</sub> O, С <sub>3</sub> H <sub>6</sub> O	. 1	0.97
Howard [7]	$H_2$ , $N_2O$	many	agreement
Howard [7]	0 <sub>2</sub> , N <sub>2</sub> 0	many	agreement
Hunt and Burch [8]	He, SF <sub>6</sub>	6	1.17 +- 0.14

The results quoted are the mean values of the ratios of the measured air change rates.

The ratio is formed by dividing the air change rate of the heavier gas by the air change rate of the lighter gas.

TABLE 2

Research Group	Tracer Gas	<u>Detector</u>
Honeywell	сн <sub>4</sub>	IR Analyzer
Princeton	sf <sub>6</sub>	Elctron Capture
LBL	$N_2O$	IR Analyzer
LBL	<sup>C</sup> 2 <sup>H</sup> 6	IR Analyzer
LBL	sf <sub>6</sub>	Electron Capture

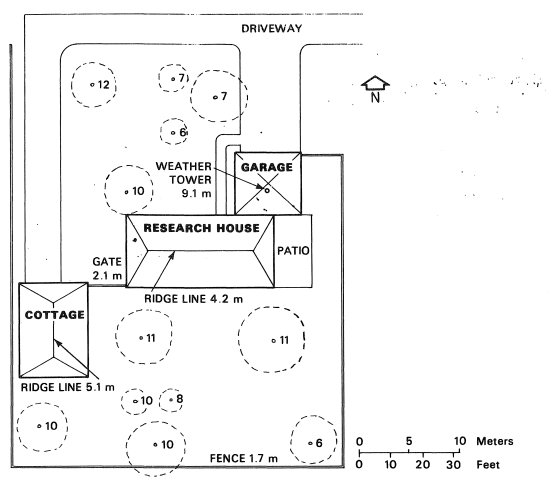
Interference between  $\text{CH}_4$  and  $\text{N}_2\text{O}$  prevented simultaneous measurements of tracer gas concentrations using these gases; therefore the tests were organized using the schedule shown in Table 3.

TABLE 3

<u>Date</u>	<u>Test Number</u>	<u>N20</u>	<u>sf</u> 6	<u>сн<sub>4</sub></u>	<u>с<sub>2</sub>н<sub>6</sub></u>
5/9/78	1	x			
5/9/78	2			x	
5/9/78	3	x	x(P)*		
5/9/78	4		x(P)	x	
5/9/78	5	<b>X</b>	<b>x(B)</b>		
5/10/78	6	x	x(B)		
5/10/78	7		<b>x(B)</b>	x	
5/10/78	8		x(B)	x	
5/10/78	9	<b>x</b>	<b>x</b> (B)		
5/10/78	10		<b>x(B)</b>	x	
5/10/78	11	x	x(B)		
5/12/78	12	x	x(B)		x
5/12/78	13	x	x(B)		

\*Test 3 and 4 using  $\rm SF_6$  were analyzed at Princeton (indicated by (P)) while the  $\rm SF_6$  tests 5 through 14 were analyzed at Berkeley (indicated by (B)).

The samples analyzed at Princeton were collected at the test site in aluminized mylar sample bags which were then shipped from California to New Jersey.



(Numbers are height of trees in meters)

Fig. 1

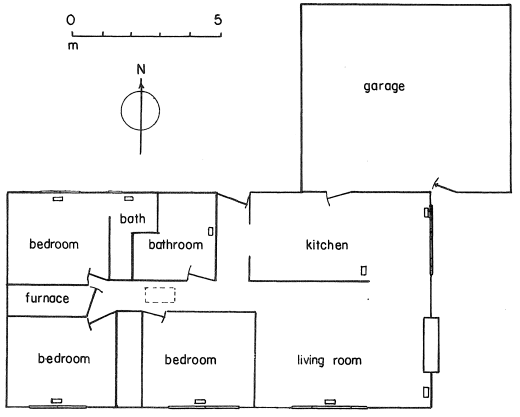


Fig. 2

#### ABSTRACT

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All tests in this intercomparison used a tracer gas concentration decay technique to measure the air exchange rate of the test space.

During all tests except test 8 and 9 (cf. Table 3,) the doors and windows were closed and the furnace was off. During tests 8 and 9 windows were open 1 cm. The furnace blower, however, ran continuously to provide mixing for the injection of the tracer gas and to provide a suitable location for sampling the air-tracer gas mixture. The blower capacity of the furnace fan was measured and was found to be  $0.4~\rm m^3/s$ ; this is equivalent to a volume flow rate of 6 house volumes/hr.

Indoor temperature and relative humidity were measured using a hygrothermograph located in the living room. Outdoor weather conditions, dry-bulb temperature, wind speed and direction, were measured using a weather tower mounted on the roof of the garage (cf. Fig. 1).

The fireplace chimney and the stove vent in the kitchen were both covered with plastic and taped closed.

The groups conducting the tests, and the tracer gases used are shown in Table 2.

#### ANALYSIS PROCEDURE

After injection, the rate of change of tracer gas concentration is the product of the air change rate, A, and the concentration in the test

space.

$$\frac{dC}{dt} = -AC \tag{1}$$

If the air change rate is constant the solution of Eq 1 is simply

$$C(t) = C_0 e^{-At}$$
 (2)

where  $C_0$  is the concentration of tracer at time t = 0.

Several procedures can be used to analyze data relating concentration and time. In these investigations, concentration was plotted as a function of time using semilog graph paper.

Three  ${
m SF}_6$  decays, trials 6, 11 and 14 were rejected after examining the graphs. Either non-uniform mixing, non-constant decay rates or instrumentation malfunctions caused this behavior.

#### EXPERIMENTAL RESULTS

Table 4 shows the results of the measurements.

The mean value of the ratio for all the tests was 1.10  $\pm$  0.10. The mean value for the comparisons of SF<sub>6</sub> with N<sub>2</sub>O was 1.09  $\pm$  0.09 while the value for the comparisons with CH<sub>4</sub> was 1.16  $\pm$  0.09.

The uncertainties listed with each of the ratios is the standard deviation of a single measurement. The t distribution with 9 degrees of freedom was used to calculate the expected range of the ratio. This result predicts that the actual range of the ratio,  $\overline{r}$ , lies within the range

$$1.01 \le \overline{r} \le 1.20$$

at the 99% level of confidence [9]. The t distribution assumes sampling from an infinite sample in which the scatter of results is only due to random effects. Our results, therefore, suggest that:

- (a) Systematic errors exist in the measurement procedure which results in  ${\rm SF}_6$  concentration decays that are too large, or  ${\rm N}_2{\rm O}$  and  ${\rm CH}_4$  decays which are too small; or
- (b) Sulfur hexaflouride overestimates the "true" air infiltration rate when used as a tracer gas.

We remind the reader that the results above refer to  $\,2\,$  independent sets of measurements of the concentration decay of SF $_6$  that were made using equipment at Princeton and LRL.

A source of additional information to help resolve this problem is contained in the paper of Hunt and Burch [8]. As discussed above, these authors compared tracer gas measurements with the measurements we have described, we obtain a ratio  $^{A}SF_{c}/^{A}$  lighter gas of 1.13 +- 0.12. Again, if we use the t- distribution with 15 degrees of freedon we predict that the actual ratio,  $\overline{r}$ , will lie within the range

$$1.04 \le \overline{r} \le 1.22$$

with 99% confidence.

Adding an additional set of independent measurements to the sample of measurements described in this report did not increase the likelihood that the discrepancy seen is the result of systematic errors in the measurements. Rather it supports hypothesis (b).

The results show that a difference exists between air exchange rates measured using SF<sub>6</sub> and air exchange rates measured using lighter tracer gases. The difference, however, is small and represents the range of uncertainty which we estimate is present in any tracer gas measurement (5% to 10%). An example of this is test 12 in the measurements reported above. The air exchange rates measured using  $\rm C_{2H_6}$ , SF<sub>6</sub> and N<sub>2</sub>O were 0.68 hr<sup>-1</sup>, 0.66 hr<sup>-1</sup>, and 0.61 hr<sup>-1</sup>. These values represent the range of values seen whenever air exchange rates are measured; consequently the differences seen in this intercomparison are scarcely large enough to be significant.

We have examined the measurement process for physical effects which would bias the data in the direction observed. On this basis two effects, molecular diffusion and absorption of  $N_{20}$  by water vapor, can be eliminated immediately.

If molecular diffusion were important in air infiltration, air change rates measured with light gases which have larger thermal speeds would be larger than those measured with heavy gases. This is the opposite of what we have observed.

If a significant amount of  $N_2O$  were absorbed by water vapor in the test space, air exchange rates measured with  $N_2O$  would tend to be larger than those using SF<sub>6</sub>. Again the opposite result was actually seen.

Another possibility to consider is settling of the tracer gas in the test space. Since the tracer is sampled at the return duct of the furnace, which is located on the ceiling in the test space, settling of the heavy gases during the course of the measurement would appear to increase the air exchange rate measured using a heavy tracer gas such as  $SF_6$ .

However, a simple calculation will show that this is quite unlikely. The tracer gases were injected into the return duct of a forced air heating system and are well mixed after a short time. Other results [10] allow us to quote a mixing time of the order of 5 min. for this house. Therefore, after 5 min. the tracer gas is well mixed throughout the test space — and this mixing continues throughout the concentration

decay measurement.

Buoyancy effects occur on the macroscopic, not microscopic, level. The fractional difference in density between a macroscopic volume of gas containing air and one containing 1 ppb (part per billion) SF  $_6$  is 4 x 10 $^{-9}$ . Since the acceleration due to buoyant forces is  $(\frac{\Delta \rho}{4})$  g, the effective acceleration of our volume element containing SF  $_6$  is 4 x 10 $^{-10}$  g. It would take about 3 hr for such an element to settle 2m in still air under an acceleration of that magnitude. However, since the furnace blower moves 6 volumes of house air through it each hour, forced mixing and convective mixing certainly dominate buoyancy effects. We conclude that stratification due to the heavy SF  $_6$  molecule is unlikely after the gas is initially mixed with room air.

Other possibilities exist. Rather than comparing tracer gases we may, in fact, be comparing instrumentation system. The  ${\rm SF}_6$  is detected with electron capture gas chromatographic techniques while the concentrations of the lighter gases are measured using infrared absorption techniques or the change in thermal conductivity of helium-air mixtures.

Another possibility may be absorption rate could have a significant effect on measurements in the parts per billion range while they would not be noticed in measurements of concentrations of parts per million.

We must be careful to emphasize, after this extensive discussion, that we are searching for an explanation to a discrepancy that is typical of the uncertainties seen in infiltration measurements. While the difference may be real, it should not preclude use of one gas in preference for another when chosing a tracer gas.

#### CONCLUSIONS

A direct comparison of air change rates measured using  $SF_6$  and either  $N_2O$  or  $CH_4$  shows that  $SF_6$  gives a slightly larger value than the lighter gases. The difference cannot be explained using ideas of molecular diffusion. Stratification due to the settling of the heavier tracer gas seems to be an unlikely cause of the effect. While the difference appears to be real, it is small enough so that it is likely to be unnoticed because of the large uncertainty associated with a single tracer gas measurement.

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TABLE 1
Published Comparisons between Tracer Gases

Reference	Tracer Gases	Number of Tests	<u>Results</u> *
Warner [3]	Coal Gas, CO <sub>2</sub>	3	1.05 <u>+</u> 0.18
Collins and Smith [4]	H <sub>2</sub> , 41 <sub>A</sub>	2	0.93 <u>+</u> 0.01
Howland, et al. [5]	co <sub>2</sub> , <sup>85</sup> Kr	3	1.00 <u>+</u> 0.09
Lidwell [6]	N <sub>2</sub> O, С <sub>3</sub> H <sub>6</sub> O	1	0.97
Howard [7]	$H_2$ , $N_2O$	many	agreement
Howard [7]	0 <sub>2</sub> , N <sub>2</sub> 0	many	agreement
Hunt and Burch [8]	He, SF <sub>6</sub>	6	1.17 +- 0.14

The results quoted are the mean values of the ratios of the measured air change rates.

The ratio is formed by dividing the air change rate of the heavier gas by the air change rate of the lighter gas.

TABLE 2

Research Group	Tracer Gas	<u>Detector</u>
Honeywell	СH <sub>4</sub>	IR Analyzer
Princeton	sf <sub>6</sub>	Elctron Capture
LBL	N <sub>2</sub> O	IR Analyzer
LBL	<sup>с</sup> 2 <sup>н</sup> 6	IR Analyzer
LBL	SF <sub>6</sub>	Electron Capture

Interference between  $\mathrm{CH}_4$  and  $\mathrm{N}_2\mathrm{O}$  prevented simultaneous measurements of tracer gas concentrations using these gases; therefore the tests were organized using the schedule shown in Table 3.

TABLE 3

Date	Test Number	<u>N20</u>	<u>sf</u> 6	<u>СН</u> 4	<u>С<sub>2</sub>Н6</u>	
5/9/78	1	x				
5/9/78	2			x		
5/9/78	3	x	x(P)*			ı
5/9/78	4		x(P)	x		
5/9/78	5	x	x(B)			
5/10/78	6	×	x(B)			
5/10/78	7		x(B)	x		
5/10/78	8		x (B)	x		
5/10/78	9	x	x(B)			
5/10/78	10		x(B)	x		
5/10/78	-11	x	x(B)			•
5/12/78	12	x	x(B)		x	ż.
5/12/78	13	x	x(B)			Û

\*Test 3 and 4 using  $SF_6$  were analyzed at Princeton (indicated by (P)) while the  $SF_6$  tests 5 through 14 were analyzed at Berkeley (indicated by (B)).

The samples analyzed at Princeton were collected at the test site in aluminized mylar sample bags which were then shipped from California to New Jersey.

•

19

Table 4

Wind Speed

6

-1

-3

-6

1.06

1.25

0.95

1.11

1.08

1.21

Ratio\* Number  $(hr^{-1})$ 0.49 1 1 \_ 2 0.75 0.71 1.11 3 0.64 -2 0.63 0.76 -4 1.21 5 0.76 7 7 1.10 0.69

0.94

1.59

1.19

0.80

0.66

0.70

Air Change Rates

0.89

0.72

. 1.27

0.91

1.25

0.51

0.61

0.58

0.47

Test

6

7

9

10

11

13

14

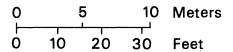
12\*\*

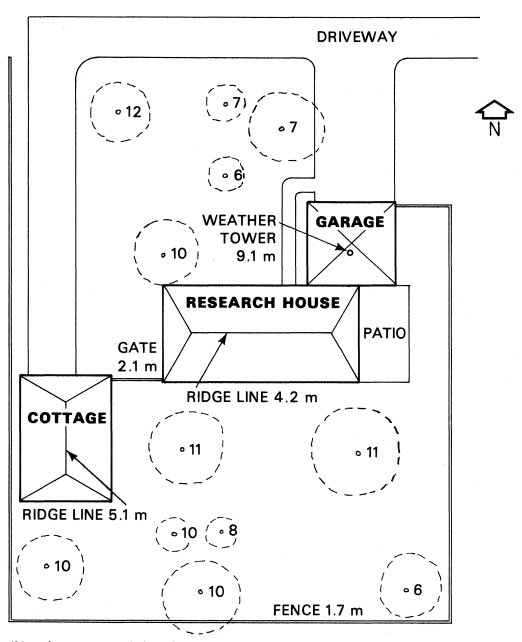
 $<sup>^*</sup>$ The ratio quoted is the air change rate of  ${\rm SF}_6$  divided by the air change rate of the lighter tracer gas

<sup>\*\*</sup>During test 12 ethane,  $C_2H_6$ , was also used as a tracer gas. The air exchange rate measured was 0.68 hr<sup>-1</sup>; this yields a ratio of 0.97.

#### FIGURE CAPTIONS

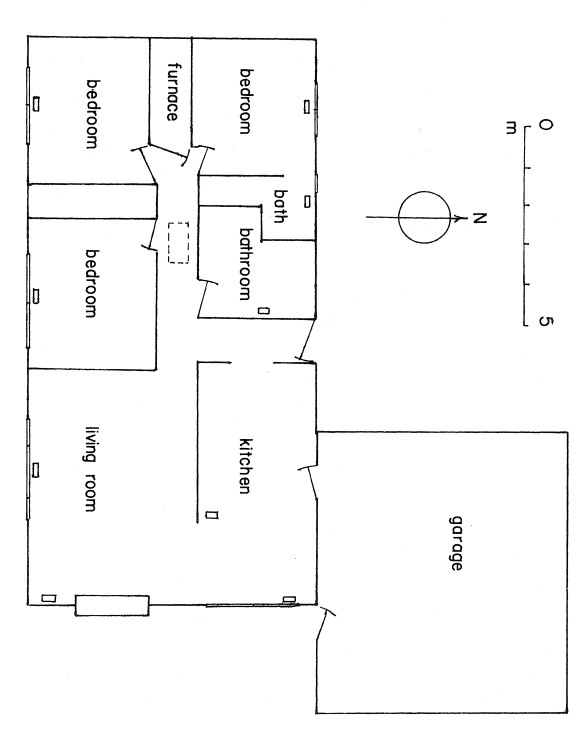
- Fig. 1 Plan view of the Walnut Creek Research House and its surroundings.
- Fig. 2 Floor plan of the Walnut Creek House. The return duct is located as shown in the hallway ceiling; the supply duct diffusers are located on the floor.





(Numbers are height of trees in meters)

XBL 788-1537



XBL 7910-12575

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