

MEASUREMENT AND INSTRUMENTS

<i>Terminology</i>	14.1	<i>Electric Measurement</i>	14.24
<i>Uncertainty Analysis</i>	14.2	<i>Rotative Speed Measurement</i>	14.24
<i>Temperature Measurement</i>	14.3	<i>Sound and Vibration Measurement</i>	14.24
<i>Humidity Measurement</i>	14.9	<i>Lighting Measurement</i>	14.28
<i>Pressure Measurement</i>	14.12	<i>Thermal Comfort Measurement</i>	14.28
<i>Velocity Measurement</i>	14.14	<i>Moisture Content and Transfer Measurement</i>	14.29
<i>Flow Rate Measurement</i>	14.17	<i>Heat Transfer Through Building Materials</i>	14.30
<i>Air Infiltration, Airtightness, and Outdoor Air</i>		<i>Air Contaminant Measurement</i>	14.31
<i>Ventilation Rate Measurement</i>	14.21	<i>Combustion Analysis</i>	14.31
<i>Carbon Dioxide Measurement</i>	14.22	<i>Data Acquisition and Recording</i>	14.31

HEATING, refrigerating, and air-conditioning engineers and technicians require instruments for both laboratory work and fieldwork. Precision is more essential in the laboratory, where research and development are undertaken, than in the field, where acceptance and adjustment tests are conducted. This chapter describes the characteristics and uses of some of these instruments.

TERMINOLOGY

The following definitions are generally accepted.

Accuracy. The capability of an instrument to indicate the true value of measured quantity. This is often confused with inaccuracy, which is the departure from the true value to which all causes of error (e.g., hysteresis, nonlinearity, drift, temperature effect, and other sources) contribute.

Amplitude. The magnitude of variation from its zero value in an alternating quantity.

Average. The sum of a number of values divided by the number of values.

Bandwidth. The range of frequencies over which a given device is designed to operate within specified limits.

Bias. The tendency of an estimate to deviate in one direction from a true value (a systematic error).

Calibration. (1) The process of comparing a set of discrete magnitudes or the characteristic curve of a continuously varying magnitude with another set or curve previously established as a standard. Deviation between indicated values and their corresponding standard values constitutes the correction (or calibration curve) for inferring true magnitude from indicated magnitude thereafter; (2) the process of adjusting an instrument to fix, reduce, or eliminate the deviation defined in (1).

Calibration curve. (1) The path or locus of a point that moves so that its coordinates on a graph correspond to values of input signals and output deflections; (2) the plot of error versus input (or output).

Confidence. The degree to which a statement (measurement) is believed to be true.

Dead band. The range of values of the measured variable to which an instrument will not effectively respond.

Deviate. Any item of a statistical distribution that differs from the selected measure of control tendency (average, median, mode).

Deviation, standard. The square root of the average of the squares of the deviations from the mean (root mean square deviation). A measure of dispersion of a population.

Distortion. An unwanted change in wave form. Principal forms of distortion are inherent nonlinearity of the device, nonuniform response at different frequencies, and lack of constant proportionality between phase-shift and frequency. (A wanted or intentional change might be identical, but it is called **modulation**.)

Drift. A gradual, undesired change in output over a period of time that is unrelated to input, environment, or load. Drift is gradual; if variation is rapid, the fluctuation is referred to as **cycling**.

Dynamic error band. The spread or band of output-amplitude deviation incurred by a constant amplitude sine wave as its frequency is varied over a specified portion of the frequency spectrum (see *Static error band*).

Error. The difference between the true or actual value to be measured (input signal) and the indicated value (output) from the measuring system. Errors can be systematic or random.

Error, accuracy. See *Error, systematic*.

Error, fixed. See *Error, systematic*.

Error, instrument. The error of an instrument's measured value that includes random or systematic errors.

Error, precision. See *Error, random*.

Error, probable. An error with a 50% or higher chance of occurrence. A statement of probable error is of little value.

Error, random. A statistical error caused by chance and not recurring. This term is a general category for errors that can take values on either side of an average value. To describe a random error, its distribution must be known.

Error, root mean square, or RMS. An accuracy statement of a system comprising several items. For example, a laboratory potentiometer, volt box, null detector, and reference voltage source have individual accuracy statements assigned to them. These errors are generally independent of one another, so a system of these units displays an accuracy given by the square root of the sum of the squares of the individual limits of error. For example, four individual errors of 0.1% yield a calibrated accuracy of 0.4% but an RMS error of only 0.2%.

Error, systematic. A persistent error not due to chance; systematic errors are causal. A systematic error is likely to have the same magnitude and sign for every instrument constructed with the same components and procedures. Errors in calibrating equipment cause systematic errors because all instruments calibrated are biased in the direction of the calibrating equipment error. Voltage and resistance drifts over time are generally in one direction and are classed as systematic errors.

Frequency response (flat). The portion of the frequency spectrum over which the measuring system has a constant value of amplitude response and a constant value of time lag. Input signals that have frequency components within this range are indicated by the measuring system (without distortion).

The preparation of this chapter is assigned to TC 1.2, Instruments and Measurements.

Hysteresis. The summation of all effects, under constant environmental conditions, that cause the output of an instrument to assume different values at a given stimulus point when that point is approached with increasing stimulus and with decreasing stimulus. Hysteresis includes backlash. It is usually measured as a percent of full scale when input varies over the full increasing and decreasing range. In instrumentation, hysteresis and dead band are similar quantities.

Linearity. The straight-lineness of the transfer curve between an input and an output; that condition prevailing when output is directly proportional to input (see *Nonlinearity*).

Loading error. A loss of output signal from a device caused by a current drawn from its output. It increases the voltage drop across the internal impedance, where no voltage drop is desired.

Mean. See *Average*.

Median. The middle value in a distribution, above and below which lie an equal number of values.

Noise. Any unwanted disturbance or spurious signal that modifies the transmission, measurement, or recording of desired data.

Nonlinearity. The prevailing condition (and the extent of its measurement) under which the input-output relationship (known as the input-output curve, transfer characteristic, calibration curve, or response curve) fails to be a straight line. Nonlinearity is measured and reported in several ways, and the way, along with the magnitude, must be stated in any specification.

Minimum-deviation-based nonlinearity. The maximum departure between the calibration curve and a straight line drawn to give the greatest accuracy; expressed as a percent of full-scale deflection.

Slope-based nonlinearity. The ratio of maximum slope error anywhere on the calibration curve to the slope of the nominal sensitivity line; usually expressed as a percent of nominal slope.

Most variations beyond these two definitions result from the many ways in which the straight line can be arbitrarily drawn. All are valid as long as construction of the straight line is explicit.

Population. A group of individual persons, objects, or items from which samples may be taken for statistical measurement.

Precision. The repeatability of measurements of the same quantity under the same conditions; not a measure of absolute accuracy. The precision of a measurement is used here to describe the relative tightness of the distribution of measurements of a quantity about their mean value. Therefore, precision of a measurement is associated more with its repeatability than its accuracy. It combines uncertainty caused by random differences in a number of identical measurements and the smallest readable increment of the scale or chart. Precision is given in terms of deviation from a mean value.

Primary calibration. A calibration procedure in which the instrument output is observed and recorded while the input stimulus is applied under precise conditions—usually from a primary external standard traceable directly to the National Institute of Standards and Technology (NIST).

Range. A statement of the upper and lower limits between which an instrument's input can be received and for which the instrument is calibrated.

Reliability. The probability that an instrument's precision and accuracy will continue to fall within specified limits.

Repeatability. See *Precision*.

Reproducibility. In instrumentation, the closeness of agreement among repeated measurements of the output for the same value of input made under the same operating conditions over a period of time, approaching from both directions; it is usually measured as a nonreproducibility and expressed as reproducibility in percent of span for a specified time period. Normally, this implies a long period of time, but under certain conditions, the period may be a short time so that drift is not included. Reproducibility includes hysteresis, dead band, drift, and repeatability. Between repeated measurements, the input may vary over the range, and operating conditions may vary within normal limits.

Resolution. The smallest change in input that produces a detectable change in instrument output. Resolution differs from precision in that it is a psychophysical term referring to the smallest increment of humanly perceptible output (rated in terms of the corresponding increment of input). The precision, the resolution, or both may be better than the accuracy. An ordinary six-digit (or dial) instrument has a resolution of one part per million (ppm) of full scale; however, it is possible that the accuracy is no better than 25 ppm (0.0025%). Note that the practical resolution of an instrument cannot be any better than the resolution of the indicator or detector, whether internal or external.

Scale factor. (1) The amount by which a measured quantity must change to produce unity output; (2) the ratio of real to analog values.

Sensitivity. The property of an instrument that determines scale factor. The word is often short for maximum sensitivity or the minimum scale factor with which an instrument can respond. The minimum input signal strength required to produce a desired value of output signal (e.g., full scale or unit output or the ratio of output to input values).

Sensitivity inaccuracy. The maximum error in sensitivity displayed as a result of the summation of the following: frequency response; attenuator inaccuracy; hysteresis or dead band; amplitude distortion (sensitivity nonlinearity); phase distortion (change in phase relationship between input signal and output deflection); and gain instability. Only by taking into account all these factors can nominal sensitivity, as indicated by the numeral on the attenuator readout, be discounted for accurate interpretation.

Stability. (1) Independence or freedom from changes in one quantity as the result of a change in another; (2) the absence of drift.

Static error band. (1) The spread of error present if the indicator (pen, needle) stopped at some value (e.g., at one-half of full scale). It is normally reported as a percent of full scale; (2) a specification or rating of maximum departure from the point where the indicator must be when an on-scale signal is stopped and held at a given signal level. This definition stipulates that the stopped position can be approached from either direction in following any random waveform. Therefore, it is a quantity that includes hysteresis and nonlinearity but excludes items such as chart paper accuracy or electrical drift (see *Dynamic error band*).

Step-function response. The characteristic curve or output plotted against time resulting from the input application of a step function (a function that is zero for all values of time before a certain instant, and a constant for all values of time thereafter).

Threshold. The smallest stimulus or signal that results in a detectable output.

Time constant. The time required for an exponential quantity to change by an amount equal to 0.632 times the total change required to reach steady state for first-order systems.

Transducer. A device for translating the changing magnitude of one kind of quantity into corresponding changes of another kind of quantity. The second quantity often has dimensions different from the first and serves as the source of a useful signal. The first quantity may be considered an input and the second an output. Significant energy may or may not transfer from the transducer's input to output.

Uncertainty. An estimated value for the error (i.e., what an error might be if it were measured by calibration). Although uncertainty may be the result of both systematic and precision errors, only precision error can be treated by statistical methods.

Zero shift. Drift in the zero indication of an instrument without any change in the measured variable.

UNCERTAINTY ANALYSIS

Uncertainty Sources

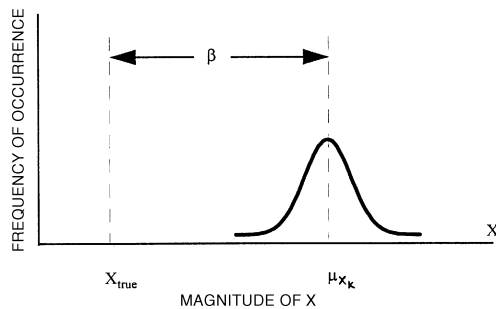
Measurement generally consists of a sequence of operations or steps. Virtually every step introduces a conceivable source of uncertainty, the effect of which must be assessed. The following list is representative of the most common, but not all, sources of uncertainty.

- Inaccuracy in the mathematical model that describes the physical quantity
- Inherent stochastic variability of the measurement process
- Uncertainties in measurement standards and calibrated instrumentation
- Time-dependent instabilities due to gradual changes in standards and instrumentation
- Effects of environmental factors such as temperature, humidity, and pressure
- Values of constants and other parameters obtained from outside sources
- Uncertainties arising from interferences, impurities, inhomogeneity, inadequate resolution, and incomplete discrimination
- Computational uncertainties and data analysis
- Incorrect specifications and procedural errors
- Laboratory practice, including handling techniques, cleanliness, and operator techniques, etc.
- Uncertainty in corrections made for known effects, such as installation effect corrections

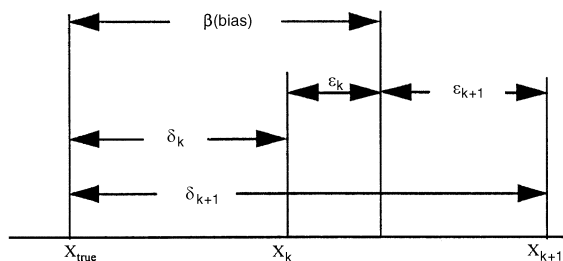
Uncertainty of a Measured Variable

For a measured variable X_i , the total error is caused by both **precision (random)** and **systematic (bias) errors**. This relationship is shown in Figure 1. The possible measurement values of the variable are scattered in a distribution around the parent population mean μ_i (Figure 1A). The curve is the normal or Gaussian distribution and is the theoretical distribution function for the infinite population of measurements that generated X_i . The parent population mean differs from $(X_i)_{true}$ by an amount called the systematic (or bias) error β_i (Figure 1B). The quantity β_i is the total fixed error that remains after all calibration corrections have been made. In general, there are several sources of bias error, such as calibration standard errors, data acquisition errors, data reduction errors, and test technique errors. There is usually no direct way to measure these errors. These errors are unknown and are assumed to be zero; otherwise, an additional correction would be applied to reduce them to as close to zero as possible.

The **precision uncertainty** for a variable, which is an estimate of the possible error associated with the repeatability of a particular



A. INFINITE NUMBER OF READINGS



B. TWO READINGS

Fig. 1 Errors in the Measurement of a Variable X

measurement, is determined from the sample standard deviation, or the estimate of the error associated with the repeatability of a particular measurement. Unlike the systematic error, the precision error varies from reading to reading. As the number of readings of a particular variable tends to infinity, the distribution of these possible errors becomes Gaussian.

For each bias error source, the experimenter must estimate a **systematic uncertainty**. Systematic uncertainties are usually estimated from previous experience, calibration data, analytical models, and engineering judgment. For a discussion on estimating systematic uncertainties (bias limits), see Coleman and Steele (1989).

For further information on measurement uncertainty, see ASME Standards MFC-2M and PTC 19.1 and Coleman and Steele (1995).

TEMPERATURE MEASUREMENT

Instruments for measuring temperature are listed in Table 1. Temperature sensor output must be related to an accepted temperature scale. This is achieved by manufacturing the instrument according to certain specifications or by calibrating it against a temperature standard. To help users conform to standard temperatures and temperature measurements, the International Committee of Weights and Measures (CIPM) has adopted the International Temperature Scale of 1990 (ITS-90).

The unit of temperature of the ITS-90 is the kelvin (K) and has a size equal to the fraction 1/273.16 of the thermodynamic temperature of the triple point of water.

The ITS-90 is maintained in the United States by the National Institute of Standards and Technology (NIST), and any laboratory may obtain calibrations from NIST based on this scale.

Benedict (1984), Considine (1985), Quinn (1990), Schooley (1986, 1992), and DeWitt and Nutter (1988) cover temperature measurement in more detail.

Sampling and Averaging

Although temperature is usually measured within, and is associated with, a relatively small volume (depending on the size of the thermometer), it can also be associated with an area (e.g., on a surface or in a flowing stream). To determine average stream temperature, the cross section must be divided into smaller areas and the temperature of each area measured. The temperatures measured are then combined into a weighted mass flow average by either (1) using equal areas and multiplying each temperature by the fraction of total mass flow in its area or (2) using areas of size inversely proportional to mass flow and taking a simple arithmetic average of the temperatures in each. A means of mixing or selective sampling may be preferable to these cumbersome procedures. While mixing can occur from turbulence alone, **transposition** is much more effective. In transposition, the stream is divided into parts determined by the type of stratification, and alternate parts pass through one another.

Static Temperature Versus Total Temperature

When a fluid stream impinges on a temperature-sensing element such as a thermometer or thermocouple, the element is at a temperature greater than the true stream temperature. The difference is a fraction of the temperature equivalent of the stream velocity t_e .

$$t_e = \frac{3600 V^2}{2g_c J c_p} \tag{1}$$

where

t_e = temperature equivalent of stream velocity, °F

V = velocity of stream, fpm

g_c = gravitational constant = 32.174 lb_m · ft/lb_f · s²

J = mechanical equivalent of heat, 778.3 ft · lb_f/Btu

c_p = specific heat of stream at constant pressure, Btu/lb_m · °F

Table 1 Temperature Measurement

Measurement Means	Application	Approximate Range, °F	Uncertainty, °F	Limitations
Liquid-in-glass thermometers				
Mercury-in-glass	Temperature of gases and liquids by contact	-36/1000	0.05 to 3.6	In gases, accuracy affected by radiation
Organic	Temperature of gases and liquids by contact	-330/400	0.05 to 3.6	In gases, accuracy affected by radiation
Gas thermometer	Primary standard	-456/1200	Less than 0.02	Requires considerable skill to use
Resistance thermometers				
Platinum	Precision; remote readings; temperature of fluids or solids by contact	-430/1800	Less than 0.0002 to 0.2	High cost; accuracy affected by radiation in gases
Rhodium-iron	Transfer standard for cryogenic applications	-460/-400	0.0002 to 0.2	High cost
Nickel	Remote readings; temperature by contact	-420/400	0.02 to 2	Accuracy affected by radiation in gases
Germanium	Remote readings; temperature by contact	-460/-400	0.0002 to 0.2	
Thermistors	Remote readings; temperature by contact	Up to 400	0.0002 to 0.2	
Thermocouples				
Pt-Rh/Pt (type S)	Standard for thermocouples on IPTS-68, not on ITS-90	32/2650	0.2 to 5	High cost
Au/Pt	Highly accurate reference thermometer for laboratory applications	-60/1800	0.1 to 2	High cost
Types K and N	General testing of high temperature; remote rapid readings by direct contact	Up to 2300	0.2 to 18	Less accurate than listed above thermocouples
Iron/Constantan (type J)	Same as above	Up to 1400	0.2 to 10	Subject to oxidation
Copper/Constantan (type T)	Same as above, especially suited for low temperature	Up to 660	0.2 to 5	
Ni-Cr/Constantan (type E)	Same as above, especially suited for low temperature	Up to 1650	0.2 to 13	
Beckman thermometers (metastatic)	For differential temperature in same applications as in glass-stem thermometer	10°F scale, used 32 to 212°F	0.01	Must be set for temperature to be measured
Bimetallic thermometers	For approximate temperature	-4/1200	2, usually much more	Time lag; unsuitable for remote use
Pressure-bulb thermometers				
Gas-filled bulb	Remote testing	-100/1200	4	Caution must be exercised so that installation is correct
Vapor-filled bulb	Remote testing	-25/500	4	Caution must be exercised so that installation is correct
Liquid-filled bulb	Remote testing	-60/2100	4	Caution must be exercised so that installation is correct
Optical pyrometers	For intensity of narrow spectral band of high-temperature radiation (remote)	1500 and up	30	
Radiation pyrometers	For intensity of total high-temperature radiation (remote)	Any range		
Seeger cones (fusion pyrometers)	Approximate temperature (within temperature source)	1200/3600	90	
Triple points, freezing/melting points, and boiling points of materials	Standards	All except extremely high temperatures	Extremely precise	For laboratory use only

This fraction of the temperature equivalent of the velocity is the **recovery factor**, and it varies from 0.6 to 0.8°F for bare thermometers to 1.0°F for aerodynamically shielded thermocouples. For precise temperature measurement, each temperature sensor must be calibrated to determine its recovery factor. However, for most applications where air velocities are below 2000 fpm, the recovery factor can be omitted.

Various temperature sensors are available for temperature measurement in fluid streams. The principal sensors are the **static temperature thermometer**, which indicates true stream temperature but is cumbersome, and the **thermistor**, used for accurate temperature measurement within a limited range.

LIQUID-IN-GLASS THERMOMETERS

Any device that changes monotonically with temperature is a thermometer; however, the term usually signifies an ordinary liquid-in-glass temperature-indicating device. Mercury-filled thermometers have a useful range from -37.8°F , the freezing point of mercury, to about 1000°F , near which the glass usually softens. Lower temperatures can be measured with organic-liquid-filled thermometers (e.g., alcohol-filled), with ranges of -330 to 400°F . During manufacture, thermometers are roughly calibrated for at least two temperatures, often the freezing and boiling points of water; space between the calibration points is divided into desired scale divisions. Thermometers that are intended for precise measurement applications have scales etched into the glass that forms their stems. The probable error for as-manufactured, etched-stem thermometers is ± 1 scale division. The highest quality mercury thermometers may have uncertainties of ± 0.06 to $\pm 4^{\circ}\text{F}$ if they have been calibrated by comparison against primary reference standards.

Liquid-in-glass thermometers are used for many applications within the HVAC industry. Some of these uses include local temperature indication of process fluids related to HVAC systems, such as cooling and heating fluids and air.

The use of mercury-in-glass thermometers as temperature measurement standards is fairly common because of their relatively high accuracy and low cost. Such thermometers used as references must be calibrated on the ITS-90 by comparison in a uniform bath with a standard platinum resistance thermometer that has been calibrated either by the appropriate standards agency or by a laboratory that has direct traceability to the standards agency and the ITS-90. Such a calibration is necessary in order to determine the proper corrections to be applied to the scale readings. For application and calibration of liquid-in-glass thermometers, refer to NIST (1976, 1986).

Liquid-in-glass thermometers are calibrated by the manufacturer for total or partial stem immersion. If a thermometer calibrated for total immersion is used at partial immersion (i.e., with a portion of the liquid column at a temperature different from that of the bath), an emergent stem correction must be made. This correction can be calculated as follows:

$$\text{Stem correction} = Kn(t_b - t_s) \quad (2)$$

where

K = differential expansion coefficient of mercury or other liquid in glass. K is 0.00009 for Fahrenheit mercurial thermometers. For K values for other liquids and specific glasses, refer to Schooley (1992).

n = number of degrees that liquid column emerges from bath

t_b = temperature of bath, $^{\circ}\text{F}$

t_s = average temperature of emergent liquid column of n degrees, $^{\circ}\text{F}$

Sources of Thermometer Errors

A thermometer measuring gas temperatures can be affected by radiation from surrounding surfaces. If the gas temperature is

approximately the same as that of the surrounding surfaces, radiation effects can be ignored. If the temperature differs considerably from that of the surroundings, radiation effects should be minimized by shielding or aspiration (ASME *Standard* PTC 193). **Shielding** may be provided by highly reflective surfaces placed between the thermometer bulb and the surrounding surfaces such that air movement around the bulb is not appreciably restricted (Parmelee and Huebscher 1946). Improper shielding can increase errors. **Aspiration** results from passing a high-velocity stream of air or gas over the thermometer bulb.

When a **thermometer well** within a container or pipe under pressure is required, the thermometer should fit snugly and be surrounded with a high thermal conductivity material (oil, water, or mercury, if suitable). Liquid in a long, thin-walled well is advantageous for rapid response to temperature changes. The surface of the pipe or container around the well should be insulated to eliminate heat transfer to or from the well.

Industrial thermometers are available for permanent installation in pipes or ducts. These instruments are fitted with metal guards to prevent breakage and are useful for many other purposes. The considerable heat capacity and conductance of the guards or shields can cause errors, however.

Allowing ample time for the thermometer to attain temperature equilibrium with the surrounding fluid prevents excessive errors in temperature measurements. When reading a liquid-in-glass thermometer, the eye should be kept at the same level as the top of the liquid column to avoid parallax.

RESISTANCE THERMOMETERS

Resistance thermometers depend on a change of the electrical resistance of a sensing element (usually metal) with a change in temperature; resistance increases with increasing temperature. The use of resistance thermometers largely parallels that of thermocouples, although readings are usually unstable above about 1000°F . Two-lead temperature elements are not recommended because they do not permit correction for lead resistance. Three leads to each resistor are necessary to obtain consistent readings, and four leads are preferred. Wheatstone bridge circuits or 6-1/2-digit multimeters can be used for measurements.

A typical circuit used by several manufacturers is shown in [Figure 2](#). In this design, a differential galvanometer is used in which coils L and H exert opposing forces on the indicating needle. Coil L is in series with the thermometer resistance AB, and coil H is in series with the constant resistance R. As the temperature falls, the resistance of AB decreases, allowing more current to flow through coil L than through coil H. This causes an increase in the force exerted by coil L, pulling the needle down to a lower reading. Likewise, as the temperature rises, the resistance of AB increases, causing less current to flow through coil L than through coil H. This forces the indicating needle to a higher reading. Rheostat S must be adjusted occasionally to maintain a constant current.

The resistance thermometer is more costly to make and likely to have considerably longer response times than thermocouples. A resistance thermometer gives best results when used to measure steady or slowly changing temperature.

Resistance Temperature Devices

Resistance temperature devices (RTDs) are typically constructed from platinum, rhodium-iron, nickel, nickel-iron, tungsten, or copper. These devices are further characterized by their simple circuit designs, high degree of linearity, good sensitivity, and excellent stability. The choice of materials for an RTD usually depends on the intended application; temperature range, corrosion protection, mechanical stability, and cost are some of the selection criteria.

Platinum RTDs. Presently, for HVAC applications, RTDs constructed of platinum are the most widely used. Platinum is extremely

stable and corrosion-resistant. Platinum RTDs are highly malleable and can thus be drawn into fine wires; they can also be manufactured at low cost as thin films. They have a high melting point and can be refined to a high degree of purity, thus attaining highly reproducible results. Due to these properties, platinum RTDs are used to define the ITS-90 for the range of 13.8033 K (triple point of equilibrium hydrogen) to 1234.93 K (freezing point of silver).

Platinum resistance temperature devices can measure the widest range of temperatures and are the most accurate and stable temperature sensors. Their resistance-temperature relationship is one of the most linear. The higher the purity of the platinum, the more stable and accurate the sensor. With high-purity platinum, primary grade platinum RTDs are capable of achieving reproducibility of $\pm 0.00002^\circ\text{F}$, whereas the minimum uncertainty of a recently calibrated thermocouple is $\pm 0.4^\circ\text{F}$.

Platinum RTD Design. The most widely used RTD is designed with a resistance of $100\ \Omega$ at 32°F ($R_0 = 100\ \Omega$). Other RTDs are

available that use lower resistances at temperatures above 1100°F . The lower the resistance value, the faster the response time for sensors of the same size.

Thin-Film Platinum RTDs. Thin-film $1000\ \Omega$ platinum RTDs are readily available. They have the excellent linear properties of lower resistance platinum RTDs and are more cost-effective because they are mass produced and have lower platinum purity. However, the problem with many platinum RTDs with R_0 values of greater than $100\ \Omega$ is the difficulty in obtaining transmitters or electronic interface boards from sources other than the RTD manufacturer. In addition to a nonstandard interface, higher R_0 value platinum RTDs may have higher self-heating losses if the excitation current is not controlled properly.

Thin-film RTDs have the advantages of lower cost and smaller sensor size. They are specifically adapted to surface mounting. Thin-film RTDs tend to have an accuracy limitation of $\pm 0.1\%$ or $\pm 0.2^\circ\text{F}$. This may prove to be adequate for most HVAC applications; only in tightly controlled facilities may users wish to install the standard wire-wound platinum RTDs with accuracies of 0.01% or $\pm 0.02^\circ\text{F}$ (these are available upon special request for certain temperature ranges).

Assembly and Construction. Regardless of the R_0 resistance value of RTDs, their assembly and construction are relatively simple. The electrical connections come in three basic types, depending on the number of wires to be connected to the resistance measurement circuitry. Two, three, or four wires are used for electrical connection using a Wheatstone bridge or a variation of it (Figure 3).

In the basic two-wire configuration, the resistance of the RTD is measured through the two connecting wires. Because the connecting wires extend from the site of the temperature measurement, any additional changes in resistivity due to a change in temperature may affect the measured resistance. Three- and four-wire assemblies are built to compensate for the connecting lead resistance values. The original three-wire circuit improved the resistance measurement by adding a compensating wire to the voltage side of the circuit. This helps reduce part of the connecting wire resistance. When more accurate measurements (better than $\pm 0.2^\circ\text{F}$) are required, the four-wire bridge is recommended. The four-wire bridge eliminates all connecting wire resistance errors.

All the bridges discussed here are direct current (dc) circuits and were used extensively until the advent of precision alternating current (ac) circuits using microprocessor-controlled ratio transformers, dedicated analog-to-digital converters, and other solid-state

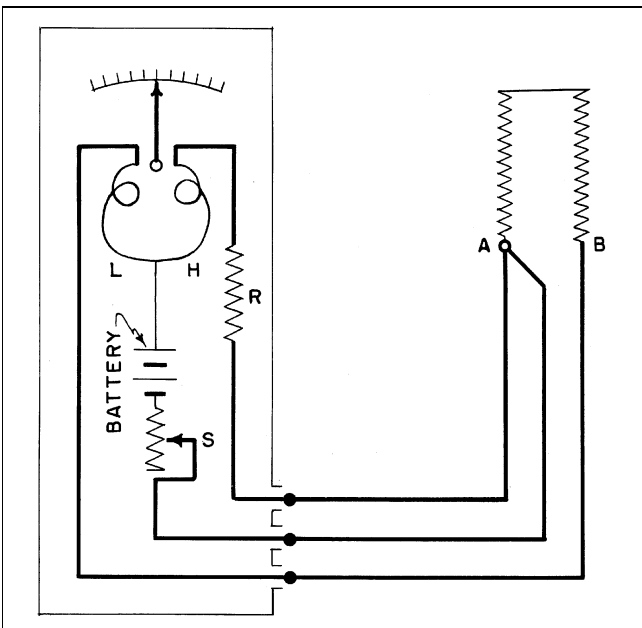


Fig. 2 Typical Resistance Thermometer Circuit

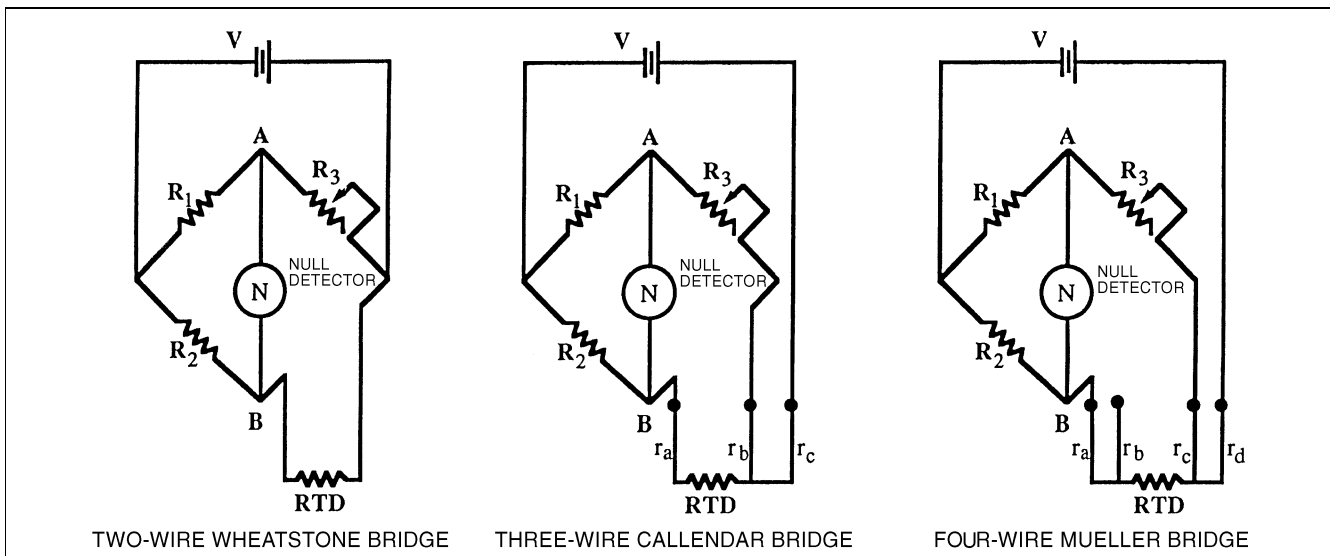


Fig. 3 Typical Resistance Temperature Device Bridge Circuits

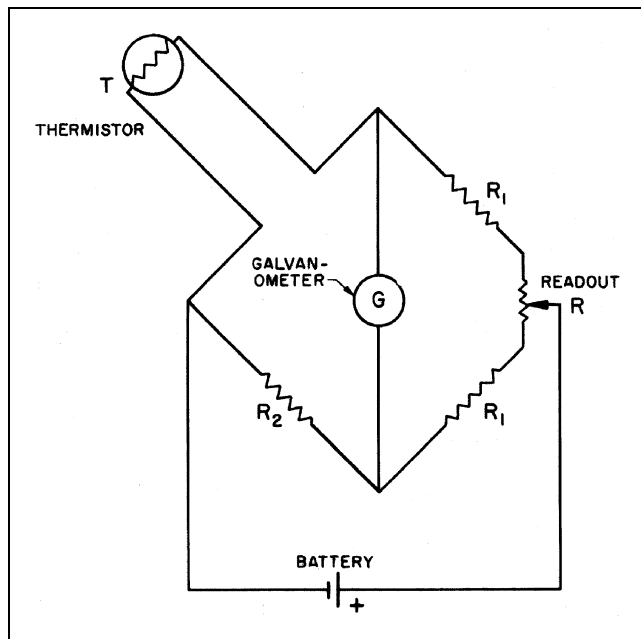


Fig. 4 Basic Thermistor Circuit

devices that measure resistance with uncertainties of less than 1 ppm. Resistance measurement technology now allows more portable thermometers, lower cost, ease of use, and high-precision temperature measurement in industrial uses.

Thermistors

Certain semiconductor compounds (usually sintered metallic oxides) exhibit large changes in resistance with temperature, usually decreasing as the temperature increases. For use, the thermistor element may be connected by lead wires into a galvanometer bridge circuit and calibrated. Alternatively, a 6-1/2-digit multimeter and a constant-current source with a means for reversing the current to eliminate thermal electromotive force (emf) effects may also be used. This method of measurement is easier and faster, and it may be more precise and accurate. Thermistors are usually applied to electronic temperature compensation circuits, such as thermocouple reference junction compensation, or to other applications where high resolution and limited operating temperature ranges exist. [Figure 4](#) illustrates a typical thermistor circuit.

Semiconductor Devices

In addition to the positive resistance coefficient RTDs and the negative resistance coefficient thermistor, there are two other types of devices that vary resistance or impedance with temperature. Although the principle of their operation has long been known, their reliability was questioned due to imprecise manufacturing techniques. Improvements in silicon microelectronics manufacturing techniques have brought semiconductors to the point where low-cost, precise temperature sensors are commercially available.

Elemental Semiconductors. Due to controlled doping of impurities into elemental germanium, a germanium semiconductor is a reliable temperature sensor for cryogenic temperature measurement in the range of 1.8 to 150°R.

Junction Semiconductors. The first simple junction semiconductor device consisted of a single diode or transistor, in which the forward-connected base emitter voltage is very sensitive to temperature. Today the more common form is a pair of diode-connected transistors, which make the device suitable for ambient temperature measurement. Applications include thermocouple reference junction compensation.

The primary advantages of silicon transistor temperature sensors are their extreme linearity and exact R_0 value. Another advantage is the incorporation of signal conditioning circuitry into the same device as the sensor element. As with thermocouples, these semiconductors require highly precise manufacturing techniques, extremely precise voltage measurements, multiple-point calibration, and temperature compensation to achieve an accuracy as high as $\pm 0.02^\circ\text{F}$, but with a much higher cost. Lower cost devices achieve accuracies of $\pm 0.2^\circ\text{F}$ using mass manufacturing techniques and single-point calibration. A mass-produced silicon temperature sensor can be interchanged easily. If one device fails, only the sensor element need be changed. Electronic circuitry can be used to recalibrate the new device.

Winding Temperature. The winding temperature of electrical operating equipment is usually determined from the resistance change of these windings in operation. With copper windings, the relation between these parameters is

$$\frac{R_1}{R_2} = \frac{148 + t_1}{148 + t_2} \quad (3)$$

where

$$\begin{aligned} R_1 &= \text{winding resistance at temperature } t_1, \Omega \\ R_2 &= \text{winding resistance at temperature } t_2, \Omega \\ t_1, t_2 &= \text{winding temperatures, } ^\circ\text{F} \end{aligned}$$

The classical method of determining winding temperature is to measure the equipment when it is inoperative and temperature-stabilized at room temperature. After the equipment has operated sufficiently to cause temperature stabilization under load conditions, the winding resistance should be measured again. The latter value is obtained by taking resistance measurements at known short time intervals after shutdown. These values may be extrapolated to zero time to indicate the winding resistance at the time of shutdown. The obvious disadvantage of this method is that the device must be shut down to determine winding temperature. A circuit described by Seely (1955), however, makes it possible to measure resistances while the device is operating.

THERMOCOUPLES

When two wires of dissimilar metals are joined by soldering, welding, or twisting, they form a thermocouple junction or **thermo-junction**. An emf that depends on the wire materials and the junction temperature exists between the wires. This is known as the **Seebeck voltage**.

Thermocouples for temperature measurement yield less precise results than platinum resistance thermometers, but except for glass thermometers, thermocouples are the most common instruments of temperature measurement for the range of 32 to 1800°F. Due to their low cost, moderate reliability, and ease of use, thermocouples continue to maintain widespread acceptance.

The most commonly used thermocouples in industrial applications are assigned letter designations. The tolerances of such commercially available thermocouples are given in [Table 2](#).

Because the measured emf is a function of the difference in temperature and the type of dissimilar metals used, a known temperature at one junction is required, whereas the remaining junction temperature may be calculated. It is common practice to call the one with a known temperature the (cold) **reference** junction and the one with the unknown temperature the (hot) **measured** junction. The reference junction is typically kept at a reproducible temperature, such as the ice point of water.

Various systems are used to maintain the reference junction temperature—a mixture of ice and water contained in an insulated flask or commercially available thermoelectric coolers to maintain the ice point temperature automatically within a reference chamber. When

Table 2 Thermocouple Tolerances on Initial Values of Electromotive Force Versus Temperature

Thermocouple Type	Material Identification	Temperature Range, °F	Reference Junction Tolerance at 32°F ^a	
			Standard Tolerance (whichever is greater)	Special Tolerance (whichever is greater)
T	Copper versus Constantan	32 to 700	±1.8°F or ±0.75%	±0.9°F or ±0.4%
J	Iron versus Constantan	32 to 1400	±4°F or ±0.75%	±2°F or ±0.4%
E	Nickel-10% Chromium versus Constantan	32 to 1600	±3.1°F or ±0.5%	±1.8°F or ±0.4%
K	Nickel-10% Chromium versus 5% Aluminum, Silicon	32 to 2300	±4°F or ±0.75%	±2°F or ±0.4%
N	Nickel-14% Chromium, 1.5% Silicon versus Nickel-4.5% Silicon, 0.1% Magnesium	32 to 2300	±4°F or ±0.75%	±2°F or ±0.4%
R	Platinum-13% Rhodium versus Platinum	32 to 2700	±2.7°F or ±0.25%	±1.1°F or ±0.1%
S	Platinum-10% Rhodium versus Platinum	32 to 2700	±2.7°F or ±0.25%	±1.1°F or ±0.1%
B	Platinum-30% Rhodium versus Platinum-6% Rhodium	1600 to 3100	±0.5%	±0.25%
T ^b	Copper versus Constantan	-328 to 32	±1.8°F or ±1.5%	c
E ^b	Nickel-10% Chromium versus Constantan	-328 to 32	±3.1°F or ±1%	c
K ^b	Nickel-10% Chromium versus 5% Aluminum, Silicon	-328 to 32	±4°F or ±2%	c

Source: ASTM Standard E 230, Temperature-Electromotive Force (EMF) Tables for Standardized Thermocouples.

^aTolerances in this table apply to new thermocouple wire, normally in the size range of 0.01 to 0.1 in. diameter and used at temperatures not exceeding the recommended limits. Thermocouple wire is available in two grades: standard and special.

^bThermocouples and thermocouple materials are normally supplied to meet the tolerance specified in the table for temperatures above 32°F. The same materials, however, may not fall within the tolerances given in the second section of the table when operated below freezing (32°F). If materials are required to meet tolerances at subfreezing temperatures, the purchase order must state so.

^cLittle information is available to justify establishing special tolerances for below-freezing temperatures. Limited experience suggests the following special tolerances for types E and T thermocouples:

Type E -328 to 32°F; ±2°F or ±0.5% (whichever is greater)

Type T -328 to 32°F; ±1°F or ±0.8% (whichever is greater)

These tolerances are given only as a guide for discussion between purchaser and supplier.

these systems cannot be used in an application, measuring instruments with automatic reference junction temperature compensation may be used.

As previously described, the principle for measuring temperature with a thermocouple is based on the accurate measurement of the Seebeck voltage. The acceptable dc voltage measurement methods are (1) millivoltmeter, (2) millivolt potentiometer, and (3) a high-input impedance digital voltmeter. Many digital voltmeters include built-in software routines for the direct calculation and display of temperature. Regardless of the method selected, many options to simplify the measurement process are available.

Solid-state digital readout devices in combination with a millivolt- or microvoltmeter, as well as packaged thermocouple readouts with built-in cold junction and linearization circuits, are available. The latter requires a proper thermocouple to provide direct meter reading of temperature. Accuracy approaching or surpassing that of potentiometers can be attained, depending on the instrument quality. This method is popular because it eliminates the null balancing requirement and reads temperature directly in a digital readout.

Wire Diameter and Composition

Thermocouple wire is selected by considering the temperature to be measured, the corrosion protection afforded to the thermocouple, and the precision and service life required. Type T thermocouples are suitable for temperatures up to 700°F; type J, up to 1400°F; and types K and N, up to 2300°F. Higher temperatures require noble metal thermocouples (type S, R, or B), which have a higher initial cost and do not develop as high an emf as the base metal thermocouples. Thermocouple wires of the same type have small compositional variation from lot to lot from the same manufacturer and especially among different manufacturers. Consequently, calibrating samples from each wire spool is essential for precision. Calibration data on wire may be obtained from the manufacturer.

Reference functions are available for relating temperature and emf of letter-designated thermocouple types. Such functions are easy to use with computers. The functions depend on thermocouple type and on the range of temperature; they are used to generate

reference tables of emf as a function of temperature but are not well suited for calculating temperatures directly from values of emf. Approximate inverse functions are available, however, for calculating temperature and are of the form

$$t = \sum_{i=0}^n a_i E^i \quad (4)$$

where t = temperature, a = thermocouple constant, and E = voltage. Burns et al. (1992) give the reference functions and approximate inverses for all letter-designated thermocouples.

The emf of a thermocouple, as measured with a high-input impedance device, is independent of the diameters of its constituent wires. Thermocouples with small-diameter wires respond faster to temperature changes and are less affected by radiation than larger ones. Large-diameter wire thermocouples, however, are necessary for high-temperature work when wire corrosion is a problem. For use in heated air or gases, thermocouples are often shielded and sometimes aspirated. An arrangement for avoiding error due to radiation involves using several thermocouples of different wire sizes and estimating the true temperature by extrapolating readings to zero diameter.

With thermocouples, temperatures can be indicated or recorded remotely on conveniently located instruments. Because thermocouples can be made of small-diameter wire, they can be used to measure temperatures within thin materials, within narrow spaces, or in otherwise inaccessible locations.

Multiple Thermocouples

Thermocouples in series, with alternate junctions maintained at a common temperature, produce an emf that, when divided by the number of thermocouples, gives the average emf corresponding to the temperature difference between two sets of junctions. This series arrangement of thermocouples, often called a **thermopile**, is used to increase sensitivity and is often used for measuring small temperature changes and differences.

Connecting a number of thermocouples of the same type in parallel with a common reference junction is useful for obtaining an average temperature of an object or volume. In such measurements, however, it is important that the electrical resistances of the individual thermocouples be the same. The use of thermocouples in series and parallel arrangements is discussed in ASTM *Manual 12*.

Surface Temperature Measurement

The thermocouple is useful in determining surface temperature. It can be attached to a metal surface in several ways. For permanent installations, soldering, brazing, or peening is suggested. For peening, a small hole is drilled and the thermocouple measuring junction is driven into it. For temporary arrangements, thermocouples can be attached by tape, adhesive, or putty-like material. For boiler or furnace surfaces, furnace cement should be used. To minimize the possibility of error caused by heat conduction along wires, a surface thermocouple should be made of fine wires placed in close contact with the surface being measured for about an inch from the junction to ensure good thermal contact. The wires must be insulated electrically from each other and from the metal surface (except at the junction).

Thermocouple Construction

The thermocouple wires are typically insulated with fibrous glass, fluorocarbon resin, or ceramic insulators. In another form of thermocouple, the thermocouple wires are insulated with compacted ceramic insulation inside a metal sheath. This form of thermocouple provides both mechanical protection and protection from stray electromagnetic fields. The measuring junction may be exposed or enclosed within the metal sheath. An enclosed junction may be either grounded or ungrounded to the metal sheath.

For the exposed junction type, the measuring junction is in direct contact with the process stream; it is therefore subject to corrosion or contamination but provides a fast temperature response. The grounded enclosed junction type, in which the thermocouple wires are welded to the metal sheath, provides electrical grounding, as well as mechanical and corrosion protection. This type, however, has a slower response time than the exposed junction type. With the ungrounded enclosed junction construction, the response time is even slower, but the thermocouple wires are isolated electrically and are less susceptible to some forms of mechanical strain than those with grounded construction.

INFRARED RADIOMETERS

Infrared radiation thermometers, also known as remote temperature sensors (Hudson 1969), permit noncontact measurement of surface temperature over a wide range. In these instruments, radiant flux from the observed object is focused by an optical system onto an infrared detector that generates an output signal proportional to the incident radiation that can be read from a meter or display unit. Point and scanning radiometers are available; the latter are able to display the temperature variation existing within the field of view.

Radiometers are usually classified according to the detector used—either thermal or photon. In thermal detectors, a change in electrical property is caused by the heating effect of the incident radiation. Examples of thermal detectors are the thermocouple, the thermopile, and metallic and semiconductor bolometers. In photon detectors, a change in electrical property is caused by the surface absorption of incident photons. Because these detectors do not require an increase in temperature for activation, their response time is much shorter than that of thermal detectors. Scanning radiometers usually use photon detectors.

A radiometer only measures the power level of the radiation incident on the detector; this incident radiation is a combination of the thermal radiation emitted by the object and the surrounding background radiation reflected from the surface of the object. An accurate measurement of the temperature, therefore, requires knowledge

of the long-wavelength emissivity of the object as well as the effective temperature of the thermal radiation field surrounding the object. Calibration against an internal or external source of known temperature and emissivity is required in order to obtain true surface temperature from the radiation measurements.

The temperature resolution of a radiometer decreases as the object temperature decreases. For example, a radiometer that can resolve a temperature difference of 0.5°F on an object near 70°F may only resolve a difference of 2°F on an object at 32°F.

INFRARED THERMOGRAPHY

Infrared thermography is the discipline concerned with the acquisition and analysis of thermal information in the form of images from an infrared imaging system. An infrared imaging system consists of (1) an infrared television camera and (2) a display unit. The infrared television camera scans a surface and senses the self-emitted and reflected radiation viewed from the surface. The display unit contains either a cathode-ray tube (CRT) that displays a gray-tone or color-coded thermal image of the surface or a color liquid crystal display (LCD) screen. A photograph of the image on the CRT is called a **thermogram**. An introductory treatise on infrared thermography is given by Paljak and Pettersson (1972).

Thermography has been used successfully to detect missing insulation and air infiltration paths in building envelopes (Burch and Hunt 1978). Standard practices for conducting thermographic inspections of buildings are given in ASTM *Standard C 1060*. A technique for quantitatively mapping the heat loss in building envelopes is given by Mack (1986).

Aerial infrared thermography of buildings is effective in identifying regions of an individual built-up roof that have wet insulation (Tobiasson and Korhonen 1985), but it is ineffective in ranking a group of roofs according to their thermal resistance (Goldstein 1978, Burch 1980). In this latter application, the emittances of the separate roofs and outdoor climate (i.e., temperature and wind speed) throughout the microclimate often produce changes in the thermal image that may be incorrectly attributed to differences in thermal resistance.

Industrial applications include locating defective or missing pipe insulation in buried heat distribution systems, surveys of manufacturing plants to quantify energy loss from equipment, and locating defects in coatings (Bentz and Martin 1987).

HUMIDITY MEASUREMENT

Any instrument capable of measuring the humidity or psychrometric state of air is a hygrometer, and many are available. The indication sensors used on the instruments respond to different moisture property contents. These responses are related to factors such as wet-bulb temperature, relative humidity, humidity (mixing) ratio, dew point, and frost point.

[Table 3](#) lists instruments for measuring humidity. Each is capable of accurate measurement under certain conditions and within specific limitations. The following sections describe various instruments used to measure humidity.

PSYCHROMETERS

A typical industrial psychrometer consists of a pair of matched electrical or mechanical temperature sensors, one of which is kept wet with a moistened wick. A blower aspirates the sensor, which lowers the temperature at the moistened temperature sensor. The lowest temperature depression occurs when the evaporation rate required to saturate the moist air adjacent to the wick is constant. This is a steady-state, open-loop, nonequilibrium process, which depends on the purity of the water, the cleanliness of the wick, the ventilation rate, radiation effects, the size and accuracy of the temperature sensors, and the transport properties of the gas.

Table 3 Humidity Sensor Properties

Type of Sensor	Sensor Category	Method of Operation	Approximate Range	Some Uses	Approximate Accuracy
Psychrometer	Evaporative cooling	Temperature measurement of wet bulb	32 to 180°F	Measurement, standard	±3 to ±7% rh
Adiabatic saturation psychrometer	Evaporative cooling	Temperature measurement of thermodynamic wet bulb	40 to 85°F	Measurement, standard	±0.2 to ±2% rh
Chilled mirror	Dew point	Optical determination of moisture formation	-110 to 200°F dp	Measurement, control, meteorology	±0.4 to ±4 °F
Heated saturated salt solution	Water vapor pressure	Vapor pressure depression in salt solution	-20 to 160°F dp	Measurement, control, meteorology	±3°F
Hair	Mechanical	Dimensional change	5 to 100% rh	Measurement, control	±5% rh
Nylon	Mechanical	Dimensional change	5 to 100% rh	Measurement, control	±5% rh
Dacron thread	Mechanical	Dimensional change	5 to 100% rh	Measurement	±7% rh
Goldbeater's skin	Mechanical	Dimensional change	5 to 100% rh	Measurement	±7% rh
Cellulosic materials	Mechanical	Dimensional change	5 to 100% rh	Measurement, control	±5% rh
Carbon	Mechanical	Dimensional change	5 to 100% rh	Measurement	±5% rh
Dunmore type	Electrical	Impedance	7 to 98% rh at 40 to 140°F	Measurement, control	±1.5% rh
Ion exchange resin	Electrical	Impedance or capacitance	10 to 100% rh at -40 to 190°F	Measurement, control	±5% rh
Porous ceramic	Electrical	Impedance or capacitance	Up to 400°F	Measurement, control	±1 to ±1.5% rh
Aluminum oxide	Electrical	Capacitance	5 to 100% rh	Measurement, control	±3% rh
Aluminum oxide	Electrical	Capacitance	-110 to 140°F dp	Trace moisture measurement, control	±2°F dp
Electrolytic hygrometer	Electrical	Capacitance			
Coulometric	Electrolytic cell	Electrolyzes due to adsorbed moisture	1 to 1000 ppm	Measurement	
Infrared laser diode	Electrical	Optical diodes	0.1 to 100 ppm	Trace moisture measurement	±0.1 ppm
Surface acoustic wave	Electrical	SAW attenuation	85 to 98% rh	Measurement, control	±1% rh
Piezoelectric	Mass sensitive	Mass changes due to adsorbed moisture	-100 to 0°F	Trace moisture measurement, control	±2 to ±10°F dp
Radiation absorption	Moisture absorption	Moisture absorption of UV or IR radiation	0 to 180°F dp	Measurement, control, meteorology	±4°F dp, ±5% rh
Gravimetric	Direct measurement of mixing ratio	Comparison of sample gas with dry airstream	120 to 20,000 ppm mixing ratio	Primary standard, research and laboratory	±0.13% of reading
Color change	Physical	Color changes	10 to 80% rh	Warning device	±10% rh

Notes:

1. This table does not include all available technology for humidity measurement.
2. The approximate range for the device types listed is based on surveys of device manufacturers.

3. The approximate accuracy is based on manufacturers' data.
4. Presently, NIST only certifies instruments whose operating range is within -103 to 212°F dew point.

ASHRAE *Standard* 41.6 recommends an airflow over both the wet and dry bulbs of 600 to 1000 fpm for transverse ventilation and 300 to 500 fpm for axial ventilation.

The **slings psychrometer** consists of two thermometers mounted side by side in a frame fitted with a handle for whirling the device through the air. The thermometers are spun until their readings become steady. In the **ventilated or aspirated psychrometer**, the thermometers remain stationary, and a small fan, blower, or syringe moves the air across the thermometer bulbs. Various designs are used in the laboratory, and commercial models are available.

Other temperature sensors, such as thermocouples and thermistors, are also used and can be adapted for recording temperatures or for use where a small instrument is required. Small-diameter wet-bulb sensors operate with low ventilation rates.

Charts and tables showing the relationship between the temperatures and humidity are available. Data are usually based on a

barometric pressure equal to one standard atmosphere. To meet special needs, charts can be produced that apply to nonstandard pressure (e.g., the ASHRAE 7500 ft psychrometric chart). Alternatively, mathematical calculations can be made (Kusuda 1965). Uncertainties of 3 to 7% rh are typical for psychrometer-based derivation. The degree of uncertainty is a function of the accuracy of the temperature measurements, wet and dry bulb, knowledge of the barometric pressure, and conformance to accepted operational procedures such as those outlined in ASHRAE *Standard* 41.6.

In air temperatures below 32°F, the water on the wick may either freeze or supercool. Because the wet-bulb temperature is different for ice and water, the state must be known and the proper chart or table used. Some operators remove the wick from the wet-bulb for freezing conditions and dip the bulb in water a few times; this allows water to freeze on the bulb between dips, forming a film of ice. Because the wet-bulb depression is slight at low temperatures,

precise temperature readings are essential. A psychrometer can be used at high temperatures, but if the wet-bulb depression is large, the wick must remain wet and water supplied to the wick must be cooled so as not to influence the wet-bulb temperature by carrying sensible heat to it (Richardson 1965, Worrall 1965).

Greenspan and Wexler (1968) and Wentzel (1961) developed devices to measure adiabatic saturation temperature.

DEW-POINT HYGROMETERS

Condensation Dew-Point Hygrometers

The condensation (chilled mirror) dew-point hygrometer is an accurate and reliable instrument with a wide humidity range. However, these features are obtained through an increase in complexity and cost compared to the psychrometer. In the condensation hygrometer, a surface is cooled (thermoelectrically, mechanically, or chemically) until dew or frost begins to condense out. The condensate surface is maintained electronically in vapor pressure equilibrium with the surrounding gas, while surface condensation is detected by optical, electrical, or nuclear techniques. The measured surface temperature is then the dew-point temperature.

The largest source of error in a condensation hygrometer stems from the difficulty in measuring condensate surface temperature accurately. Typical industrial versions of the instrument are accurate to $\pm 1.0^\circ\text{F}$ over wide temperature spans. With proper attention to the condensate surface temperature measuring system, errors can be reduced to about $\pm 0.4^\circ\text{F}$. Condensation hygrometers can be made surprisingly compact using solid-state optics and thermoelectric cooling.

Wide span and minimal errors are two of the main features of this instrument. A properly designed condensation hygrometer can measure dew points from 200°F down to frost points of -100°F . Typical condensation hygrometers can cool to 150°F below the ambient temperature, establishing lower limits of the instrument to dew points corresponding to approximately 0.5% rh. Accuracies for measurements above -40°F can be $\pm 2^\circ\text{F}$ or better, deteriorating to $\pm 4^\circ\text{F}$ at lower temperatures.

The response time of a condensation dew-point hygrometer is usually specified in terms of its cooling/heating rate, typically 4°F/s for thermoelectric cooled mirrors. This makes it somewhat faster than a heated salt hygrometer. Perhaps the most significant feature of the condensation hygrometer is its fundamental measuring technique, which essentially renders the instrument self-calibrating. For calibration, it is necessary only to manually override the surface cooling control loop, causing the surface to heat, and witness that the instrument recools to the same dew point when the loop is closed. Assuming that the surface temperature measuring system is correct, this is a reasonable check on the instrument's performance.

Although condensation hygrometers can become contaminated, they can easily be cleaned and returned to service with no impairment to performance.

Salt-Phase Heated Hygrometers

Another instrument in which the temperature varies with ambient dew-point temperature is variously designated as a self-heating salt-phase transition hygrometer or a heated electrical hygrometer. This device usually consists of a tubular substrate covered by glass fiber fabric, with a spiral bifilar winding for electrodes. The surface is covered with a salt solution, usually lithium chloride. The sensor is connected in series with a ballast and a 24 V (ac) supply. When the instrument is in operation, electrical current flowing through the salt film heats the sensor. The electrical resistance characteristics of the salt are such that a balance is reached with the salt at a critical moisture content corresponding to a saturated solution. The sensor temperature adjusts automatically so that the water vapor pressures of the salt film and ambient atmosphere are equal.

With lithium chloride, this sensor cannot be used to measure relative humidity below approximately 12% (the equilibrium relative humidity of this salt), and it has an upper dew-point limit of about 160°F . The regions of highest precision are between -10 and 93°F , and above 105°F dew point. Another problem is that the lithium chloride solution can be washed off when exposed to water. In addition, this type of sensor is subject to contamination problems, which limits its accuracy. Its response time is also very slow; it takes approximately 2 min for a 67% step change.

MECHANICAL HYGROMETERS

Many organic materials change in dimension with changes in humidity; this action is used in a number of simple and effective humidity indicators, recorders, and controllers (see [Chapter 15](#)). They are coupled to pneumatic leak ports, mechanical linkages, or electrical transduction elements to form hygrometers.

Commonly used organic materials are human hair, nylon, Dacron, animal membrane, animal horn, wood, and paper. Their inherent nonlinearity and hysteresis must be compensated for within the hygrometer. These devices are generally unreliable below 32°F . The response is generally inadequate for monitoring a changing process. Responses can be affected significantly by exposure to extremes of humidity. Mechanical hygrometers require initial calibration and frequent recalibration; however, they are useful because they can be arranged to read relative humidity directly, and they are simpler and less expensive than most other types.

ELECTRICAL IMPEDANCE AND CAPACITANCE HYGROMETERS

Many substances adsorb or lose moisture with changing relative humidity and exhibit corresponding changes in electrical impedance or capacitance.

Dunmore Hygrometers

This sensor consists of dual electrodes on a tubular or flat substrate; it is coated with a film containing salt, such as lithium chloride, in a binder to form an electrical connection between windings. The relation of sensor resistance to humidity is usually represented by graphs. Because the sensor is highly sensitive, the graphs are a series of curves, each for a given temperature, with intermediate values found by interpolation. Several resistance elements, called Dunmore elements, cover a standard range. Systematic calibration is essential because the resistance grid varies with time and contamination as well as with exposure to temperature and humidity extremes.

Polymer Film Electronic Hygrometers

These devices consist of a hygroscopic organic polymer deposited by means of thin or thick film processing technology on a water-permeable substrate. Both capacitance and impedance sensors are available. The impedance devices may be either ionic or electronic conduction types. These hygrometers typically have integrated circuits that provide temperature correction and signal conditioning. The primary advantages of this sensor technology are small size; low cost; fast response times (on the order of 1 to 120 s for 64% change in relative humidity); and good accuracy over the full range, including the low end (1 to 15% h), where most other devices are less accurate.

Ion Exchange Resin Electric Hygrometers

A conventional ion exchange resin consists of a polymer having a high relative molecular mass and polar groups of positive or negative charge in cross-link structure. Associated with these polar groups are ions of opposite charge that are held by electrostatic forces to the fixed polar groups. In the presence of water or water

vapor, the electrostatically held ions become mobile; thus, when a voltage is impressed across the resin, the ions are capable of electrolytic conduction. The **Pope cell** is one example of an ion exchange element. It is a wide-range sensor, typically covering 15 to 95% rh; therefore, one sensor can be used where several Dunmore elements would be required. The Pope cell, however, has a nonlinear characteristic from approximately 1000 Ω at 100% rh to several megohms at 10% rh.

Impedance-Based Porous Ceramic Electronic Hygrometers

Using the adsorption characteristics of oxides, humidity-sensitive ceramic oxide devices employ either ionic or electronic measurement techniques to relate adsorbed water to relative humidity. Ionic conduction is produced by dissociation of water molecules forming surface hydroxyls. The dissociation causes migration of protons such that the impedance of the device decreases with increasing water content. The ceramic oxide is sandwiched between porous metal electrodes that connect the device to an impedance-measuring circuit for linearizing and signal conditioning. These sensors have excellent sensitivity, are resistant to contamination and high temperature (up to 400°F), and may get fully wet without sensor degradation. These sensors are accurate to about $\pm 1.5\%$ rh, and $\pm 1\%$ rh when temperature compensated. These sensors have a moderate cost.

Aluminum Oxide Capacitive Sensor

This sensor consists of an aluminum strip that is anodized by a process that forms a porous oxide layer. A very thin coating of cracked chromium or gold is then evaporated over this structure. The aluminum base and the cracked chromium or gold layer form the two electrodes of what is essentially an aluminum oxide capacitor.

Water vapor is rapidly transported through the cracked chromium or gold layer and equilibrates on the walls of the oxide pores in a manner functionally related to the vapor pressure of water in the atmosphere surrounding the sensor. The number of water molecules adsorbed on the oxide structure determines the capacitance between the two electrodes.

ELECTROLYTIC HYGROMETERS

In electrolytic hygrometers, air is passed through a tube, where moisture is adsorbed by a highly effective desiccant (usually phosphorous pentoxide) and electrolyzed. The airflow is regulated to 0.0035 cfm at a standard temperature and pressure. As the incoming water vapor is absorbed by the desiccant and electrolyzed into hydrogen and oxygen, the current of electrolysis determines the mass of water vapor entering the sensor. The flow rate of the entering gas is controlled precisely to maintain a standard sample mass flow rate into the sensor. The instrument is usually designed for use with moisture-air ratios in the range of less than 1 ppm to 1000 ppm but can be used with higher humidities.

PIEZOELECTRIC SORPTION

This hygrometer compares the changes in frequency of two hygroscopically coated quartz crystal oscillators. As the mass of the crystal changes due to the absorption of water vapor, the frequency changes. The amount of water sorbed on the sensor is a function of relative humidity (i.e., partial pressure of water as well as ambient temperature).

A commercial version uses a hygroscopic polymer coating on the crystal. The humidity is measured by monitoring the change in the vibration frequency of the quartz crystal when the crystal is alternately exposed to wet and dry gas.

SPECTROSCOPIC (RADIATION ABSORPTION) HYGROMETERS

Radiation absorption devices operate on the principle that selective absorption of radiation is a function of frequency for different media. Water vapor absorbs **infrared** radiation at 2 to 3 μm wavelengths and **ultraviolet** radiation centered about the Lyman-alpha line at 0.122 μm . The amount of absorbed radiation is directly related to the absolute humidity or water vapor content in the gas mixture according to Beer's law. The basic unit consists of an energy source and optical system for isolating wavelengths in the spectral region of interest and a measurement system for determining the attenuation of radiant energy caused by the water vapor in the optical path. The absorbed radiation is measured extremely quickly and independent of the degree of saturation of the gas mixture. Response times of 0.1 to 1 s for 90% change in moisture content are common. Spectroscopic hygrometers are primarily used where a noncontact application is required; this may include atmospheric studies, industrial drying ovens, and harsh environments. The primary disadvantages of this device are its high cost and relatively large size.

GRAVIMETRIC HYGROMETERS

Humidity levels can be measured by extracting and finding the mass of water vapor in a known quantity or atmosphere. For precise laboratory work, powerful desiccants, such as phosphorous pentoxide and magnesium perchlorate, are used for the extraction process; for other purposes, calcium chloride or silica gel is satisfactory.

When the highest level of accuracy is required, the gravimetric hygrometer, developed and maintained by NIST, is the ultimate in the measurement hierarchy. The gravimetric hygrometer gives the absolute water vapor content, where the mass of the absorbed water and the precise measurement of the gas volume associated with the water vapor determine the mixing ratio or absolute humidity of the sample. This system has been chosen as the primary standard because the required measurements of mass, temperature, pressure, and volume can be made with extreme precision. However, its complexity and required attention to detail limit the usefulness of the gravimetric hygrometer.

CALIBRATION

For many hygrometers, the need for recalibration depends on the accuracy required, the stability of the sensor, and the conditions to which the sensor is being subjected. Many hygrometers should be calibrated regularly by exposure to an atmosphere maintained at a known humidity and temperature, or by comparison with a transfer standard hygrometer. Complete calibration usually requires observation of a series of temperatures and humidities. Methods for producing known humidities include saturated salt solutions (Greenspan 1977, Huang and Whetstone 1985); sulfuric acid solutions, and mechanical systems, such as the divided flow, two-pressure (Amdur 1965); two-temperature (Till and Handegord 1960); and NIST two-pressure humidity generator (Hasegawa 1976). All these systems rely on precise methods of temperature and pressure control within a controlled environment to produce a known humidity, usually with accuracies of 0.5 to 1.0%. The operating range for the precision generator is typically 5 to 95% rh.

PRESSURE MEASUREMENT

Pressure is the force exerted per unit area by a medium, generally a liquid or gas. Pressure so defined is sometimes called **absolute pressure**. Thermodynamic and material properties are expressed in terms of absolute pressures; thus, the properties of a refrigerant will be given in terms of absolute pressures. **Vacuum** refers to pressures below atmospheric.

Differential pressure is the difference between two absolute pressures. In many cases, the differential pressure can be very small compared to either of the absolute pressures (these are often referred to as low-range, high-line differential pressures). A common example of differential pressure is the pressure drop, or difference between inlet and outlet pressures, across a filter or flow element.

Gage pressure is a special case of differential pressure where one of the pressures (the reference pressure) is atmospheric pressure. Many pressure gages, including most refrigeration test sets, are designed to make gage pressure measurements, and there are probably more gage pressure measurements made than any other. Gage pressure measurements are often used as surrogates for absolute pressures. However, because of variations in atmospheric pressure due to elevation (atmospheric pressure in Denver, Colorado, is about 81% of sea-level pressure) and weather changes, the measurement of gage pressures to determine absolute pressures can significantly restrict the accuracy of the measured pressure, unless corrections are made for the local atmospheric pressure at the time of the measurement.

Pressures can be further classified as static or dynamic. **Static pressures** have a small or undetectable change with time; **dynamic pressures** include a significant pulsed, oscillatory, or other time-dependent component. Static pressure measurements are the most common, but equipment such as blowers and compressors can generate significant oscillatory pressures at discrete frequencies. Flow in pipes and ducts can generate resonant pressure changes, as well as turbulent “noise” that can span a wide range of frequencies.

Units

A plethora of pressure units, many of them poorly defined, are in common use. The international (SI) unit is the newton per square metre, called the pascal (Pa). While the bar and the standard atmosphere are used, they should not be introduced where they are not used at present. Although not internationally recognized, the pound per square inch (psi) is a widely used unit. Also widely used, but not as rigorously defined and, therefore, a potential source of error, are units based on the length of liquid columns, including inches of mercury (in. Hg), mm of mercury (mm Hg), and inches of water (in. of water). The latter is often used for low-range differential pressure measurements. In the case of pounds per square inch, the type of pressure measurement is often indicated by a modification of the unit (i.e., both psi and psia are used to indicate absolute pressure measurements, psid indicates a differential measurement, and psig indicates a gage measurement). No such standard convention exists for other units, and unless explicitly stated, reported values are assumed to be absolute pressures. Conversion factors for different pressure units can be found in [Chapter 37](#).

The difference between the conversion factors for inches of mercury and inches of water at the different temperatures is indicative of the errors that can arise from uncertainties about the definitions of these units.

Types of Pressure-Measuring Instruments

Broadly speaking, pressure instruments can be divided into three different categories—standards, mechanical gages, and electromechanical transducers. Standards instruments are used for the most accurate calibrations. The liquid-column manometer, which is the most common and potentially the most accurate standard, is used for a variety of applications, including field applications. Mechanical pressure gages are generally the least expensive and the most common pressure instruments. However, electromechanical transducers have become much less expensive and are easier to use, so they are being used more often.

PRESSURE STANDARDS

Liquid-column manometers measure pressure by determining the vertical displacement of a liquid of known density in a known gravitational field. Typically they are constructed as a U-tube of transparent material (glass or plastic). The pressure to be measured is applied to one side of the U-tube. If the other (reference) side is evacuated (zero pressure), the manometer measures absolute pressure; if the reference side is open to the atmosphere, it measures gage pressure; if the reference side is connected to some other pressure, the manometer measures the differential between the two pressures. Manometers filled with water and different oils are often used to measure low-range differential pressures. In some low-range instruments, one tube of the manometer is inclined in order to enhance the readability. Mercury-filled manometers are used for higher range differential and absolute pressure measurements. In the latter case, the reference side is evacuated, generally with a mechanical vacuum pump. Typical full-scale ranges for manometers vary from 10 in. of water to 3 atm.

For pressures above the range of manometers, standards are generally of the piston-gage, pressure-balance, or deadweight-tester type. These instruments apply pressure to the bottom of a vertical piston, which is surrounded by a close-fitting cylinder (typical clearances are millionths of an inch). The pressure generates a force approximately equal to the pressure times the area of the piston. This force is balanced by weights stacked on the top of the piston. If the mass of the weights, the local acceleration of gravity, and the area of the piston (or more properly, the “effective area” of the piston and cylinder assembly) are known, the applied pressure can be calculated. Piston gages generally generate gage pressures with respect to the atmospheric pressure above the piston. They can be used to measure absolute pressures either indirectly by separately measuring the atmospheric pressure and adding it to the gage pressure determined by the piston gage, or directly by surrounding the top of the piston and weights with an evacuated bell jar. Piston gage full-scale ranges vary from 5 to 200,000 psi.

At the other extreme, very low absolute pressures (below about 0.4 in. of water), a number of different types of standards are used. These tend to be specialized and expensive instruments found only in major standards laboratories. However, one low-pressure standard, the **McLeod gage**, has been used for field applications. Unfortunately, although the theory of the McLeod gage is simple and straightforward, it is difficult to make accurate measurements with this instrument, and major errors can occur when it is used to measure gases that condense or are adsorbed (e.g., water). In general, gages other than the McLeod gage should be used for most low-pressure or vacuum applications.

MECHANICAL PRESSURE GAGES

Mechanical pressure gages couple a pressure sensor to a mechanical readout, typically a pointer and dial. The most common type employs a **Bourdon tube** sensor, which is essentially a coiled metal tube of circular or elliptical cross section. Increasing pressure applied to the inside of the tube causes it to uncoil. A mechanical linkage translates the motion of the end of the tube to the rotation of a pointer. In most cases, the Bourdon tube is surrounded by atmospheric pressure, so that the gages measure gage pressure. A few instruments surround the Bourdon tube with a sealed enclosure that can be evacuated for absolute measurements or connected to another pressure for differential measurements. Available instruments vary widely in cost, size, pressure range, and accuracy. Full-scale ranges can vary from 5 to 100,000 psi. Accuracy of properly calibrated and used instruments can vary from 0.1 to 10% of full scale. Generally there is a strong correlation between size, accuracy, and price; larger instruments are more accurate and expensive.

To achieve better sensitivity, some low-range mechanical gages, sometimes called **aneroid gages**, employ corrugated diaphragms or capsules as sensors. The capsule is basically a short bellows sealed with end caps. These sensors are more compliant than a Bourdon tube, and a given applied pressure will cause a larger deflection of the sensor. The inside of a capsule can be evacuated and sealed in order to measure absolute pressures or connected to an external fitting to allow differential pressures to be measured. Typically, these gages are used for low-range measurements of 1 atm or less. In instruments of better quality, accuracies of 0.1% of reading or better can be achieved.

ELECTROMECHANICAL TRANSDUCERS

Mechanical pressure gages are generally limited by inelastic behavior of the sensing element, friction in the readout mechanism, and limited resolution of the pointer and dial. These effects can be eliminated or reduced by using electronic techniques to sense the distortion or stress of a mechanical sensing element and electronically convert that stress or distortion to a pressure reading. A wide variety of sensors is used, including Bourdon tubes, capsules, diaphragms, and different resonant structures whose vibration frequency varies with the applied pressure. Capacitive, inductive, and optical lever sensors are used to measure the displacement of the sensor element. In some cases, feedback techniques may be used to constrain the sensor in a null position, minimizing distortion and hysteresis of the sensing element. Temperature control or compensation is often included. Readout may be in the form of a digital display, analog voltage or current, or a digital code. Size varies, but in the case of transducers employing a diaphragm fabricated as part of a silicon chip, the sensor and signal-conditioning electronics can be contained in a small transistor package, and the largest part of the device is the pressure fitting. The best of these instruments achieve long-term instabilities of 0.01% or less of full scale, and corresponding accuracies when properly calibrated. Performance of the less expensive instruments can be more on the order of several percent.

While the dynamic response of most mechanical gages is limited by the sensor and readout, the response of some electromechanical transducers can be much faster, allowing measurements of dynamic pressures at frequencies up to 1 kHz and beyond in the case of transducers specifically designed for dynamic measurements. Manufacturers' literature should be consulted as a guide to the dynamic response of specific instruments.

As the measured pressure is reduced below about 1.5 psia, it becomes increasingly difficult to sense mechanically. A variety of gages have been developed that measure some other property of the gas that is related to the pressure. In particular, thermal conductivity gages, known as thermocouple, thermistor, Pirani, and convection gages, are used for pressures down to about 0.0004 in. of water. These gages have a sensor tube with a small heated element and a temperature sensor; the temperature of the heated element is determined by the thermal conductivity of the gas, and the output of the temperature sensor is displayed on an analog or digital electrical meter contained in an attached electronics unit. The accuracy of thermal conductivity gages is limited by their nonlinearity, dependence on gas species, and tendency to read high when contaminated. Oil contamination is a particular problem. However, these gages are small, reasonably rugged, and relatively inexpensive; in the hands of a typical user, they will give far more reliable results than a McLeod gage. They can be used to check the base pressure in a system that is being evacuated prior to being filled with refrigerant. They should be checked periodically for contamination by comparing the reading with that from a new, clean sensor tube.

GENERAL CONSIDERATIONS

Accurate values of atmospheric or barometric pressure are required for weather prediction and aircraft altimetry. In the United

States, a network of calibrated instruments, generally accurate to within 0.1% of reading and located at airports, is maintained by the National Weather Service, the Federal Aviation Administration, and local airport operating authorities. These agencies are generally cooperative in providing current values of atmospheric pressure that can be used to check the calibration of absolute pressure gages or to correct gage pressure readings to absolute pressures. However, the pressure readings generally reported for weather and altimetry purposes are not the true atmospheric pressure, but rather a value adjusted to an equivalent sea level pressure. Therefore, unless the location is near sea level, it is important to ask for the station or true atmospheric pressure rather than using the adjusted values broadcast by radio stations. Further, the atmospheric pressure decreases with increasing elevation at a rate (near sea level) of about 0.001 in. Hg/ft, and corresponding corrections should be made to account for the difference in elevation between the instruments being compared.

As noted before, gage-pressure instruments are sometimes used to measure absolute pressures, and the accuracy of these measurements can be compromised by uncertainties in the atmospheric pressure. This error can be particularly serious when gage-pressure instruments are used to measure a vacuum (negative gage pressures). For all but the most crude measurements, absolute-pressure gages should be used for vacuum measurements; for pressures below about 0.4 in. of water, a thermal conductivity gage should be used.

All pressure gages are susceptible to temperature errors. Several techniques are used to minimize these errors—sensor materials are generally chosen to minimize temperature effects, mechanical readouts can include temperature compensation elements, electromechanical transducers may include a temperature sensor and compensation circuit, and some transducers are operated at a controlled temperature. Clearly, temperature effects are of greater concern for field applications, and it is prudent to check the manufacturers' literature for the temperature range over which the specified accuracy can be maintained. Abrupt temperature changes can also cause large transient errors that may take some time to decay.

The readings of some electromechanical transducers with a resonant or vibrating sensor can depend on the gas species. Although some of these units can achieve calibrated accuracies of the order of 0.01% of reading, they are typically calibrated with dry air or nitrogen, and the readings for other gases can be in error by several percent, quite possibly much more for refrigerants and other high-density gases. High-accuracy readings can be maintained by calibrating these devices with the gas to be measured. Manufacturer's literature should be consulted.

The measurement of dynamic pressures is limited not just by the frequency response of the pressure gage, but also by the hydraulic or pneumatic time constant of the connection between the gage and the system to be monitored. As a general rule, the longer the connecting lines and the smaller their diameter, the lower the frequency response of the system. Further, even if only the static component of the pressure is of interest, and a gage with a low-frequency response is used, a significant pulsating or oscillating pressure component can cause significant errors in pressure gage readings and, in some cases, can damage the gage, particularly gages with a mechanical readout mechanism. In these cases, a filter or snubber should be used to reduce the higher frequency components.

VELOCITY MEASUREMENT

HVAC engineers measure the flow of air more often than any other gas, and the air is usually measured at or near atmospheric pressure. Under this condition, the air can be treated as an incompressible fluid, and simple formulas give sufficient precision to solve many problems. Instruments that measure fluid velocity and their application range and precision are listed in [Table 4](#).

Table 4 Velocity Measurement

Measurement Means	Application	Range, fpm	Precision	Limitations
Smoke puff or airborne solid tracer	Low air velocities in rooms; highly directional	5 to 50	10 to 20%	Awkward to use but valuable in tracing air movement
Deflecting vane anemometer	Air velocities in rooms, at outlets, etc.; directional	30 to 24,000	5%	Needs periodic check calibration
Revolving vane anemometer	Moderate air velocities in ducts and rooms; somewhat directional	100 to 3000	2 to 5%	Extremely subject to error with variations in velocities with space or time; easily damaged; needs periodic calibration
Hot-wire anemometer	a. Low air velocities; directional and nondirectional available	1 to 2000	2 to 5%	Requires accurate calibration at frequent intervals. Some are relatively costly.
	b. High air velocities	Up to 60,000	0.2 to 5%	
	c. Transient velocity and turbulence			
Pitot tube	Standard instrument for measuring duct velocities	180 to 10,000 with micromanometer; 600 to 10,000 with draft gages; 10,000 up with manometer	1 to 5%	Accuracy falls off at low end of range
Impact tube and sidewall or other static tap	High velocities, small tubes and where air direction may be variable	120 to 10,000 with micromanometer; 600 to 10,000 with draft gages; 10,000 up with manometer	1 to 5%	Accuracy depends on constancy of static pressure across stream section
Cup anemometer	Meteorological	Up to 12,000	2 to 5%	Poor accuracy at low air velocity (<500 fpm)
Laser Doppler velocimeter	Calibration of air velocity instruments	1 to 6000	1 to 3%	High cost and complexity limit LDVs to laboratory applications

AIRBORNE TRACER TECHNIQUES

Tracer techniques are suitable for measuring velocity in an open space. Typical tracers include smoke, feathers, pieces of lint, and radioactive or nonradioactive gases. Measurements are made by timing the rate of movement of solid tracers or by monitoring the change in concentration level of gas tracers.

Smoke is a useful qualitative tool in studying air movements. Smoke can be obtained from titanium tetrachloride (irritating to nasal membranes) or by mixing potassium chlorate and powdered sugar (a nonirritating smoke) and firing the mixture with a match. The latter process produces considerable heat and should be confined to a pan away from flammable materials. Titanium tetrachloride smoke works well for spot tests, particularly for leakage through casings and ducts, because it can be handled easily in a small, pistol-like ejector.

The fumes of ammonia water and sulfuric acid, if permitted to mix, form a white precipitate. Two bottles, one containing ammonia water and the other containing acid, are connected to a common nozzle by rubber tubing. A syringe forces air over the liquid surfaces in the bottles; the two streams mix at the nozzle and form a white cloud.

A satisfactory test smoke also can be made by bubbling an air-stream through ammonium hydroxide and then hydrochloric acid (Nottage et al. 1952). Smoke tubes, smoke candles, and smoke bombs are available for studying airflow patterns.

ANEMOMETERS

Deflecting Vane Anemometers

The deflecting vane anemometer consists of a pivoted vane enclosed in a case. Air exerts pressure on the vane as it passes through the instrument from an upstream to a downstream opening. A hair spring and a damping magnet resist vane movement. The instrument gives instantaneous readings of directional velocities on an indicating scale. With fluctuating velocities, it is necessary to

average the needle swings visually to obtain average velocities. This instrument is useful for studying air motion in a room; locating objectionable drafts; measuring air velocities at supply and return diffusers and grilles; and measuring laboratory hood face velocities.

Propeller or Revolving Vane Anemometers

The propeller anemometer consists of a light, revolving wind-driven wheel connected through a gear train to a set of recording dials that read linear feet of air passing in a measured length of time. It is made in various sizes—3, 4, and 6 in. are the most common. Each instrument requires individual calibration. At low velocities, the friction drag of the mechanism is considerable. To compensate for this, a gear train that overspeeds is commonly used. For this reason, the correction is often additive at the lower range and subtractive at the upper range, with the least correction in the middle range of velocities. The best of these instruments have starting speeds of 50 fpm or higher; therefore, they cannot be used below that air speed. Electronic revolving vane anemometers, with optical or magnetic pickups to sense the rotation of the vane, are available. Sizes for the vanes range as small as 1/2 in. in diameter for the electronic versions.

Cup Anemometers

The cup anemometer is primarily used to measure outdoor, meteorological wind speeds. It consists of three or four hemispherical cups mounted radially from a vertical shaft. Wind from any direction with a vector component in the plane of cup rotation causes the cups and shaft to rotate. Because the primary use of this anemometer is to make meteorological wind speed measurements, the instrument is usually constructed so that wind speeds can be recorded or indicated electrically at a remote point.

Thermal Anemometers

The thermal or hot-wire anemometer consists of a heated RTD, thermocouple junction, or thermistor sensor constructed at the end

of a probe; it is designed to provide a direct, simple method of determining air velocity at a point in the flow field. The probe is placed into an airstream, and the movement of air past the electrically heated velocity sensor tends to cool the sensor in proportion to the speed of the airflow. The electronics and sensor are commonly combined into a portable, hand-held device that interprets the sensor signal and provides a direct reading of air velocity in either analog or digital display format. Often the sensor probe also incorporates an ambient temperature-sensing RTD or thermistor, in which case the indicated air velocity is “temperature compensated” to “standard” air density conditions (typically 0.0748 lb/ft³).

Hot-wire anemometers have long been used in the fluid flow research field. Research anemometer sensors have been constructed using very fine wires in configurations that allow the researcher to characterize fluid flows in one, two, and three dimensions with sensor/electronics response rates up to several hundred kilohertz. This technology has been incorporated into more ruggedized sensors suitable for measurements in the HVAC field, primarily for unidirectional airflow measurement. Omnidirectional sensing instruments suitable for thermal comfort studies are also available.

The principal advantages of thermal anemometers are their wide dynamic range and their ability to sense extremely low velocities. Typical accuracy (including repeatability) of 2 to 5% of reading over the entire velocity range is often achieved in commercially available portable instruments.

Among the limitations of thermal anemometers are the following: (1) the unidirectional sensor must be carefully aligned in the airstream (typically to within ±20° rotation) to achieve accurate results; (2) the velocity sensor must be kept clean because contaminant buildup will cause the calibration to change; and (3) due to the inherent high speed of response of thermal anemometers, measurements in turbulent flows can yield fluctuating velocity measurements. Electronically controlled time-integrated functions are now available in many digital air velocity meters to help smooth these turbulent flow measurements.

In the HVAC field, thermal anemometers are suitable for use in a variety of applications. They are particularly well-suited to the low velocities associated with laboratory fume hood face velocity measurements (typically in the 60 to 200 fpm range). Thermal anemometers can also be used for taking multipoint traverse measurements in ventilation ductwork.

Laser Doppler Velocimeters (or Anemometers)

The laser Doppler velocimeter (LDV) or laser Doppler anemometer (LDA) is an extremely complex system that collects scattered light produced by a particle passing through the intersection volume of two intersecting laser beams of the same light frequency (Mease et al. 1992). The scattered light consists of bursts containing a regularly spaced fringe pattern whose frequency is linearly proportional to the speed of the particle. Due to the cost and complexity of these systems, they are usually not suitable for in situ field measurements. Rather, the primary application of LDV systems in the HVAC industry is the calibration of systems used to calibrate other air velocity instruments.

The greatest advantage of an LDV is its performance at low air speeds. It is capable of reading air speeds as low as 15 fpm with uncertainty levels of 1% or less (Mease et al. 1992). In addition, it is nonintrusive in the flow—only optical access is required. It can be used to measure fluctuating components as well as mean speeds and is available in one-, two-, and even three-dimensional configurations. Its biggest disadvantages are its high cost and extreme technological complexity, which requires highly skilled operators. Modern fiber optic systems require less-skilled operators but at a considerable increase in cost.

PITOT-STATIC TUBES

The pitot-static tube, in conjunction with a suitable manometer or differential pressure transducer, provides a simple method of determining air velocity at a point in a flow field. Figure 5 shows the construction of a standard pitot tube (ASHRAE Standard 51) and the method of connecting it with inclined manometers to display both static pressure and velocity pressure. The equation for determining air velocity from measured velocity pressure is

$$V = C \sqrt{\frac{2p_w g_c}{\rho}} \quad (5)$$

where

V = velocity, fpm

p_w = velocity pressure (pitot-tube manometer reading), in. of water

ρ = density of air, lb_m/ft³

g_c = gravitational constant = 32.174 lb_m·ft/lb_f·s²

C = unit conversion factor = 136.8

The type of manometer or differential pressure transducer used with a pitot-static tube depends on the magnitude of velocity pressure being measured and on the desired accuracy. At velocities greater than 1500 fpm, a draft gage of appropriate range is usually satisfactory. If the pitot-static tube is used to measure air velocities lower than 1500 fpm, a precision manometer or comparable pressure differential transducer is essential.

Other pitot-static tubes have been used and calibrated. To meet special conditions, various sizes of pitot-static tubes geometrically similar to the standard tube can be used. For relatively high velocities in ducts of small cross-sectional area, total pressure readings can be obtained with an impact (pitot) tube. Where static pressure across the stream is relatively constant, as in turbulent flow in a straight duct, a sidewall tap to obtain static pressure can be used with the impact tube to obtain the velocity pressure head. One form of impact tube is a small streamlined tube with a fine hole in its upstream end and with its axis placed parallel to the stream.

If the Mach number of the flow is greater than about 0.3, the effects of compressibility should be included in the computation of the air speed from pitot-static and impact (stagnation or pitot) tube measurements (Mease et al. 1992).

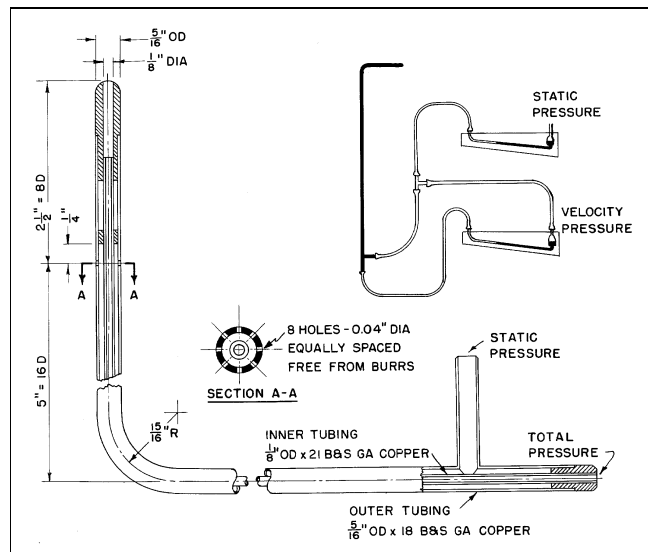


Fig. 5 Standard Pitot Tube

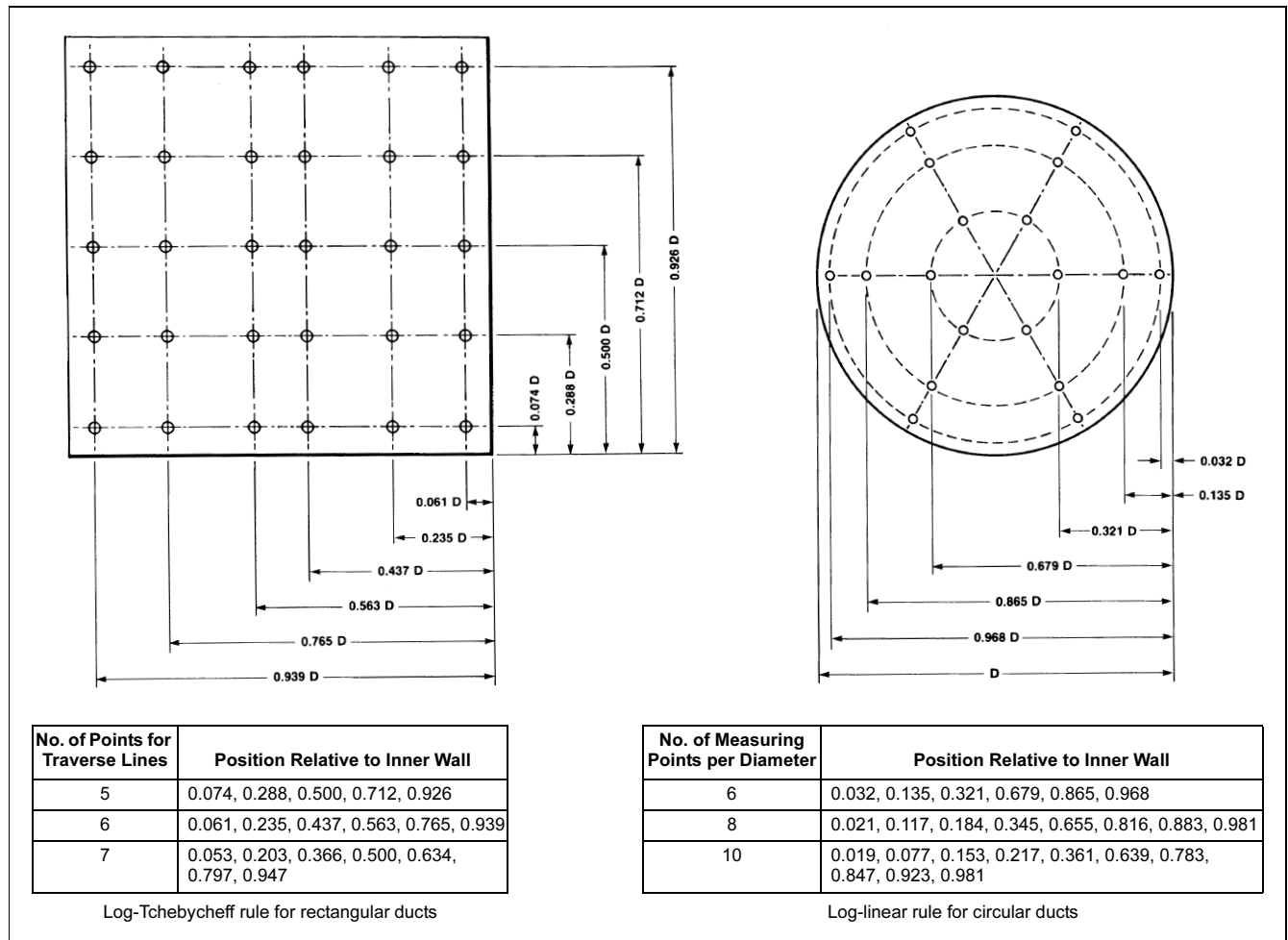


Fig. 6 Measuring Points for Rectangular and Round Duct Traverse

MEASURING FLOW IN DUCTS

Because velocity in a duct is seldom uniform across any section, and a pitot tube reading or thermal anemometer indicates velocity at only one location, a traverse is usually made to determine average velocity. Generally, velocity is lowest near the edges or corners and greatest at or near the center.

To determine the velocity in a traverse plane, a straight average of individual point velocities will give satisfactory results when point velocities are determined by the **log-Tchebycheff rule** (ISO Standard 3966) or, if care is taken, by the **equal area method**. [Figure 6](#) shows suggested sensor locations for traversing round and rectangular ducts. The log-Tchebycheff rule provides the greatest accuracy because its location of traverse points accounts for the effect of wall friction and the fall-off of velocity near wall ducts. This method is now recommended for rectangular ducts, although for circular ducts the log-Tchebycheff and log-linear traverse methods are similar. Log-Tchebycheff minimizes the positive error (measured greater than actual) caused by the failure to account for losses at the duct wall. This error can occur when using the older method of equal subareas to traverse rectangular ducts.

For a rectangular duct traverse, a minimum of 25 points should be measured. For a duct size less than 18 in., the points should be located at the center of equal areas not more than 6 in. apart, and a minimum of 2 points per side should be used. For a duct side greater than 56 in., the maximum distance between points is 8 in. For a cir-

cular duct traverse, the log-linear rule and three symmetrically disposed diameters may be used ([Figure 6](#)). Points on two perpendicular diameters may be used where access is limited.

If possible, measuring points should be located at least 7.5 diameters downstream and 3 diameters upstream from a disturbance (e.g., caused by a turn). Compromised traverses as close as 2 diameters downstream and 1 diameter upstream can be performed with an increase in measurement error. Because field-measured airflows are rarely steady and uniform, particularly near disturbances, accuracy can be improved by increasing the number of measuring points. Straightening vanes (ASHRAE Standard 51) located 1.5 duct diameters ahead of the traverse plane improve measurement precision.

When velocities at a traverse plane fluctuate, the readings should be averaged on a time-weighted basis. Two traverse readings in short succession also help to average out velocity variations that occur with time. If negative velocity pressure readings are encountered, they are considered a measurement value of zero and calculated in the average velocity pressure. ASHRAE Standard 111 has further information on measuring flow in ducts.

FLOW RATE MEASUREMENT

Various means of measuring fluid flow rate are listed in [Table 5](#). The values for volumetric or mass flow rate measurement (ASME Standard PTC 19.5, Benedict 1984) are often determined

Table 5 Volumetric or Mass Flow Rate Measurement

Measurement Means	Application	Range	Precision	Limitations
Orifice and differential pressure measurement system	Flow through pipes, ducts, and plenums for all fluids	Above Reynolds number of 5000	1 to 5%	Discharge coefficient and accuracy influenced by installation conditions
Nozzle and differential pressure measurement system	Flow through pipes, ducts, and plenums for all fluids	Above Reynolds number of 5000	0.5 to 2.0%	Discharge coefficient and accuracy influenced by installation conditions
Venturi tube and differential pressure measurement system	Flow through pipes, ducts, and plenums for all fluids	Above Reynolds number of 5000	0.5 to 2.0%	Discharge coefficient and accuracy influenced by installation conditions
Timing given mass or volumetric flow	Liquids or gases; used to calibrate other flowmeters	Any	0.1 to 0.5%	System is bulky and slow
Rotameters	Liquids or gases	Any	0.5 to 5.0%	Should be calibrated for fluid being metered
Displacement meter	Relatively small volumetric flow with high pressure loss	As high as 1000 cfm depending on type	0.1 to 2.0% depending on type	Most types require calibration with fluid being metered
Gasometer or volume displacement	Short-duration tests; used to calibrate other flowmeters	Total flow limited by available volume of containers	0.5 to 1.0%	—
Thomas meter (temperature rise of stream due to electrical heating)	Elaborate setup justified by need for good accuracy	Any	1%	Uniform velocity; usually used with gases
Element of resistance to flow and differential pressure measurement system	Used for check where system has calibrated resistance element	Lower limit set by readable pressure drop	1 to 5%	Secondary reading depends on accuracy of calibration
Turbine flowmeters	Liquids or gases	Any	0.25 to 2.0%	Uses electronic readout
Instrument for measuring velocity at point in flow	Primarily for installed systems with no special provision for flow measurement	Lower limit set by accuracy of velocity measurement	2 to 4%	Accuracy depends on uniformity of flow and completeness of traverse
Heat input and temperature changes with steam and water coil	Check value in heater or cooler tests	Any	1 to 3%	—
Laminar flow element and differential pressure measurement system	Measure liquid or gas volumetric flow rate; nearly linear relationship with pressure drop; simple and easy to use	0.0001 to 2000 cfm	1%	Fluid must be free of dirt, oil, and other impurities that could plug meter or affect its calibration
Magnetohydrodynamic flowmeter (electromagnetic)	Measures electrically conductive fluids, slurries; meter does not obstruct flow; no moving parts	0.1 to 10,000 gpm	1%	At present state of the art, conductivity of fluid must be greater than 5 $\mu\text{mho/cm}$
Swirl flowmeter and vortex shedding meter	Measure liquid or gas flow in pipe; no moving parts	Above Reynolds number of 10^4	1%	—

by measuring pressure difference across an orifice, nozzle, or venturi tube. The various meters have different advantages and disadvantages. For example, the orifice plate is more easily changed than the complete nozzle or venturi tube assembly. However, the nozzle is often preferred to the orifice because its discharge coefficient is more precise. The venturi tube is a nozzle followed by an expanding recovery section to reduce net pressure loss. Differential pressure-type flow measurement has benefited through workshops addressing fundamental issues, textbooks, research, and improved standards (Miller 1983, DeCarlo 1984, Mattingly 1984, ASME *Standard* B40.1, ASME *Standard* MFC-9M, ASME *Standard* MFC-10M, ASME *Standard* MFC-1M).

Fluid meters use a wide variety of physical techniques to make flow measurements (ASME *Standard* PTC 19.5, Miller 1983, DeCarlo 1984); those more prevalently used are described in this section. To assure and validate the accuracy of flow rate measurement instruments, appropriate calibration procedures should include documentation of traceability to the calibration facility. The calibration facility should, in turn, provide documentation of traceability to national standards.

Flow Measurement Methods

Direct. Both gas and liquid flow can be measured quite accurately by timing a collected amount of fluid that is measured gravimetrically

or volumetrically. While this method is commonly used for calibrating other metering devices, it is particularly useful where the flow rate is low or intermittent and where a high degree of accuracy is required. These systems are generally large and slow, but in their simplicity, they can be considered primary devices.

The variable area meter or rotameter is a convenient direct-reading flowmeter for liquids and gases. This is a vertical, tapered tube in which the flow rate is indicated by the position of a float suspended in the upward flow. The position of the float is determined by its buoyancy and the upwardly directed fluid drag.

Displacement meters measure total liquid or gas flow over time. The two major types of displacement meters used for gases are the conventional gas meter, which uses a set of bellows, and the wet test meter, which uses a water displacement principle.

Indirect. The **Thomas meter** is used in laboratories to measure high gas flow rates with low pressure losses. The gas is heated by electric heaters, and the temperature rise is measured by two resistance thermometer grids. When the heat input and temperature rise are known, the mass flow of gas is calculated as the quantity of gas that will remove the equivalent heat at the same temperature rise.

A velocity traverse (made using a pitot tube or other velocity-measuring instrument) measures airflow rates in the field or calibrates large nozzles. This method can be imprecise at low velocities and impracticable where many test runs are in progress.

Another field-estimating method measures the pressure drop across elements with known pressure drop characteristics, such as heating and cooling coils or fans. If the pressure drop/flow rate relationship has been calibrated, the results can be precise. If the method depends on rating data, it should be used for check purposes only.

VENTURI, NOZZLE, AND ORIFICE FLOWMETERS

Flow in a pipeline can be measured by a venturi meter (Figure 7), flow nozzle (Figure 8), or orifice plate (Figure 9). American Society of Mechanical Engineers (ASME) *Standard* MFC-3M describes measurement of fluid flow in pipes using the orifice, nozzle, and venturi; ASME *Standard* PTC 19.5 specifies their construction.

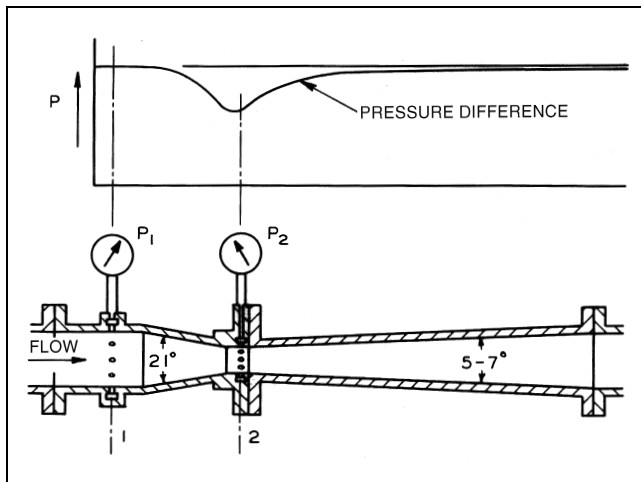


Fig. 7 Typical Herschel Type Venturi Meter

Assuming an incompressible fluid (liquid or slow-moving gas), uniform velocity profile, frictionless flow, and no gravitational effects, the principle of conservation of mass and energy can be applied to the venturi and nozzle geometries to give

$$w = \rho V_1 A_1 = \rho V_2 A_2 = A_2 \sqrt{\frac{2g_c \rho (p_1 - p_2)}{1 - \beta^4}} \quad (6)$$

where

- w = mass flow rate, lb_m/s
- V = velocity of stream, fps
- A = flow area, ft²
- g_c = gravitational constant = 32.174 lb_m · ft/lb_f · s²
- ρ = density of fluid, lb_m/ft³
- p = absolute pressure, lb_f/ft²
- β = (D_2/D_1) for venturi and sharp edge orifice and d/D for flow nozzle

Note: Subscript 1 refers to the entering conditions; subscript 2 refers to the throat conditions.

Because the flow through the meter is not frictionless, a correction factor C is defined to account for friction losses. If the fluid is at a high temperature, an additional correction factor F_a should be included to account for thermal expansion of the primary element. Because this amounts to less than 1% at 500°F, it can usually be omitted. Equation (6) then becomes

$$w = CA_2 \sqrt{\frac{2g_c \rho (p_1 - p_2)}{1 - \beta^4}} \quad (7)$$

where C is the friction loss correction factor.

The factor C is a function of geometry and Reynolds number. Values of C are given in ASME *Standard* PTC 19.5. The jet passing through an orifice plate contracts to a minimum area at the vena contracta located a short distance downstream from the orifice

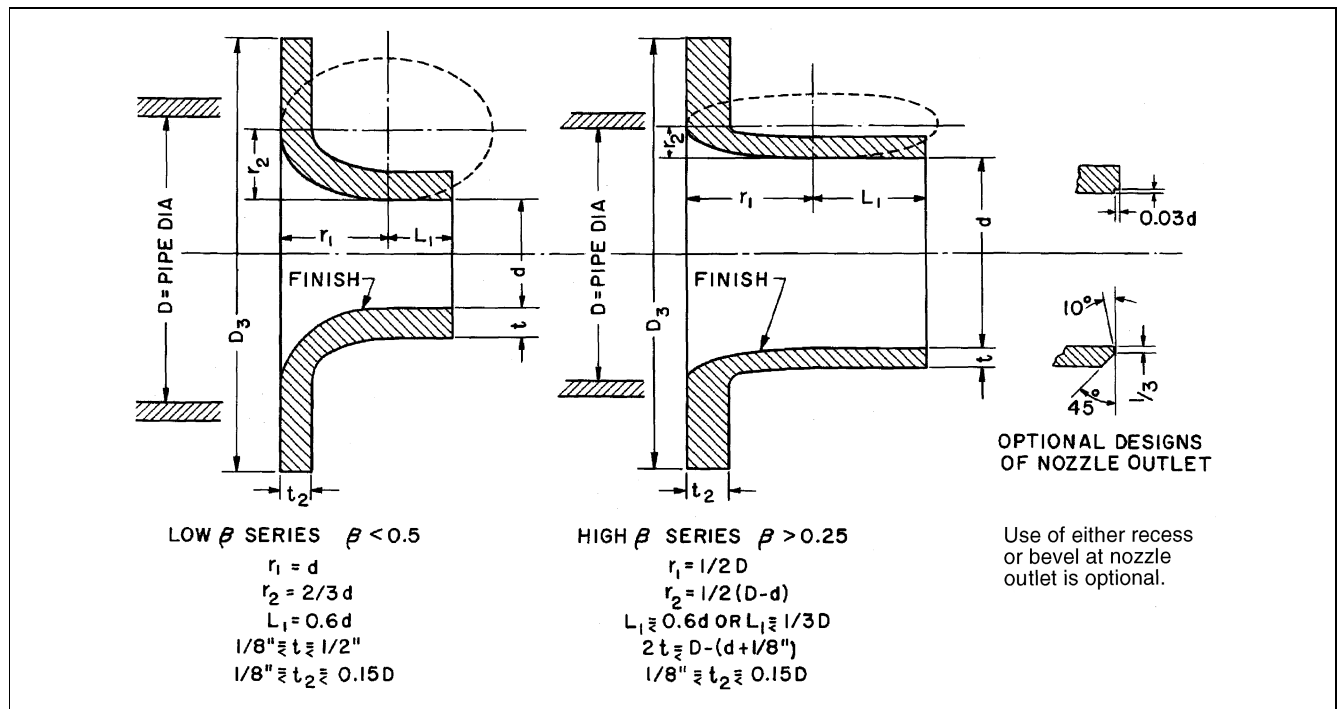


Fig. 8 Dimensions of ASME Long-Radius Flow Nozzles
From ASME PTC 19.5. Reprinted with permission of ASME.

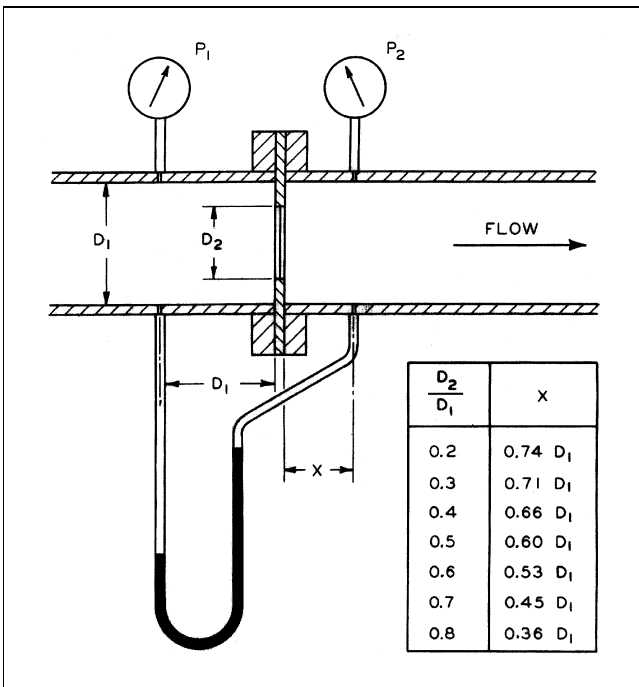


Fig. 9 Sharp Edge Orifice with Pressure Tap Locations
From ASME PTC 19.5. Reprinted with permission of ASME.

plate. The contraction coefficient, friction loss coefficient C , and approach factor $1/(1 - \beta^4)^{0.5}$ can be combined into a single constant K , which is a function of geometry and Reynolds number. The orifice flow rate equations then become

$$Q = KA_2 \sqrt{\frac{2g_c(p_1 - p_2)}{\rho}} \tag{8}$$

where

Q = discharge flow rate, cfs

A_2 = orifice area, ft²

$p_1 - p_2$ = pressure drop as obtained by pressure taps, lb_f/ft²

Values of K are shown in ASME Standard PTC 19.5.

Valves, bends, and fittings upstream from the flowmeter can cause errors. Long, straight pipes should be installed upstream and downstream from the flow devices to assure fully developed flow for proper measurement. ASHRAE Standard 41.8 specifies upstream and downstream pipe lengths for measuring flow of liquids with an orifice plate. ASME Standard PTC 19.5 gives the piping requirements between various fittings and valves and the venturi, nozzle, and orifice. If these conditions cannot be met, flow conditioners or straightening vanes can be used (ASME Standard PTC 19.5, ASME Standard MFC-10M, Mattingly 1984, Miller 1983).

Compressibility effects must be considered for gas flow if the pressure drop across the measuring device is more than a few percent of the initial pressure.

Nozzles are sometimes arranged in parallel pipes from a common manifold; thus, the capacity of the testing equipment can be changed by shutting off the flow through one or more nozzles. An apparatus designed for testing airflow and capacity of air-conditioning equipment is described by Wile (1947), who also presents pertinent information on nozzle discharge coefficients, Reynolds numbers, and resistance of perforated plates. Some laboratories refer to this apparatus as a code tester.

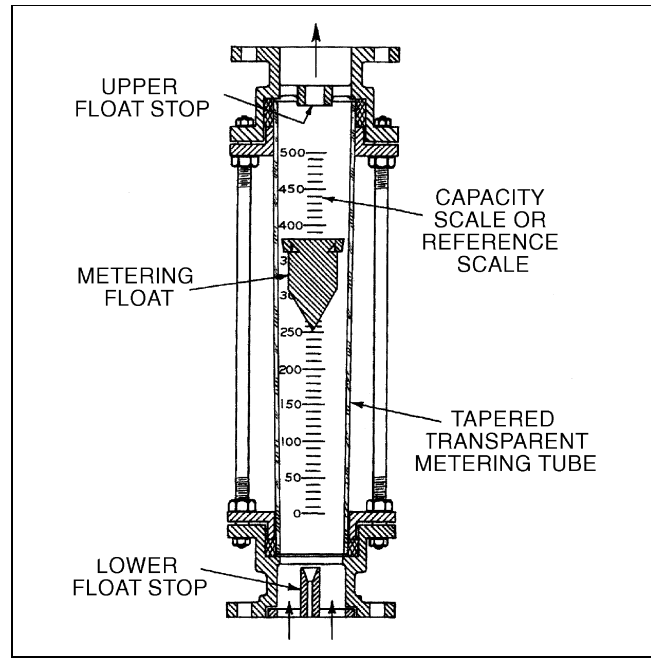


Fig. 10 Variable Area Flowmeter

VARIABLE AREA FLOWMETERS (ROTAMETERS)

In permanent installations where high precision, ruggedness, and operational ease are important, the variable area flowmeter is satisfactory. It is frequently used to measure liquids or gases in small-diameter pipes. For ducts or pipes over 6 in. in diameter, the expense of this meter may not be warranted. In larger systems, however, the meter can be placed in a bypass line and used with an orifice.

The variable area meter (Figure 10) commonly consists of a float that is free to move vertically in a transparent tapered tube. The fluid to be metered enters at the narrow bottom end of the tube and moves upward, passing at some point through the annulus formed between the float and the inside wall of the tube. At any particular flow rate, the float assumes a definite position in the tube; a calibrated scale on the tube shows the float's location and the fluid flow rate.

The position of the float is established by a balance between the fluid pressure forces across the annulus and gravity on the float. The buoyant force supporting the float, $v_f(\rho_f - \rho)g/g_c$, is balanced by the pressure difference acting on the cross-sectional area of the float $A_f \Delta p$, where ρ_f , A_f , and v_f are, respectively, the float density, float cross-sectional area, and float volume. The pressure difference across the annulus is

$$\Delta p = \frac{v_f(\rho_f - \rho)g}{A_f g_c} \tag{9}$$

The mass flow follows from Equation (8) as

$$w = KA_2 \sqrt{\frac{2v_f(\rho_f - \rho)g\rho}{A_f}} \tag{10}$$

The flow for any selected fluid is nearly proportional to the area, so that calibration of the tube is convenient. To use the meter for different fluids, the flow coefficient variation for any float must be known. Float design can reduce variation of the flow coefficient with Reynolds number; float materials can reduce the dependence of mass flow calibration on fluid density.

POSITIVE DISPLACEMENT METERS

Many positive displacement meters are available for measuring total liquid or gas volumetric flow rates. The fluid measured in these meters flows progressively into compartments of definite size. As the compartments are filled, they are rotated so that the fluid discharges from the meter. The flow rate through the meter is equal to the product of the compartment volume, the number of compartments, and the rotation rate of the rotor. Most of these meters have a mechanical register calibrated to show total flow.

TURBINE FLOWMETERS

Turbine flowmeters are volumetric flow rate sensing meters with a magnetic stainless steel turbine rotor suspended in the flow stream of a nonmagnetic meter body. The fluid stream exerts a force on the blades of the turbine rotor, setting it in motion and converting the fluid's linear velocity to an angular velocity. Design motivation for turbine meters is to have the rotational speed of the turbine proportional to the average fluid velocity and thus to the volume rate of fluid flow (Miller 1983, DeCarlo 1984, Mattingly 1992).

The rotational speed of the rotor is monitored by an externally mounted pickoff assembly. Magnetic and radio frequency are the most commonly used pickoffs. The magnetic pickoff contains a permanent magnet and coil. As the turbine rotor blades pass through the field produced by the permanent magnet, a shunting action induces ac voltage in the winding of the coil wrapped around the magnet. A sine wave with a frequency proportional to the flow rate develops. With the radio frequency pickoff, an oscillator applies a high-frequency carrier signal to a coil in the pickoff assembly. The rotor blades pass through the field generated by the coil and modulate the carrier signal by shunting action on the field shape. The carrier signal is modulated at a rate corresponding to the rotor speed, which is proportional to the flow rate. With both pickoffs, frequency of the pulses generated becomes a measure of flow rate, and the total number of pulses measures total volume (Woodring 1969, Shafer 1961, Mattingly 1992).

Because output frequency of the turbine flowmeter is proportional to flow rate, every pulse from the turbine meter is equivalent to a known volume of fluid that has passed through the meter; the sum of these pulses yields total volumetric flow. Summation is accomplished by electronic counters designed for use with turbine flowmeters; they combine a mechanical or electronic register with the basic electronic counter.

Turbine flowmeters should be installed with straight lengths of pipe upstream and downstream from the meter. The length of the inlet and outlet pipes should be according to manufacturers' recommendations or pertinent standards. Where recommendations of standards cannot be accommodated, the meter installation should be calibrated. Some turbine flowmeters can be used in bidirectional flow applications. A fluid strainer, used with liquids of poor or marginal lubricity, minimizes bearing wear.

The lubricity of the process fluid and the type and quality of rotor bearings determine whether the meter is satisfactory for the particular application. When choosing turbine flowmeters for use with fluorocarbon refrigerants, attention must be paid to the type of bearings used in the meter and to the oil content of the refrigerant. For these applications, sleeve-type rather than standard ball bearings are recommended. The amount of oil in the refrigerant can severely affect calibration and bearing life.

In metering liquid fluorocarbon refrigerants, the liquid must not flash to a vapor (cavitate). This would cause a tremendous increase in flow volume. Flashing results in erroneous measurements and rotor speeds that can damage the bearings or cause a failure. Flashing can be avoided by maintaining an adequate back pressure on the downstream side of the meter (Liptak 1972).

AIRFLOW-MEASURING HOODS

Flow-measuring hoods are portable instruments designed to measure supply or exhaust airflow through diffusers and grilles in HVAC systems. A flow-measuring hood assembly typically consists of a fabric hood section, a plastic or metal base, an airflow-measuring manifold, a meter, and handles for carrying and holding the hood in place.

For volumetric airflow measurements, the flow-measuring hood is placed over a diffuser or grille. The fabric hood captures and directs airflow from the outlet or inlet across the flow-sensing manifold in the base of the instrument. The manifold consists of a number of tubes containing upstream and downstream holes in a grid pattern designed to simultaneously sense and average multiple velocity points across the base of the flow-measuring hood. Air from the upstream holes flows through the tubes past a sensor and then exits through the downstream holes. Sensors employed by different manufacturers include swinging vane anemometers, electronic micromanometers, and thermal anemometers. In the case of the electronic micromanometer sensor, air does not actually flow through the manifold, but the airtight sensor senses the pressure differential from the upstream to downstream series of holes. The meter on the base of the flow-measuring hood interprets the signal from the sensor and provides a direct reading of volumetric flow in either an analog or digital display format.

As a performance check in the field, the indicated flow of a measuring hood can be compared to a duct traverse flow measurement (using a pitot-tube or a thermal anemometer). All flow-measuring hoods induce some back pressure on the air-handling system because the hood restricts the flow coming out of the diffuser. This added resistance alters the true amount of air coming out of the diffuser. In most cases, this error is negligible and is less than the accuracy of the instrument. For proportional balancing, this error need not be taken into account because all similar diffusers will have about the same amount of back pressure. To determine whether back pressure is significant, a velocity traverse can be made in the duct ahead of the diffuser with and without the flow-measuring hood in place. The difference in the average velocity of the traverse indicates the degree of back-pressure compensation required on similar diffusers in the system. For example, if the average velocity is 800 fpm with the hood in place and 820 fpm without the hood, the indicated flow reading can be multiplied by 1.025 on similar diffusers in the system ($820/800 = 1.025$). As an alternative, the designer of the air-handling system can predict the reduction in airflow due to the additional pressure of the hood by using a curve supplied by the flow-measuring hood manufacturer. This curve indicates the pressure drop through the hood for different flow rates.

AIR INFILTRATION, AIRTIGHTNESS, AND OUTDOOR AIR VENTILATION RATE MEASUREMENT

Two major characteristics describe air infiltration in buildings—air exchange rate and envelope air leakage. The measurement approaches used to determine these factors are described in [Chapter 26](#). The air exchange rate of a building refers to the rate at which outdoor air enters the building under normal weather and ventilation system operation. In general, the air change rate includes both outdoor air taken in through the air handlers and air leakage through the building envelope (infiltration). The outdoor air intake rate is determined by the design, installation, and operation of the mechanical ventilation system. Infiltration is determined by the extent and distribution of leaks over the building envelope and the pressure differences across these leaks. These pressure differences are induced by wind, inside-outside temperature differences, and the operation of building mechanical equipment. To fully characterize the air

exchange performance of a building, the air exchange rate must be measured over a range of weather and equipment operation.

The outdoor air ventilation rate is an indicator of the rate of dilution of occupant- and building-generated contaminants. Building air exchange rates can be measured by injecting a tracer gas into a building and monitoring and analyzing the tracer gas concentration response. The equipment required for tracer testing includes (1) a means of injecting the tracer gas and (2) a tracer gas monitor. A variety of tracer gas techniques are used. They are distinguished by their injection strategy and analysis approach. These techniques include the constant concentration (equilibrium tracer), tracer decay (ASTM *Standard E 741*), and outside air fraction (air ratio) methods. Tracer decay is the simplest and most accurate of these techniques (as per [Chapter 26](#)), but the other methods may be satisfactory if care is taken. Carbon dioxide is often used as a tracer gas because CO₂ gas monitors are relatively inexpensive and easy to use, and occupant-generated CO₂ can be used for most tracer gas techniques. Bottled CO₂ or CO₂ fire extinguishers are also readily available for tracer gas injection.

The airtightness of a building envelope can be measured relatively quickly using building pressurization techniques, which are described in [Chapter 26](#). In the pressurization technique, a large fan or blower mounted in a door or window induces a large and roughly uniform pressure difference across the building shell. The airflow required to maintain this pressure difference is then measured. The more leakage in the building, the more airflow is required to induce a specific indoor-outdoor pressure difference. The building airtightness is characterized by the airflow rate at a reference pressure, normalized by the building volume or surface area. Under proper test conditions, the results of a pressurization test are independent of weather conditions. The instrumentation requirements for pressurization testing include air-moving equipment, a device to measure airflow, and a differential pressure gage.

CARBON DIOXIDE MEASUREMENT

Carbon dioxide has become an important measurement parameter for air-conditioning, heating, and refrigerating engineers, particularly for use in indoor air quality (IAQ) applications. Although CO₂ is generally not of concern as a specific toxin in indoor air, it is used as a surrogate indicator of odor related to human occupancy. ASHRAE *Standard 62* states that maintaining CO₂ concentrations below 1000 ppm (based on a differential of 700 ppm between indoor and outdoor CO₂ concentrations) usually results in conditions conducive to comfort and reduced odor from human-generated pollutants. *Standard 62* also recommends specific minimum outdoor air ventilation rates to ensure adequate indoor air quality. Carbon dioxide is often used as a tracer gas when quantifying outdoor air ventilation rates. Carbon dioxide sensors are also used in building control strategies to optimize ventilation as a function of occupancy.

NONDISPERSIVE INFRARED CO₂ DETECTORS

The technology in most widespread use for IAQ applications is the nondispersive infrared (NDIR) sensor ([Figure 11](#)). This device makes use of the strong absorption band that CO₂ produces at 4.2 μm when excited by an infrared light source. Indoor air quality-specific NDIR instruments, when calibrated between 0 and 5000 ppm, are typically accurate within 150 ppm, but the accuracy of some sensors can be improved to within 50 ppm if the instrument is calibrated for a narrower range. Portable NDIR meters are available with direct-reading digital displays; however, response time varies significantly among different instruments. While most NDIR cell designs facilitate very rapid CO₂ sample diffusion, some of the instruments now in widespread use for indoor air quality measurement exhibit slower sensor response, resulting in stabilization times greater than 5 min (up to 15 min), which may complicate walk-through inspections.

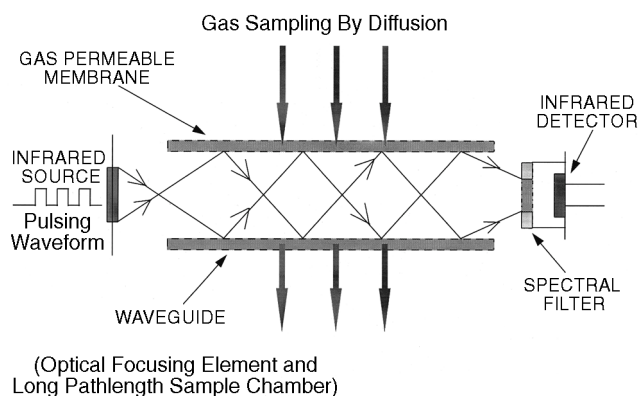


Fig. 11 Nondispersive Infrared Carbon Dioxide Sensor

Calibration

In a clean, stable environment, NDIR sensors can hold calibration for months, but condensation, dust, dirt, and mechanical shock may offset calibration. As with all other CO₂ sensor technologies, NDIR sensor readings are proportional to pressure due to the change in density of the gas molecules that results from a change in the sample pressure. This leads to errors in CO₂ readings when the barometric pressure changes from the calibration pressure. Weather-induced errors will be small, but all CO₂ instruments should be recalibrated if used at an altitude that is significantly different from the calibration altitude. Some NDIR sensors are sensitive to cooling effects when placed in an airstream. This is an important consideration when locating a fixed sensor or when using a portable system to evaluate air-handling system performance because airflow in supply and return ducts may significantly shift readings.

Applications

Nondispersive infrared sensors are well suited for equilibrium tracer and tracer decay ventilation studies, and faster response models are ideal for a quick, basic evaluation of human-generated pollution and ventilation adequacy. When properly located, these sensors are also appropriate for continuous monitoring and for control strategies using equilibrium tracer and air fraction tracer calculations.

AMPEROMETRIC ELECTROCHEMICAL CO₂ DETECTORS

Amperometric electrochemical CO₂ sensors ([Figure 12](#)) use a measured current driven between two electrodes by the reduction of CO₂ that diffuses across a porous membrane. Unlike NDIR sensors, which normally last the lifetime of the instrument, electrochemical CO₂ sensors may change in electrolyte chemistry over time (typically 12 to 18 months); so sensors should be replaced periodically. These sensors typically hold their calibration for several weeks, but they may drift more if exposed to low humidity; this drift makes them less suitable for continuous monitoring applications. At low humidity (below 30% rh), the sensors must be kept moist to maintain specified accuracy.

Amperometric electrochemical sensors have a lower power requirement than NDIR sensors, usually operating continuously for weeks where NDIR instruments typically operate for 6 hours (older models) to 150 hours (newer models). The longer battery life can be advantageous for spot checks and walk-throughs and for measuring CO₂ distribution throughout a building and within a zone. Unlike most NDIR sensors, amperometric electrochemical sensors are not affected by high humidity, although readings may be affected if condensate is allowed to form on the sensor.

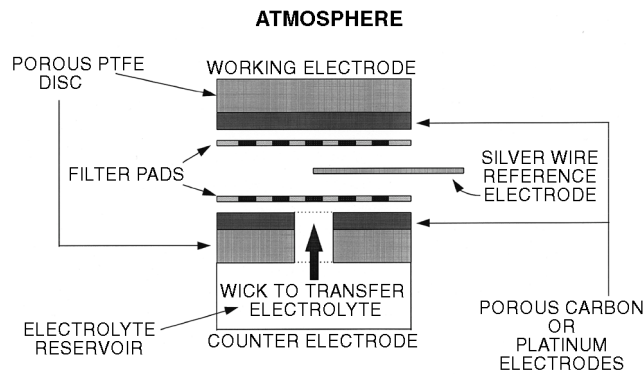


Fig. 12 Amperometric Carbon Dioxide Sensor

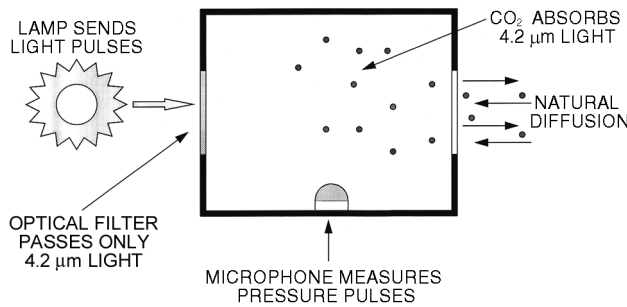


Fig. 13 Open-Cell Photoacoustic Carbon Dioxide Sensor

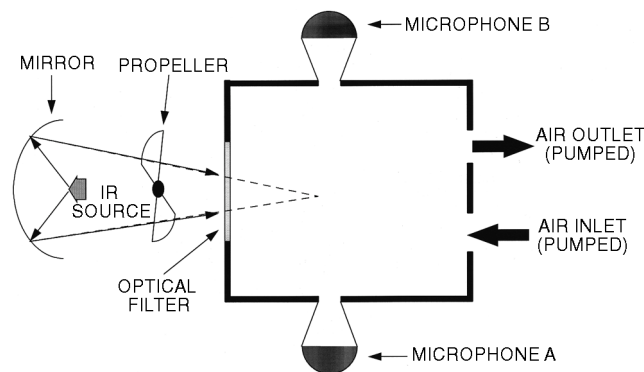


Fig. 14 Closed-Cell Photoacoustic Carbon Dioxide Sensor

PHOTOACOUSTIC CO₂ DETECTORS

Both open- and closed-cell photoacoustic sensors are available.

Open-Cell Sensors

Open-cell photoacoustic CO₂ sensors (Figure 13) operate as air diffuses through a permeable membrane into a chamber that is pulsed with filtered light at the characteristic CO₂ absorption frequency of 4.2 μm. The light energy absorbed by the CO₂ heats the sample chamber, causing a pressure pulse, which is sensed by a piezoresistor. Open-cell photoacoustic CO₂ sensors are presently unavailable in portable instruments, in part because any vibration that might occur during transportation would affect calibration and might affect the signal obtained for a given concentration of CO₂. Ambient acoustical noise may also influence readings. For continuous monitoring, vibration is a concern, as are temperature and airflow cooling effects. However, if a sensor is located properly and if the optical filter is kept

relatively clean, photoacoustic CO₂ sensors may be very stable. Commercially available open-cell photoacoustic transmitters do not allow recalibration to adjust for pressure differences, so an offset should be incorporated in any control system using these sensors at an altitude or duct pressure other than calibration conditions.

Closed-Cell Sensors

Closed-cell photoacoustic sensors (Figure 14) operate under the same principle as the open-cell version, except that samples are pumped into a sample chamber that is sealed and environmentally stabilized. Two acoustic sensors are sometimes used in the chamber to minimize vibration effects. Closed-cell units, available as portable or fixed monitors, come with particle filters that are easily replaced (typically at 3- to 6-month intervals) if dirt or dust accumulates on them. Closed-cell photoacoustic monitors permit recalibration to correct for drift, pressure effects, or other environmental factors that might influence accuracy.

POTENTIOMETRIC ELECTROCHEMICAL CO₂ DETECTORS

Potentiometric electrochemical CO₂ sensors use a porous fluorocarbon membrane that is permeable to CO₂, which diffuses into a carbonic acid electrolyte, changing the electrolyte pH. This pH change is monitored by a pH electrode inside the cell. The pH electrode isopotential drift prohibits long-term monitoring to the accuracy and resolution required for continuous measurement or control or for detailed IAQ evaluations, although accuracy within 100 ppm, achievable short-term over the 2000 ppm range, may be adequate for basic ventilation and odor evaluations. In addition, this type of sensor exhibits slow response, which increases the operator time necessary for field applications or for performing a walk-through of a building.

COLORIMETRIC DETECTOR TUBES

Colorimetric detector tubes contain a chemical compound that discolors in the presence of CO₂ gas, with the amount of discoloration related to the CO₂ concentration. These detector tubes are often used to spot check CO₂ levels; when used properly, they are accurate to within 25%. If numerous samples are taken (i.e., six or more), uncertainty may be reduced; this may render the tubes, if used in the late afternoon, adequate as a very basic determination of odor and discomfort related to human occupation. However, CO₂ detector tubes are generally not appropriate for specific ventilation assessment because of their inaccuracy and inability to record concentration changes over time.

LABORATORY MEASUREMENTS

Laboratory techniques for measuring CO₂ concentration include mass spectroscopy, thermal conductivity, infrared spectroscopy, and gas chromatography. These techniques typically require taking on-site **grab samples** for laboratory analysis. Capital costs for each piece of equipment are high, and significant training is required. A considerable drawback to grab sampling is that CO₂ levels change significantly during the day and over the course of a week, making it sensible to place sensors on site with an instrument capable of recording or data logging measurements continuously over the course of a workweek. An automated grab sampling system capturing many samples of data would be quite cumbersome and expensive if designed to provide CO₂ trend information over time. However, an advantage to laboratory techniques is that they can be highly accurate. A mass spectrometer, for example, can measure CO₂ concentration to within 5 ppm from 0 to 2000 ppm. All laboratory measurement techniques are subject to errors resulting from interfering agents. A gas chromatograph is typically used in conjunction with the mass spectrometer to eliminate interference from

nitrous oxide (N₂O), which has an equivalent mass, if samples are collected in a hospital or in another location where N₂O might be present.

ELECTRIC MEASUREMENT

Ammeters

Ammeters are low-resistance instruments for measuring current. They should be connected in series with the circuit being measured (Figure 15). Ideally, they have the appearance of a short circuit, but in practice, all ammeters have a nonzero input impedance that influences the measurement to some extent.

Ammeters often have several ranges, and it is good practice when measuring unknown currents to start with the highest range and then reduce the range to the appropriate value to obtain the most sensitive reading. Ammeters with range switches maintain circuit continuity during switching. On some older instruments, it may be necessary to short-circuit the ammeter terminals when changing the range.

Current transformers are often used to increase the operating range of ammeters. They may also provide isolation and thus protection from a high-voltage line. Current transformers have at least two separate windings on a magnetic core (Figure 16). The primary winding is connected in series with the circuit in which the current is measured. In the case of a clamp-on probe, the transformer core is actually opened and then connected around a single conductor carrying the current to be measured. That conductor serves as the primary winding. The secondary winding carries a scaled-down version of the primary current, which is connected to an ammeter. Depending on the type of instrument, the ammeter reading may have to be multiplied by the ratio of the transformer.

When using an auxiliary current transformer, the secondary circuit must not be open when current is flowing in the primary winding; dangerous high voltage may exist across the secondary terminals. A short-circuiting blade between the secondary terminals should be closed before the secondary circuit is opened at any point.

Transformer accuracy can be impaired by the residual magnetism in the core when the primary circuit is opened at an instant when the flux is large. The transformer core may be left magnetized, resulting in ratio and phase angle errors. The primary and secondary windings should be short-circuited before making changes.

Voltmeters

Voltmeters are high-resistance instruments that should be connected across the load (in parallel), as shown in Figure 17. Ideally they have the appearance of an open circuit, but in practice all voltmeters have some finite impedance that influences the measurement to some extent.

Voltage transformers are often used to increase the operating range of a voltmeter (Figure 18). They also provide isolation from high voltages and prevent injury to the operator. Like current transformers, voltage transformers consist of two or more windings on a magnetic core. The primary winding is generally connected across the high voltage to be measured, and the secondary winding is connected to the voltmeter. It is important not to short-circuit the secondary winding of a voltage transformer.

Wattmeters

Wattmeters are instruments that measure the active power of an ac circuit, which equals the voltage multiplied by that part of the current in phase with the voltage. There are generally two sets of terminals—one to connect the load voltage and the other to connect in series with the load current. Current and voltage transformers can be

used to extend the range of a wattmeter or to isolate it from high voltage. Figure 19 and Figure 20 show connections for single-phase wattmeters, and Figure 21 shows use of current and voltage transformers with a single-phase wattmeter.

Wattmeters with multiple current and voltage elements are available to measure polyphase power. Polyphase wattmeter connections are shown in Figure 22 and Figure 23.

Power-Factor Meters

Power-factor meters measure the ratio of the active power to the apparent power (product of the voltage and current). The connections for power-factor meters and wattmeters are similar, and current and voltage transformers can be used to extend their range. Connections for single-phase and polyphase power-factor meters are shown in Figure 24 and Figure 25, respectively.

ROTATIVE SPEED MEASUREMENT

Tachometers

Tachometers, or direct-measuring rpm counters, vary from hand-held mechanical or electric meters to shaft-driven and electronic pulse counters. They are used in general laboratory and shop work to check the rotative speeds of motors, engines, and turbines.

Stroboscopes

Optical rpm counters work by producing a controlled high-speed electronic flashing light. The operator directs the light on a rotating member and increases the rate of flashes until the optical effect of stopping rotation of the member is achieved. At this point, the rpm measured is equal to the flashes per minute emitted by the strobe unit. Care must be taken to start at the bottom of the instrument scale and work up because multiples of the rpm produce almost the same optical effect as true synchronism. Multiples can be indicated by positioning suitable marks on the shaft, such as a bar on one side and a circle on the opposite side. If, for example, the two are seen superimposed, then the strobe light is flashing at an even multiple of the true rpm.

AC Tachometer-Generators

A tachometer-generator consists of a rotor and a stator. The rotor is a permanent magnet driven by the equipment. The stator is a winding with a hole through the center for the rotor. Concentricity is not critical; bearings are not required between rotor and stator. The output can be a single-cycle-per-revolution signal whose voltage is a linear function of rotor speed. The polypole configuration that generates 10 cycles per revolution permits measurement of speeds as low as 20 rpm without causing the indicating needle to flutter. The output of the ac tachometer-generator is rectified and connected to a dc voltmeter.

SOUND AND VIBRATION MEASUREMENT

Measurement systems for determining sound pressure, sound intensity, and mechanical vibration generally involve the use of transducers to convert mechanical signals into electrical signals, which are then processed electronically in order to characterize the measured mechanical signals. These measurement systems contain one or more of the following elements, which may or may not be contained in a single instrument:

1. A transducer, or an assembly of transducers, to convert sound pressure, sound intensity, or mechanical vibration (time-varying strain, displacement, velocity, acceleration, or force) into an electrical signal that is quantitatively related to the mechanical quantity being measured.

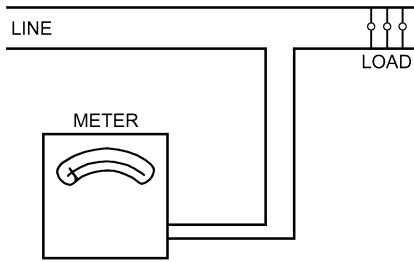


Fig. 15 Ammeter Connected in Power Circuit

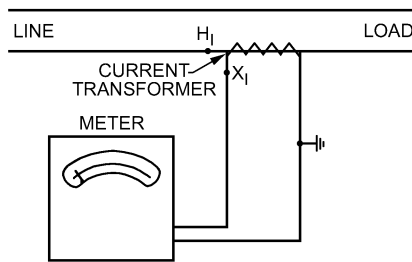


Fig. 16 Ammeter with Current Transformer

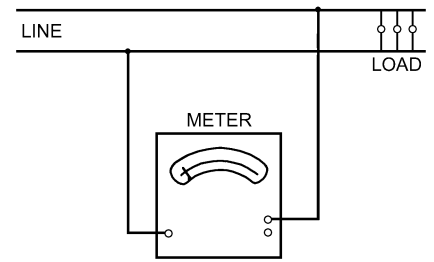


Fig. 17 Voltmeter Connected Across Load

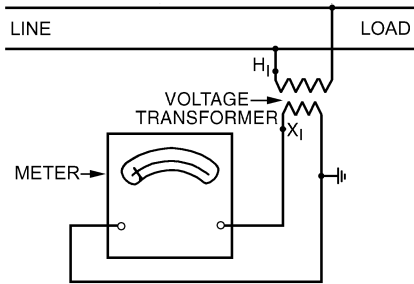


Fig. 18 Voltmeter with Potential Transformer

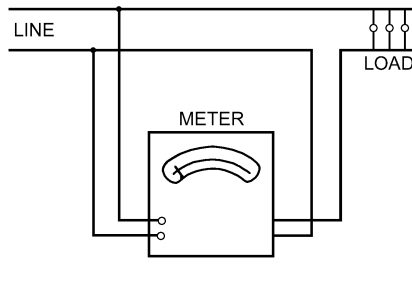


Fig. 19 Wattmeter in Single-Phase Circuit Measuring Power Load plus Loss in Current-Coil Circuit

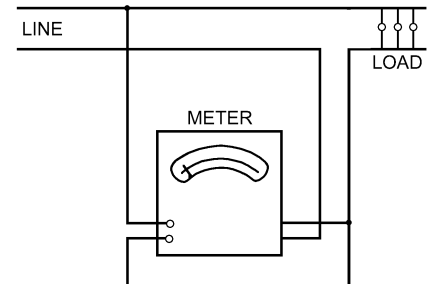


Fig. 20 Wattmeter in Single-Phase Circuit Measuring Power Load plus Loss in Potential-Coil Circuit

2. Amplifiers and networks to provide such functions as electrical impedance match-

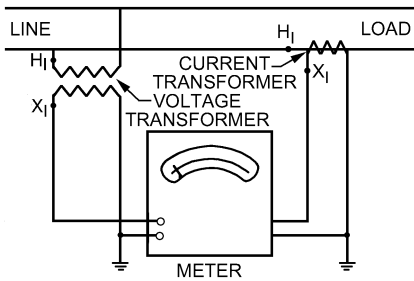


Fig. 21 Wattmeter with Current and Potential Transformer

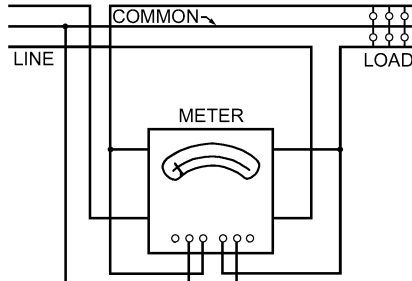


Fig. 22 Polyphase Wattmeter in Two-Phase, Three-Wire Circuit with Balanced or Unbalanced Voltage or Load

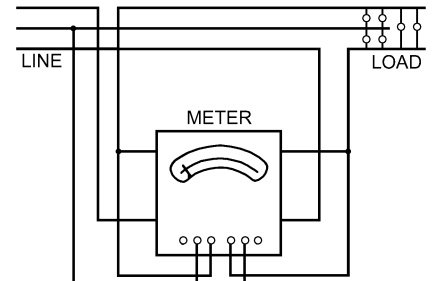


Fig. 23 Polyphase Wattmeter in Three-Phase, Three-Wire Circuit

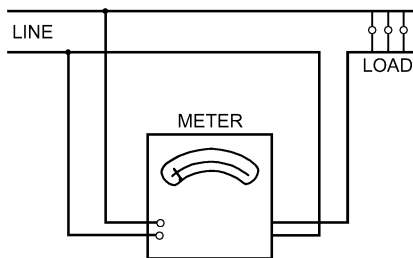


Fig. 24 Single-Phase Power-Factor Meter

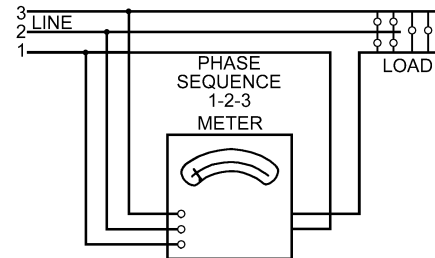


Fig. 25 Three-Wire, Three-Phase Power-Factor Meter

ing, signal conditioning, integration, differentiation, frequency weighting, and gain.

3. Signal-processing equipment to detect and quantify those aspects of the signal that are being measured (peak value, rms value, time-weighted average level, power spectral density, or magnitude or phase of a complex linear spectrum or transfer function).
4. A device such as a meter, oscilloscope, digital display, or level recorder to display the signal or the aspects of it that are being quantified.

The relevant range of sound and vibration signals can vary over more than 12 orders of magnitude in amplitude and more than 8 orders of magnitude in frequency, depending on the application. References on instrumentation, measurement procedures, and signal analysis are given in the section on Bibliography. Product and application notes, technical reviews, and books published by instrumentation manufacturers are an excellent source of additional reference material. See [Chapter 47 of the ASHRAE Handbook—Applications](#) and [Chapter 7](#) of this volume for further information on sound and vibration.

SOUND MEASUREMENT

Microphones

A microphone is an electroacoustical transducer that transforms an acoustical signal into an electrical signal. The two predominant transduction principles used in the measurement of sound (as opposed to broadcasting or recording) are the electrostatic and the piezoelectric. **Electrostatic (capacitor) microphones** are available either as electric microphones, which do not require an external polarizing voltage, or as condenser microphones, which do require an external polarizing voltage, typically in the range of 28 to 200 V (dc). **Piezoelectric microphones** may be manufactured using either natural piezoelectric crystals or poled ferroelectric crystals. The types of response characteristics of measuring microphones are pressure, free field, and random incidence (diffuse field).

A microphone with a uniform pressure response characteristic maintains uniform sensitivity over its operating frequency range when exposed to a sound pressure that is uniform over the surface of the sensing element. A microphone with a uniform free-field response characteristic maintains uniform sensitivity over its operating frequency range when exposed to a plane progressive sound wave at a specified angle of incidence to the surface of the sensing element. A microphone with a uniform random-incidence response characteristic maintains uniform sensitivity over its operating frequency range when exposed to a diffuse sound field.

The sensitivity and the frequency range over which the microphone has uniform sensitivity (flat frequency response) vary with both the diameter (surface area) of the sensing element and the microphone type. Other factors that may critically affect the performance or response of a measuring microphone and preamplifier in a given measurement application are atmospheric pressure, temperature, relative humidity, external magnetic and electrostatic fields, mechanical vibration, and radiation. A microphone should be selected based on its long- and short-term stability; the match between its performance characteristics (e.g., sensitivity, frequency response, amplitude linearity, self-noise) and the expected amplitude of sound pressure, frequency, range of analysis, and expected environmental conditions of measurement; and any other pertinent considerations, such as size and directional characteristics.

Sound Measurement Systems

Microphone preamplifiers, amplifiers, weighting networks (see [Chapter 7](#)), filters, and displays are available either separately or

integrated into a measuring instrument such as a sound level meter, personal noise exposure meter, measuring amplifier, or real-time fractional octave or Fourier [e.g., fast Fourier transform (FFT)] signal analyzer. The instrument(s) included in a sound measurement system depends on the purpose of the measurement and the frequency range and resolution of signal analysis. In the case of community and industrial noise measurements for regulatory purposes, the instrument, signal processing, and quantity to be measured are usually dictated by the pertinent regulation. The optimal set of instruments generally varies for measurement of different characteristics such as sound power in HVAC ducts, sound power emitted by machinery, noise criteria (NC) numbers, sound absorption coefficients, sound transmission loss of building partitions, and reverberation times (T_{60}).

Frequency Analysis

Measurement criteria often dictate the use of filters to analyze the signal in order to indicate the spectrum of the sound being measured. Filters of different bandwidths for different purposes include fractional octave band (one, one-third, one-twelfth, etc.), constant percentage bandwidth, and constant (typically narrow) bandwidth. The filters may be analog or digital and, if digital, may or may not be capable of real-time data acquisition during the measurement period, depending on the bandwidth of frequency analysis. FFT signal analyzers are generally used in situations that require very narrow band signal analysis when the amplitudes of the sound spectra vary significantly with respect to frequency. This may occur in regions of resonance or when it is necessary to identify narrow-band or discrete sine-wave signal components of a spectrum in the presence of other such components or of broadband noise.

Sound Chambers

Special rooms and procedures are required in order to characterize and calibrate sound sources and receivers. The rooms are generally classified into three types—anechoic, semianechoic, and reverberant. The ideal **anechoic** room or chamber would have boundary surfaces that completely absorb sound energy at all frequencies. The ideal **semianechoic** room or chamber would be identical to the ideal anechoic room, except that one surface would totally reflect sound energy at all frequencies. The ideal **reverberant** room or chamber would have boundary surfaces that totally reflect sound energy at all frequencies.

Anechoic chambers are used to perform measurements under conditions approximating those of a free sound field. They can be used in calibrating and characterizing individual microphones, microphone arrays, acoustic intensity probes, reference sound power sources, loudspeakers, sirens, and other individual or complex sources of sound.

Semianechoic chambers are built with a hard reflecting floor in order to accommodate heavy machinery or to simulate large factory floor or outdoor conditions. They can be used in calibrating and characterizing reference sound power sources, obtaining sound power levels of noise sources, and characterizing the sound output of emergency vehicle sirens when mounted on an emergency motor vehicle.

Reverberation chambers are used to perform measurements under conditions approximating those of a diffuse sound field. They can be used in calibrating and characterizing random-incidence microphones and reference sound power sources, obtaining sound power ratings of equipment and sound power levels of noise sources, measuring sound absorption coefficients of building materials and panels, and measuring the transmission loss through building partitions and components such as doors and windows.

Calibration

A measurement system should be calibrated as a system from microphone or probe to indicating device before it is used to per-

form absolute measurements of sound. Acoustic calibrators and pistonphones of fixed or variable frequency and amplitude are available for this purpose. These calibrators should be used at a frequency low enough that the pressure, free-field, and random-incidence response characteristics of the measuring microphone(s) are, for practical purposes, equivalent, or at least related in a known quantitative manner for that specific measurement system. In general, the sound pressure produced by these calibrators may vary, depending on the microphone type, whether the microphone has a protective grid, atmospheric pressure, temperature, and relative humidity. Correction factors and coefficients are required for conditions of use that differ from those existing during the calibration of the acoustic calibrator or pistonphone. For demanding applications, precision sound sources and measuring microphones should periodically be sent to the manufacturer, a private testing laboratory, or a national standards laboratory for calibration.

VIBRATION MEASUREMENT

With the exception of seismic instruments that record or indicate vibration directly via a mechanical or optomechanical device connected to the test surface, vibration measurements involve the use of an electromechanical or interferometric vibration transducer. Here, the term vibration transducer refers to a generic mechanical vibration transducer. Electromechanical and interferometric vibration transducers belong to a large and varied group of transducers that detect mechanical motion and furnish an electrical signal that is quantitatively related to a particular physical characteristic of the motion. Depending on the design of the transducer, the electrical signal may be related to mechanical strain, displacement, velocity, acceleration, or force. The operating principles of vibration transducers may involve optical interference; electrodynamic coupling; piezoelectric (including poled ferroelectric) or piezoresistive crystals; or variable capacitance, inductance, reluctance, or resistance. A considerable variety of vibration transducers with a wide range of sensitivities and bandwidths is commercially available. Vibration transducers may be contacting (e.g., seismic transducers) or non-contacting (e.g., interferometric or capacitive).

Transducers

Seismic transducers use a spring mass resonator within the transducer. At frequencies much greater than the fundamental natural frequency of the mechanical resonator, the relative displacement between the base and the seismic mass of the transducer is nearly proportional to the displacement of the transducer base. At frequencies much lower than the fundamental resonant frequency, the relative displacement between the base and the seismic mass of the transducer is nearly proportional to the acceleration of the transducer base. Therefore, seismic displacement transducers and seismic electrodynamic velocity transducers tend to have a relatively compliant suspension with a low resonant frequency; piezoelectric accelerometers and force transducers have a relatively stiff suspension with a high resonant frequency.

Strain transducers include the metallic resistance gage and the piezoresistive strain gage. For dynamic strain measurements, these are usually of the bonded type, where the gages are bonded directly to the test surface. The accuracy with which a bonded strain gage replicates strain occurring in the test structure is largely a function of how well the strain gage was oriented and bonded to the test surface.

Displacement transducers include the capacitance gage, fringe-counting interferometer, seismic displacement transducer, and linear variable differential transformer (LVDT). Velocity transducers include the reluctance (magnetic) gage, laser Doppler interferometer, and seismic electrodynamic velocity transducer. Accelerometers and force transducers include the piezoelectric, piezoresistive, and force-balance servo.

Vibration Measurement Systems

The sensitivity, frequency limitations, bandwidth, and amplitude linearity of vibration transducers vary greatly with the transduction mechanism and the manner in which the transducer is applied in a given measurement apparatus. The performance of contacting transducers can be significantly affected by the mechanical mounting methods and points of attachment of the transducer and connecting cable and by the mechanical impedance of the structure loading the transducer. Amplitude linearity varies significantly over the operating range of the transducer, with some transducer types or configurations being inherently more linear than others. Other factors that may critically affect the performance or response of a vibration transducer in a given measurement application are temperature; relative humidity; external acoustic, magnetic, and electrostatic fields; transverse vibration; base strain; chemicals; and radiation. A vibration transducer should be selected based on its long- and short-term stability; the match between its performance characteristics (e.g., sensitivity, frequency response, amplitude linearity, self-noise) and the expected amplitude of vibration, frequency range of analysis, and expected environmental conditions of measurement; and any other pertinent considerations (e.g., size, mass, and resonant frequency).

Vibration exciters, or **shakers**, are used in structural analysis, vibration analysis of machinery, fatigue testing, mechanical impedance measurements, and vibration calibration systems. Vibration exciters have a table or moving element with a drive mechanism that may be mechanical, electrodynamic, piezoelectric, or hydraulic. They range from relatively small, low-power units for calibrating transducers such as accelerometers to relatively large, high-power units for structural and fatigue testing.

Conditioning amplifiers, power supplies, preamplifiers, charge amplifiers, voltage amplifiers, power amplifiers, filters, controllers, and displays are available either separately or integrated into a measuring instrument or system, such as a structural analysis system, vibration analyzer, vibration monitoring system, vibration meter, measuring amplifier, multichannel data-acquisition and modal analysis system, or real-time fractional-octave or FFT signal analyzer. The choice of instrument(s) to include in a vibration measurement system depends on the mechanical quantity to be determined, the purpose of the measurement, and the frequency range and resolution of signal analysis. In the case of vibration measurements, the signal analysis is relatively narrow in bandwidth and may be relatively low in frequency in order to accurately characterize structural resonances. Accelerometers with internal integrated circuitry are available to provide impedance matching or servo control for measuring very low frequency acceleration (servo accelerometers). Analog integration and differentiation of vibration signals is available through integrating and differentiating networks and amplifiers, and digital is available through FFT analyzers. Vibration measurements made for different purposes (e.g., machinery diagnostics and health monitoring, balancing rotating machinery, analysis of torsional vibration, analysis of machine-tool vibration, modal analysis, analysis of vibration isolation, stress monitoring, industrial control) will generally each dictate different mechanical measurement requirements and a different optimal set of instrumentation.

Calibration

Because of their inherent long- and short-term stability, amplitude linearity, wide bandwidth, wide dynamic range, low noise, and wide range of sensitivities, seismic accelerometers have traditionally been used as a reference standard for dynamic mechanical measurements. A measurement system should be calibrated as a system from transducer to indicating device before it is used to perform absolute dynamic measurements of mechanical quantities. Calibrated reference vibration exciters, standard reference accelerometers, precision conditioning amplifiers, and precision

calibration exciters are available for this purpose. These exciters and standard reference accelerometers can be used to transfer a calibration to another transducer. For demanding applications, either a calibrated exciter or a standard reference accelerometer with connecting cable and conditioning amplifier should periodically be sent to the manufacturer, a private testing laboratory, or a national standards laboratory for calibration.

LIGHTING MEASUREMENT

Light level, or illuminance, is usually measured with a photocell made from a semiconductor such as silicon or selenium. Such photocells produce an output current proportional to incident luminous flux; when linked with a microammeter, color- and cosine-corrected filters, and multirange switches, they are used in inexpensive hand-held light meters and more precise instruments. Different cell heads allow multirange use in precision meters.

Cadmium sulfide photocells, in which the resistance varies with illumination, are also used in light meters. Both gas-filled and vacuum photoelectric cells are in use.

Small survey-type meters are not as accurate as laboratory meters; their readings should be considered approximate, although consistent, for a given condition. Their range is usually from 5 to 5000 footcandles. Precision low-level meters have cell heads with ranges down to 0 to 2 footcandles.

A photometer installed in a revolving head is called a goniophotometer and is used to measure the distribution of light sources or luminaires. To measure total luminous flux, the luminaire is placed in the center of a sphere painted inside with a high-reflectance white with a near perfect diffusing matte surface. Total light output is measured through a small baffled window in the sphere wall.

To measure irradiation from germicidal lamps, a filter of fused quartz with fluorescent phosphor is placed over the light meter cell.

If meters are used to measure the number of lumens per unit area diffusely leaving a surface, luminance (cd/in^2) instead of illumination (footcandles) is read. Light meters can be used to measure luminance; or electronic lux meters containing a phototube, an amplifier, and a microammeter can read luminance directly. In the case of a perfectly diffuse reflecting surface, which has a constant luminance regardless of viewing angle, the unit of footlamberts in lumens/ ft^2 is sometimes applied.

THERMAL COMFORT MEASUREMENT

Thermal comfort depends on the combined influence of clothing, activity, air temperature, air velocity, mean radiant temperature, and air humidity. Thermal comfort is influenced by heating or cooling of particular body parts. This is due to radiant temperature asymmetry (plane radiant temperature), draft (air temperature, air velocity, turbulence), vertical air temperature differences, and floor temperature (surface temperature).

A general description of thermal comfort is given in [Chapter 8](#), and guidelines for an acceptable thermal environment are given in *ASHRAE Standard 55* and *ISO Standard 7730*. *ASHRAE Standard 55* also includes required measuring accuracy. In addition to specified accuracy, *ISO Standard 7726* includes recommended measuring locations and a detailed description of instruments and methods.

Clothing and Activity Level

These values are estimated from tables ([Chapter 8](#), *ISO Standard 9920*, *ISO Standard 8996*). The thermal insulation of clothing (clo-

value) can be measured on a thermal mannequin (McCullough et al. 1985, Olesen 1985). The activity (met-value) can be estimated from measuring CO_2 and O_2 in a person's expired air.

Air Temperature

Various types of thermometers may be used to measure air temperature. Placed in a room, the sensor registers a temperature between air temperature and mean radiant temperature. One way of reducing the radiant error is to make the sensor as small as possible because the convective heat transfer coefficient increases as the size decreases while the radiant heat transfer coefficient is constant. A smaller sensor also provides a favorably low time constant. The radiant error can also be reduced by using a shield (an open, polished aluminum cylinder) around the sensor, by using a sensor with a low-emittance surface, or by artificially increasing the air velocity around the sensor (aspirating air through a tube in which the sensor is placed).

Air Velocity

In occupied zones, air velocities are usually small (0 to 100 fpm) but have an effect on human thermal sensation. Because the velocity fluctuates, the mean value should be measured over a suitable period, typically 3 min. Velocity fluctuations with frequencies up to 1 Hz significantly increase human discomfort due to draft, which is a function of air temperature, mean air velocity, and turbulence (see [Chapter 8](#)). The fluctuations can be given as the standard deviation of the air velocity over the measuring period (3 min) or as the turbulence intensity (standard deviation divided by mean air velocity). Velocity direction may change and is difficult to identify at low air velocities. An omnidirectional sensor with a short response time should be used. A thermal anemometer is suitable. If a hot-wire anemometer is used, the direction of the flow being measured must be perpendicular to the hot wire. Smoke puffs can be used to identify the direction.

Plane Radiant Temperature

This refers to the uniform temperature of an enclosure in which the radiant flux on one side of a small plane element is the same as in the actual nonuniform environment. It describes the radiation in one direction. The plane radiant temperature can be calculated from the surface temperatures of the environment (half-room) and the angle factors between the surfaces and a plane element (*ASHRAE Standard 55*). The plane radiant temperature may also be measured by a net-radiometer or a radiometer with a sensor consisting of a reflective disk (polished) and an absorbent disk (painted black) (Olesen et al. 1989).

Mean Radiant Temperature

This is the uniform temperature of an imaginary black enclosure in which an occupant would exchange the same amount of radiant heat as in the actual nonuniform enclosure. The mean radiant temperature can be calculated from measured surface temperatures and the corresponding angle factors between the person and the surfaces ([Chapter 8](#)).

The mean radiant temperature can also be determined from the plane radiant temperature in six opposite directions weighted according to the projected area factors for a person ([Chapter 8](#)).

Because of its simplicity, the instrument most commonly used to determine the mean radiant temperature is a **black globe thermometer** (Vernon 1932, Bedford and Warner 1935). This thermometer consists of a hollow sphere usually 6 in. in diameter coated in flat black paint with a thermocouple or thermometer bulb at its center. The temperature assumed by the globe at equilibrium results from a balance between heat gained and lost by radiation and convection.

The mean radiant temperatures are calculated from

$$\bar{t}_r = \left[(t_g + 460)^4 + \frac{4.74 \times 10^7 V_a^{0.6}}{\varepsilon D^{0.4}} (t_g - t_a) \right]^{1/4} - 460 \quad (11)$$

where

- \bar{t}_r = mean radiant temperature, °F
- t_g = globe temperature, °F
- V_a = air velocity, fpm
- t_a = air temperature, °F
- D = globe diameter, ft
- ε = emissivity (0.95 for black globe)

According to Equation (11), air temperature and air velocity around the globe must also be determined. The globe thermometer is spherical, while mean radiant temperature is defined in relation to the human body. For sedentary people, the globe represents a good approximation. For people who are standing, the globe, in a radiant nonuniform environment, overestimates the radiation from floor or ceiling. An ellipsoid-shaped sensor gives a closer approximation to the human shape. A black globe will also overestimate the influence of short-wave radiation (e.g., sunshine). A flat gray color better represents the radiant characteristic of normal clothing (Olesen et al. 1989). The hollow sphere is usually made of copper, which results in an undesirable high time constant. This can be overcome by using lighter materials (e.g., a thin plastic bubble).

Air Humidity

The water vapor pressure (absolute humidity) is usually uniform in the occupied zone of a space; therefore, it is sufficient to measure absolute humidity at one location. Many of the instruments listed in Table 3 are applicable. At ambient temperatures that provide comfort or slight discomfort, the thermal effect of humidity is only moderate, and highly accurate humidity measurements are unnecessary.

CALCULATING THERMAL COMFORT

When the thermal parameters have been measured, their combined effect can be calculated by the thermal indices in Chapter 8. For example, the effective temperature (Gagge et al. 1971) can be determined from air temperature and humidity. Based on the four environmental parameters and an estimation of clothing and activity, the **predicted mean vote (PMV)** can be determined with the aid of tables (Fanger 1982, ISO Standard 7730, Chapter 8). The PMV is an index predicting the average thermal sensation that a group of occupants may experience in a given space.

For certain types of normal activity and clothing, the environmental parameters measured can be compared directly with those described in ASHRAE Standard 55 or ISO Standard 7730.

INTEGRATING INSTRUMENTS

Several instruments have been developed to evaluate the combined effect of two or more thermal parameters on human comfort. Madsen (1976) developed an instrument that gives information on the occupants' expected thermal sensation by direct measurement of the PMV value. The comfort meter has a heated ellipsoid-shaped sensor that simulates the body (Figure 26). The estimated clothing (insulation value), activity in the actual space, and humidity are set on the instrument. The sensor then integrates the thermal effect of the air temperature, mean radiant temperature, and air velocity in approximately the same way the body does. The electronic instrument gives the measured operative and equivalent temperature, calculated PMV, and predicted percentage of dissatisfied (PPD).

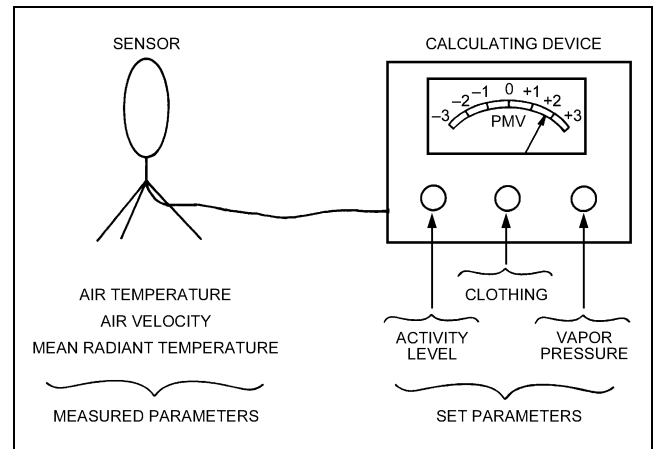


Fig. 26 Madsen's Comfort Meter
(Madsen 1976)

MOISTURE CONTENT AND TRANSFER MEASUREMENT

Little off-the-shelf instrumentation exists to measure the moisture content of porous materials or the moisture transfer through those materials. However, many measurements can be set up with a small investment of time and money. Three moisture properties are most commonly sought—(1) the sorption isotherm, a measure of the amount of water vapor a hygroscopic material will adsorb from humid air; (2) vapor permeability, a measure of the rate at which water vapor will pass through a given material; and (3) liquid diffusivity, a measure of the rate at which liquid water will pass through a porous material.

Sorption Isotherm

A sorption isotherm relates the **equilibrium moisture content (EMC)** of a hygroscopic material to the ambient relative humidity under conditions of constant temperature. Moisture content is the ratio of the total mass of water in a sample to the dry mass of the sample. Determining a sorption isotherm involves exposing a sample of material to a known relative humidity at a known temperature and then measuring the sample's moisture content after a sufficient period of time has elapsed for the sample to reach equilibrium with its surroundings. Hysteresis in the sorption behavior of most hygroscopic materials requires that measurements be made both for increasing relative humidity (the adsorption isotherm) and for decreasing relative humidity (the desorption isotherm).

The ambient relative humidity can be controlled using saturated salt solutions or mechanical refrigeration equipment (Tveit 1966, Cunningham and Sprott 1984, Carotenuto et al. 1991). Precise measurements of the relative humidity produced by various salt solutions have been reported by Greenspan (1977). ASTM Standard E 104 describes the use of saturated salt solutions. The EMC of a sample is usually determined gravimetrically using a precision balance. The sample dry mass, necessary to calculate moisture content, can be found by oven drying or desiccant drying. Oven dry mass may be lower than desiccant dry mass because of the loss of volatiles other than water in the oven (Richards et al. 1992).

A major difficulty in the measurement of the sorption isotherms of engineering materials is the long time required for many materials to reach equilibrium—often as long as weeks or months. The rate-limiting mechanism for these measurements is usually the slow process of vapor diffusion into the pores of the material. The use of smaller samples can help reduce the diffusion time.

Vapor Permeability

The diffusive transfer of water vapor through porous materials is often described by a modified form of Fick's law:

$$w''_v = -\mu \frac{dp}{dx} \quad (12)$$

where

- w''_v = mass of vapor diffusing through unit area in unit time, gr/h·ft²
- dp/dx = vapor pressure gradient, in. Hg/in.
- μ = vapor permeability, gr·in/h·ft²·in. Hg

In engineering practice, permeance may be used instead of permeability. **Permeance** is simply permeability divided by the material thickness in the direction of the flow of vapor; thus, while permeability is a material property, permeance depends on thickness.

Measurement of permeability is made with wet-cup, dry-cup, or modified cup tests. Wet- and dry-cup tests are described in [Chapter 23](#).

For many engineering materials, vapor permeability is a strong function of mean relative humidity. Wet and dry cups cannot adequately characterize this dependence on relative humidity. Instead, a modified cup method can be used (McLean et al. 1990, Burch et al. 1992). In the modified cup method, the pure water or desiccant within a cup is replaced with a saturated salt solution. A second saturated salt solution is used to condition the environment external to the cup. With such an arrangement, the relative humidities on both sides of the sample material can be varied from 0 to 100%. Several cups with a range of mean relative humidities are used to map out the dependence of vapor permeability on relative humidity.

In measuring materials of high permeability, the finite rate of vapor diffusion through air in the cup may become a factor. The air-film resistance could then be a significant fraction of the resistance to vapor flow presented by the sample material. An accurate measurement of high-permeability materials may require an accounting of diffusive rates across all air gaps (Fanney et al. 1991).

Liquid Diffusivity

The transfer of liquid water through porous materials may be characterized as a diffusion-like process:

$$w''_l = -\rho D_l \frac{dy}{dx} \quad (13)$$

where

- w''_l = mass of liquid transferred through unit area per unit time, lb/h·ft²
- ρ = liquid density, lb/ft³
- D_l = liquid diffusivity, ft²/h
- dy/dx = moisture content gradient, ft⁻¹

D_l typically shows a strong dependence on moisture content.

Transient measurement methods deduce the functional form of D_l by observing the evolution of a one-dimensional moisture content profile over time. An initially dry specimen is brought into contact with liquid water. The free water will migrate into the specimen, drawn in by surface tension. The resulting moisture content profile, which changes with time, must be differentiated to find the liquid diffusivity of the material (Bruce and Klute 1956).

Determining the transient moisture content profile typically involves the use of a noninvasive and nondestructive method of measuring local moisture content. Gamma ray absorption (Freitas et al. 1991, Kumaran and Bomberg 1985, Quenard and Sallee 1989), X-ray radiography (Ambrose et al. 1990), neutron radiography (Prazak et al. 1990), and nuclear magnetic resonance (NMR) (Gummerson et al. 1979) have all been employed.

Uncertainty in the resulting measurement of the liquid diffusivity is often large because of the necessity to differentiate noisy experimental data.

HEAT TRANSFER THROUGH BUILDING MATERIALS

Thermal Conductivity

The thermal conductivity of a heat insulator, as defined in [Chapter 23](#), is a unit heat transfer factor. Two methods of determining the thermal conductivity of flat insulation are the **guarded hot plate** and the **heat flow meter apparatus**, according to ASTM *Standards* C 177 and C 518, respectively. Both methods use parallel isothermal plates to induce a steady temperature gradient across the thickness of the specimen(s). The guarded hot plate is considered an absolute method for determining thermal conductivity. The heat flow meter apparatus requires calibration with a specimen having a known thermal conductivity, usually determined in the guarded hot plate. The heat flow meter apparatus is calibrated by determining the voltage output of its heat flux transducer(s) as a function of the heat flux through the transducer(s).

The basic design of the guarded hot plate consists of an electrically heated plate and two liquid-cooled plates. Two similar specimens of a material are required for a test; one is mounted on each side of the hot plate. A cold plate is then pressed against the outside of each specimen by a clamp screw. The heated plate consists of two sections separated by a small gap. During tests, the central (metering) section and the outer (guard) section are maintained at the same temperature to minimize errors caused by edge effects. The electric energy required to heat the metering section is measured carefully and converted to heat flow. The thermal conductivity of the material can be calculated under steady-state conditions using this heat flow quantity, the area of the metering section, the temperature gradient, and the specimen thickness. The thermal conductivity of cylindrical or pipe insulation ([Chapter 23](#)) is determined in a similar manner, but an equivalent thickness must be calculated to account for the cylindrical shape (ASTM *Standard* C 335). Transient methods have been developed by D'Eustachio and Schreiner (1952), Hooper and Lepper (1950), and Hooper and Chang (1953) using a line heat source within a slender probe. These instruments are available commercially and have the advantages of rapidity and a small test specimen requirement. The probe is a useful research and development tool, but it has not been as accepted as the guarded hot plate, heat flow meter apparatus, or pipe insulation apparatus.

Thermal Conductance and Resistance

Thermal conductances (C-factors) and resistances (R-values) of many building assemblies can be calculated from the conductivities and dimensions of their components, as described in [Chapter 25](#). Test values can also be determined experimentally using large specimens representative of the building assemblies tested in the hot box apparatus described in ASTM *Standards* C 236 and C 976. This laboratory apparatus allows measurement of heat transfer through a specimen under controlled air temperature, air velocity, and radiation conditions. It is specially suited for large nonhomogeneous specimens.

For in situ measurements, heat flux and temperature transducers are useful in measuring the dynamic or steady-state behavior of opaque building components (ASTM *Standard* C 1046). A heat flux transducer is simply a differential thermopile within a core or substrate material. Two types of construction are used: (1) multiple thermocouple junctions wrapped around a core material, or (2) printed circuits with a uniform array of thermocouple junctions. The transducer is calibrated by determining its voltage output as a function of the heat flux through the transducer. For in situ measurements, the transducer is installed in either the wall or roof, or mounted on an exterior surface with tape or adhesive. The data obtained can be used to compute the thermal conductance or resistance of the building component (ASTM *Standard* C 1155).

AIR CONTAMINANT MEASUREMENT

Three measures of particulate air contamination include the number, projected area, and mass of particles per unit volume of air (ASTM Volume 11.03). Each requires an appropriate sampling technique.

Particles are counted by capturing them in impingers, impactors, membrane filters, or thermal or electrostatic precipitators. Counting may be done by microscope, using stage counts if the sample covers a broad range of sizes.

Electronic particle counters can give rapid data on particle size distribution and concentration. However, their accuracy depends on careful calibration, appropriate maintenance, and proper application. Particle counters have been used in indoor office environments as well as in clean rooms.

Projected area determinations are usually made by sampling onto a filter paper and comparing the light transmitted or scattered by this filter to a standard filter. The staining ability of dusts depends on the projected area and refractive index per unit volume. For sampling, filters must collect the minimum sized particle of interest. In this respect, membrane or glass fiber filters are recommended.

To determine particle mass, a measured quantity of air is drawn through filters, preferably of membrane or glass fiber, and the filter mass is compared to the mass before sampling. Electrostatic or thermal precipitators and various impactors have also been used. For further information, see ACGIH (1983), Lundgren et al. (1979), and Lodge (1989).

[Chapter 45 of the ASHRAE Handbook—Applications](#) presents information on measuring and monitoring gaseous contaminants. Relatively costly analytical equipment, which must be calibrated and operated carefully by experienced personnel, is needed. Numerous methods of sampling the contaminants, as well as the laboratory analysis techniques used after sampling, are specified. Some of the analytical methods are specific to a single pollutant; others are capable of presenting a concentration spectrum for many compounds simultaneously.

COMBUSTION ANALYSIS

Two approaches are used to measure the thermal output or capacity of a boiler, furnace, or other fuel-burning device. The direct or **calorimetric test** measures change in enthalpy or heat content of the fluid, air, or water heated by the device, and multiplies this by the flow rate to arrive at the unit's capacity. The indirect test or **flue gas analysis** method determines the heat losses in the flue gases and the jacket and deducts them from the heat content (higher heating value) of the measured fuel input to the appliance. A **heat balance** simultaneously applies both tests to the same device. The indirect test usually indicates the greater capacity, and the difference is credited to radiation from the casing or jacket and unaccounted-for losses.

With small equipment, the expense of the direct test is usually not justified, and the indirect test is used with an arbitrary radiation and unaccounted-for loss factor.

FLUE GAS ANALYSIS

The flue gases from burning fossil fuels generally contain carbon dioxide (CO₂), water, and hydrogen (H₂) with some small amounts of carbon monoxide (CO), nitrogen oxides (NO_x), sulfur oxides (SO_x), and unburned hydrocarbons. However, generally only CO₂ (or O₂) and CO are measured to determine completeness of combustion and efficiency.

In the laboratory, the instruments most commonly used to measure CO and CO₂ are nondispersive infrared (NDIR) analyzers. The NDIR instruments have several advantages: (1) they are not very

sensitive to flow rate, (2) no wet chemicals are required, (3) they have a relatively fast response, (4) measurements can be made over a wide range of concentrations, and (5) they are not sensitive to the presence of contaminants in the ambient air.

In the laboratory, oxygen is generally measured with an instrument that makes use of the paramagnetic properties of oxygen. The paramagnetic instruments are generally used because of their excellent accuracy and because they can be made specific to the measurement of oxygen.

For field testing and burner adjustment, portable combustion testing equipment is available. These instruments generally measure O₂ and CO with electrochemical cells. The CO₂ is then calculated by an on-board microprocessor and, together with temperature, is used to calculate thermal efficiency. If a less expensive approach is required, a portable Orsat apparatus can be used to measure CO₂, and a length-of-stain tube to measure CO.

DATA ACQUISITION AND RECORDING

Almost every type of transducer and sensor is available with the necessary interface system to make it computer-compatible. The transducer itself begins to lose its identity when integrated into a system that incorporates such features as linearization, offset correction, self-calibration, and so forth. This has eliminated the concern regarding the details of signal conditioning and amplification of basic transducer outputs. The personal computer is integrated into every aspect of data recording, including sophisticated graphics, acquisition and control, and analysis. Modems connected to the Internet or an internal network allow easy access to remote personal computer-based data-recording systems from virtually any locale.

Other means of recording, such as chart recorders, which can be either multipurpose or specifically designed for a given sensor, are available. Chart recorders provide a visual indication and a hard copy record of the data. Rarely is the output of a chart recorder used to process data. Simple indicators and readouts are used mostly to monitor the output of a sensor visually. In most situations, analog indicators such as d'Arsonval movement meters have been rendered obsolete by modern digital indicators. Industrial environments commonly employ signal transmitters for control or computer data-handling systems to convert the signal output of the primary sensor into a compatible common signal span (e.g., the standard 4-20 mA current loop). All signal conditioning (ranging, zero suppression, reference-junction compensation) is provided at the transmitter. Thus, all recorders and controllers in the system can have an identical electrical span, with variations only in charts and scales offering the advantages of interchangeability and economy in equipment cost. Long signal transmission lines can be used, and receiving devices can be added to the loop without degrading performance.

The vast selection of available hardware, an often confusing set of terminology, and the challenge of optimizing the performance/cost ratio for a specific application make the task of configuring a data acquisition system difficult. A system specifically configured to meet a particular measurement need can quickly become obsolete if it has inadequate flexibility. Memory size, recording speed, and signal processing capability are major considerations in determining the correct recording system. Thermal, mechanical, electromagnetic interference, portability, and meteorological factors also influence the selection.

Digital Recording

A digital data acquisition system must contain an interface, which is a system involving one or several analog-to-digital converters, and in the case of multichannel inputs, circuitry for multiplexing. The interface may also provide excitation for transducers, calibration, and conversion of units. The digital data are arranged

into one or several standard digital bus formats. Many data acquisition systems are designed to acquire data rapidly and store large records of data for later recording and analysis. Once the input signals have been digitized, the digital data are essentially immune to noise and can be transmitted over great distances.

Information is transferred to a computer/recorder from the interface as a pulse train, which can be transmitted as 4-, 8-, 12-, 16-, or 32-bit words. An 8-bit word is a byte; many communications methods are rated according to their bytes per second transfer rate. Digital data are transferred in either serial or parallel mode. Serial transmission means that the data are sent as a series of pulses, one bit at a time. Although slower than parallel systems, serial interfaces require only two wires, which lowers their cabling cost. The speed of serial transmissions is rated according to the symbols per second rate, or baud rate. In parallel transmission, the entire data word is transmitted at one time. To do this, each bit of a data word has to have its own transmission line; other lines are needed for clocking and control. Parallel mode is used for short distances or when high data transmission rates are required. Serial mode must be used for long-distance communications where wiring costs are prohibitive.

The two most popular interface bus standards currently used for data transmission are the IEEE 488, or general-purpose interface bus (GPIB), and the RS232 serial interface. The IEEE 488 bus system feeds data down eight parallel wires, one data byte at a time. This parallel operation allows it to transfer data rapidly at up to 1 million characters per second. However, the IEEE 488 bus is limited to a cable length of 65 ft and requires an interface connection on every meter for proper termination. The RS232 system feeds data serially down two wires, one bit at a time. An RS232 line may be over 1000 ft long. For longer distances, it may feed a modem to send data over standard telephone lines. A local area network (LAN) may be available in a facility for transmitting information. With appropriate interfacing, transducer data are available to any computer connected to the network.

Bus measurements can greatly simplify three basic applications—data gathering, automated limit testing, and computer-controlled processes. In data gathering applications, readings are collected over time. The most common applications include aging tests in quality control, temperature tests in quality assurance, and testing for intermittents in service. A controller can monitor any output indefinitely and then display the data directly on its screen or record it on magnetic tape or disks for future use.

In automated limit testing, the computer simply compares each measurement with programmed limits. The controller converts the readings to a good/bad readout. Automatic limit testing becomes highly cost-effective when working with large number of parameters of a particular unit under test.

In computer-controlled processes, the IEEE 488 bus system becomes a permanent part of a larger, completely automated system. For example, a large industrial process may require many electrical sensors that feed a central computer controlling many parts of the manufacturing process. An IEEE 488 bus controller collects readings from several sensors and saves the data until asked to dump an entire batch of readings to a larger central computer at one time. Used in this manner, the IEEE 488 bus controller serves as a slave of the central computer.

Dynamic range and accuracy are two important parameters that must be considered in a digital recording system. **Dynamic range** refers to the ratio of the maximum input signal for which the system is useful to the noise floor of the system. The **accuracy** figure for a system is impacted by the signal noise level, nonlinearity, temperature, time, crosstalk, and so forth. In selecting an 8-, 12-, or 16-bit analog-to-digital converter, the designer cannot assume that the system accuracy will necessarily be determined by the resolution of the encoders (i.e., 0.4%, 0.025%, and 0.0016%, respectively). If the sensor preceding the converter is limited to 1% full-scale accuracy, for example, no significant benefits are gained by using a 12-bit system

over an 8-bit system and suppressing the least significant bit. However, a greater number of bits may be required to cover a larger dynamic range.

Data Logging Devices

Data loggers digitally store electrical signals (analog or digital) to an internal memory storage component. The signal from connected sensors is typically stored to memory at timed intervals ranging from MHz to hourly sampling. Some data loggers store data based on an event (e.g., button push, contact closure). Many data loggers can perform linearization, scaling, or other signal conditioning and permit logged readings to be either instantaneous or averaged values. Most data loggers have built-in clocks that record the time and date together with transducer signal information. Data loggers range from single-channel input to 256 or more channels. Some are general-purpose devices that will accept a multitude of analog and/or digital inputs, while others are more specialized to a specific measurement (e.g., a portable anemometer with built-in data logging capability) or for a specific application (e.g., a temperature, relative humidity, CO₂, and CO monitor with data logging for IAQ applications). Stored data are generally downloaded using a serial interface with a temporary direct connection to a personal computer. Remote data loggers may also download via modem through telephone lines. Some data loggers are designed to allow downloading directly to a printer.

With the reduction in size of personal computers (laptops, notebooks, hand-held PCs, and palmtops), the computer itself is now being used as the data logger. These “mobile” computers may be left in the field storing measurements from sensors directly interfaced into the computer.

Chart Recorders

Chart recorders convert electrical signals (analog or digital) to records of the data versus time on a hard copy, usually paper. Mechanical styluses use ink, hot wire, pressure, or electrically sensitive paper to provide a continuous trace. They are useful up to a few hundred hertz. Thermal and ink recorders are confined to chart speeds of a few inches per second for recording relatively slow processes. Newer advances in portable recorders provide multichannel inputs and up to 25 kHz real-time frequency response without using a pen or pen motor. Both *x-y* recorders and plotters allow two variables to be recorded with respect to one another. Their response times are generally limited to that of thermal and ink recorders. Oscillographic recorders have largely been made obsolete by digital oscilloscopes.

STANDARDS

- ASA. 1980. Techniques of machinery vibration measurement. *ANSI Standard S2.17-80* (R 1986). Acoustical Society of America, New York.
- ASA. 1984. Mechanical vibration of rotating and reciprocating machinery—Requirements for instruments for measuring vibration severity. *ANSI Standard S2.40-84* (R 1990).
- ASA. 1984. Specification for acoustical calibrators. *ANSI Standard S1.40-84* (R 1994).
- ASA. 1985. Statistical methods for determining and verifying stated noise emission values of machinery and equipment. *ANSI Standard S12.3-85* (R 1990).
- ASA. 1987. Methods for determination of insertion loss of outdoor noise barriers. *ANSI Standard S12.8-87*.
- ASA. 1989. Guide to the mechanical mounting of accelerometers. *ANSI Standard S2.61-89* (R 1991).
- ASA. 1989. Method for the designation of sound power emitted by machinery and equipment. *ANSI Standard S12.23-89*.
- ASA. 1989. Reference quantities for acoustical levels. *ANSI Standard S1.8-89*.
- ASA. 1990. Survey methods for the determination of sound power levels of noise sources. *ANSI Standard S12.36-90*.

- ASA. 1990. Vibrations of buildings—Guidelines for the measurements of vibrations and evaluation of their effects on buildings. *ANSI Standard S2.47-90*.
- ASA. 1995. Measurement of sound pressure levels in air. *ANSI Standard S1.13-95*.
- ASHRAE. 1984. Standard method for measurement of flow of gas. *ANSI/ASHRAE Standard 41.7-1984 (RA 91)*.
- ASHRAE. 1985. Laboratory methods of testing fans for rating. *ANSI/ASHRAE Standard 51-1985*, also *ANSI/AMCA Standard 210-85*.
- ASHRAE. 1986. Engineering analysis of experimental data. *Guideline 2-1986 (RA 96)*.
- ASHRAE. 1986. Laboratory method of testing in-duct sound power measurement procedure for fans. *ANSI/ASHRAE Standard 68-1986*, also *ANSI/AMCA Standard 330-86*.
- ASHRAE. 1986. Standard method for temperature measurement. *ANSI/ASHRAE Standard 41.1-1986 (RA 91)*.
- ASHRAE. 1987. Standard methods for laboratory air flow measurement. *ANSI/ASHRAE Standard 41.2-1987 (RA 92)*.
- ASHRAE. 1988. Practices for measurement, testing, adjusting, and balancing of building heating, ventilation, air-conditioning and refrigeration systems. *ANSI/ASHRAE Standard 111-1988*.
- ASHRAE. 1988. A standard calorimeter test method for flow measurement of a volatile refrigerant. *ANSI/ASHRAE Standard 41.9-1988*.
- ASHRAE. 1989. Standard method for pressure measurement. *ANSI/ASHRAE Standard 41.3-1989*.
- ASHRAE. 1989. Standard methods of measurement of flow of liquids in pipes using orifice flowmeters. *ANSI/ASHRAE Standard 41.8-1989*.
- ASHRAE. 1992. Thermal environmental conditions for human occupancy. *ANSI/ASHRAE Standard 55-1992*.
- ASHRAE. 1994. Standard method for measurement of moist air properties. *ANSI/ASHRAE Standard 41.6-1994*.
- ASHRAE. 1996. Method for measurement of proportion of lubricant in liquid refrigerant. *Standard 41.4-1996*.
- ASME. 1972. Application of fluid meters, Part II, 6th ed. (Interim Supplement 19.5 on Instruments and Apparatus) *Standard PTC 19.5-72*. American Society of Mechanical Engineers, New York.
- ASME. 1974. Temperature measurement instruments and apparatus. *ANSI/ASME Standard PTC 19.3-74 (R 1986)*.
- ASME. 1983. Measurement uncertainty for fluid flow in closed conduits. *ANSI/ASME Standard MFC-2M-83*.
- ASME. 1985. Part I: Measurement uncertainty instruments and apparatus. *ANSI/ASME Standard PTC 19.1-85*.
- ASME. 1988. Measurement of liquid flow in closed conduits by weighing methods. *ANSI/ASME Standard MFC-9M-88*.
- ASME. 1989. Measurement of fluid flow in pipes using orifice, nozzle, and venturi. *Standard MFC-3M-85*.
- ASME. 1991. Gauges—Pressure indicating dial type—Elastic elements. *ANSI/ASME Standard B40.1-91*.
- ASME. 1991. Glossary of terms used in the measurement of fluid flow in pipes. *ANSI/ASME Standard MFC-1M-91*.
- ASME. 1994. Method for establishing installation effects on flowmeters. *ANSI/ASME Standard MFC-10M-94*.
- ASTM. 1985. Standard practice for maintaining constant relative humidity by means of aqueous solutions. *Standard E 104-85 (R 1996)*. American Society for Testing and Materials, West Conshohocken, PA.
- ASTM. 1989. Standard test method for steady-state thermal performance of building assemblies by means of a guarded hot box. *Standard C 236-89 (1993)e1*.
- ASTM. 1990. Standard practice for thermographic inspection of insulation installations in envelope cavities of frame buildings. *Standard C 1060-90 (1997)e1*.
- ASTM. 1990. Standard test method for thermal performance of building assemblies by means of a calibrated hot box. *Standard C 976-90 (1996)e1*.
- ASTM. 1993. Temperature-electromotive force (EMF) tables for standardized thermocouples. *Standard E 230-93*.
- ASTM. 1995. Standard practice for in-situ measurement of heat flux and temperature on building envelope components. *Standard C 1046-95*.
- ASTM 1995. Standard test method for determining air change in a single zone by means of gas dilution. *Standard E 741-95*.
- ASTM. 1995. Standard test method for steady-state heat transfer properties of horizontal pipe insulation. *Standard C 335-95*.
- ASTM. 1995. Standard practice for determining thermal resistance of building envelope components from the in-situ data. *Standard C 1155-95*.
- ASTM. 1995. Standard test methods for water vapor transmission of materials. *Standard E 96-95*.
- ASTM. 1997. Standard test method for steady-state heat flux measurements and thermal transmission properties by means of the guarded-hot-plate apparatus. *Standard C 177-97*.
- ASTM. 1998. Standard test method for steady-state thermal transmission properties by means of the heat flow meter apparatus. *Standard C 518-98*.
- ASTM. 2000. *Atmospheric analysis; occupational health and safety; protective clothing*. Vol. 11.03. (182 standards).
- ISO. 1977. Measurement of fluid flow in closed conduits—Velocity area method using Pitot static tubes. *Standard 3966*. International Organization for Standardization, Geneva.
- ISO. 1985. Thermal environments—Instruments and methods for measuring physical quantities. *Standard 7726*.
- ISO. 1990. Ergonomics—Determination of metabolic heat production. *Standard 8996*.
- ISO. 1994. Moderate thermal environments—Determination of the PMV and PPD indices and specification of the conditions for thermal comfort. *Standard 7730*.
- ISO. 1995. Ergonomics of the thermal environment—Estimation of the thermal insulation and evaporative resistance of a clothing ensemble. *Standard 9920*.

REFERENCES

- ACGIH. 1983. *Air sampling instruments for evaluation of atmospheric contaminants*, 6th ed. American Conference of Governmental Industrial Hygienists, Cincinnati, OH.
- ASTM. 1993. *Manual on the use of thermocouples in temperature measurement*. Manual 12.
- Ambrose, J.H., L.C. Chow, and J.E. Beam. 1990. Capillary flow properties of mesh wicks. *AIAA Journal of Thermophysics* 4:318-24.
- Amdur, E.J. 1965. Two-pressure relative humidity standards. In *Humidity and Moisture* 3:445. Reinhold Publishing Corporation, New York.
- Bedford, T. and C.G. Warmer. 1935. The globe thermometer in studies of heating and ventilating. *Journal of the Institution of Heating and Ventilating Engineers* 2:544.
- Benedict, R.P. 1984. *Fundamentals of temperature, pressure and flow measurements*, 3rd ed. John Wiley and Sons, New York.
- Bentz, D.P. and J.W. Martin. 1987. Using the computer to analyze coating defects. *Journal of Protective Coatings and Linings* 4(5).
- Bruce, R.R. and A. Klute. 1956. The measurement of soil moisture diffusivity. *Proceedings of the Soil Science Society of America* 20:458-62.
- Burch, D.M. 1980. Infrared audits of roof heat loss. *ASHRAE Transactions* 86(2).
- Burch, D.M. and C.M. Hunt. 1978. Retrofitting an existing residence for energy conservation—An experimental study. *Building Science Series 105*. National Institute of Standards and Technology, Gaithersburg, MD.
- Burch, D.M., W.C. Thomas, and A.H. Fanney. 1992. Water vapor permeability measurements of common building materials. *ASHRAE Transactions* 98(1).
- Burns, G.W., M.G. Scroger, G.F. Strouse, M.C. Croarkin, and W.F. Guthrie. 1992. Temperature-electromotive force reference functions and tables for the letter-designated thermocouple types based on the ITS-90. *NIST Monograph 175*. U.S. Government Printing Office, Washington, D.C.
- Carotenuto, A., F. Fucci, and G. LaFianzi. 1991. Adsorption phenomena in porous media in the presence of moist air. *International Journal of Heat and Mass Transfer* 18:71-81.
- Cohen, E.R. 1990. The expression of uncertainty in physical measurements. 1990 Measurement Science Conference Proceedings, Anaheim, CA.
- Coleman, H.W. and W.G. Steele. 1989. *Experimentation and uncertainty analysis for engineers*. John Wiley and Sons, New York.
- Coleman, H.W. and W.G. Steele. 1995. Engineering application of experimental uncertainty analysis. *ALAA Journal* 33(10).
- Considine, D.M. 1985. *Process instruments and controls handbook*, 3rd ed. McGraw-Hill, New York.
- Cunningham, M.J. and T.J. Sprott. 1984. Sorption properties of New Zealand building materials. Building Research Association of New Zealand *Research Report No. 45*, Judgeford.
- D'Eustachio, D. and R.E. Schreiner. 1952. A study of transient heat method for measuring thermal conductivity. *ASHVE Transactions* 58:331.

- DeCarlo, J.P. 1984. *Fundamentals of flow measurement*. Instrumentation Society of America, Research Triangle Park, NC.
- DeWitt, D.P. and G.D. Nutter. 1988. *Theory and practice of radiation thermometry*. John Wiley and Sons, New York.
- Fanger, P.O. 1982. *Thermal comfort*. Robert E. Krieger, Malabar, FL.
- Fanney, A.H., W.C. Thomas, D.M. Burch, and L.R. Mathena. 1991. Measurements of moisture diffusion in building materials. *ASHRAE Transactions* 97:99-113.
- Freitas, V., P. Crausse, and V. Abrantes. 1991. Moisture diffusion in thermal insulating materials. In *Insulation materials: Testing and applications*, Vol. 2. ASTM Special Technical Publication STP 1116. American Society for Testing and Materials, West Conshohocken, PA.
- Gagge, A.P., J.A.J. Stolwijk, and Y. Nishi. 1971. An effective temperature scale based on a simple model of human physiological regulatory response. *ASHRAE Transactions* 77(1).
- Goldstein, R.J. 1978. Application of aerial infrared thermography. *ASHRAE Transactions* 84(1).
- Greenspan, L. 1977. Humidity fixed points of binary saturated aqueous solutions. *Journal of Research of the National Bureau of Standards* 81A:89-95.
- Greenspan, L. and A. Wexler. 1968. An adiabatic saturation psychrometer. *Journal of Research of the National Bureau of Standards* 72C(1):33.
- Gummerson, R.J., C. Hall, W.D. Hoff, R. Hawkes, G.N. Holland, and W.S. Moore. 1979. Unsaturated water flow within porous materials observed by NMR imaging. *Nature* 281:56-57.
- Hasegawa, S. 1976. The NBS two-pressure humidity generator, Mark 2. *Journal of Research of the National Bureau of Standards* 81A:81.
- Hooper, F.C. and F.C. Lepper. 1950. Transient heat flow apparatus for the determination of thermal conductivity. *ASHVE Transactions* 56:309.
- Hooper, F.C. and S.C. Chang. 1953. Development of thermal conductivity probe. *ASHVE Transactions* 59:463.
- Hudson, R.D., Jr. 1969. *Infrared system engineering*. John Wiley & Sons, NY.
- Kumaran, M.K. and M. Bomberg. 1985. A gamma-spectrometer for determination of density distribution and moisture distribution in building materials. Proceedings of the International Symposium on Moisture and Humidity, Washington, D.C., pp. 485-90.
- Kusuda, T. 1965. Calculation of the temperature of a flat-plate wet surface under adiabatic conditions with respect to the Lewis relation. *Humidity and Moisture I:16*. Reinhold Publishing Corporation, New York.
- Liptak, B.G., ed. 1972. *Instrument engineers handbook*, Vol. 1. Chilton Book Company, Philadelphia, PA.
- Lodge, J.P., ed. 1989. *Methods of air sampling and analysis*, 3rd ed. Lewis Publishers, MI.
- Lundgren, D.A. et al., eds. 1979. *Aerosol measurement*. University Presses of Florida, Gainesville.
- Mack, R.T. 1986. Energy loss profiles: Foundation for future profit in thermal imager, sales, and service. Proceedings of the 5th Infrared Information Exchange, Book 1, AGEMA Infrared Systems, Secaucus, NJ.
- Madsen, T.L. 1976. Thermal comfort measurements. *ASHRAE Transactions* 82(1).
- Mattingly, G.E. 1984. Workshop on fundamental research issues in orifice metering. GRI Report 84/0190. Gas Research Institute, Chicago, IL.
- Mattingly, G.E. 1992. The characterization of a piston displacement-type flowmeter calibration facility and the calibration and use of pulsed output type flowmeters. *Journal of Research of the National Institute of Standards and Technology* 97(5):509.
- McCullough, E.A., B.W. Jones, and J. Huck. 1985. A comprehensive data base for estimating clothing insulation. *ASHRAE Transactions* 92:29-47.
- McLean, R.C., G.H. Galbraith, and C.H. Sanders. 1990. Moisture transmission testing of building materials and the presentation of vapour permeability values. *Building Research and Practice*. 82-103.
- Mease, N.E., W.G. Cleveland, Jr., G.E. Mattingly, J.M. Hall. 1992. Air speed calibrations at the National Institute of Standards and Technology. Proceedings of the 1992 Measurement Science Conference, Anaheim, CA.
- Miller, R.W. 1983. *Measurement engineering handbook*. McGraw-Hill, NY.
- NIST. 1976. Liquid-in-glass thermometry. NIST Monograph 150. National Institute of Standards and Technology, Gaithersburg, MD.
- NIST. 1986. Thermometer calibrations. NIST Monograph 174.
- Nottage, H.B., J.G. Slaby, and W.P. Gojsza. 1952. A smoke-filament technique for experimental research in room air distribution. *ASHVE Transactions* 58:399.
- Olesen, B.W. 1985. A new and simpler method for estimating the thermal insulation of a clothing ensemble. *ASHRAE Transactions* 92:478-92.
- Olesen, B.W., J. Rosendahl, L.N. Kalisperis, L.H. Summers, and M. Steinman. 1989. Methods for measuring and evaluating the thermal radiation in a room. *ASHRAE Transactions* 95(1).
- Paljak, I. and B. Pettersson. 1972. *Thermography of buildings*. National Swedish Institute for Materials Testing, Stockholm.
- Parmelee, G.V. and R.G. Huebscher. 1946. The shielding of thermocouples from the effects of radiation. *ASHVE Transactions* 52:183.
- Prazak, J., J. Tywoniak, F. Peterka, and T. Slonc. 1990. Description of transport of liquid in porous media—A study based on neutron radiography data. *International Journal of Heat and Mass Transfer* 33:1105-20.
- Quenard, D. and H. Sallee. 1989. A gamma-ray spectrometer for measurement of the water diffusivity of cementitious materials. Proceedings of the Materials Research Society Symposium, Vol. 137.
- Quinn, T.J. 1990. *Temperature*, 2nd ed. Academic Press, New York.
- Richards, R.F., D.M. Burch, W.C. Thomas. 1992. Water vapor sorption measurements of common building materials. *ASHRAE Transactions* 98(1).
- Richardson, L. 1965. A thermocouple recording psychrometer for measurement of relative humidity in hot, arid atmosphere. In *Humidity and Moisture I:101*. Reinhold Publishing Corporation, New York.
- Schooley, J.F. 1986. *Thermometry*. CRC Press, Boca Raton, FL.
- Schooley, J.F., ed. 1992. *Temperature: Its measurement and control in science and in industry*, Vol. 6. American Institute of Physics, New York.
- Seely, R.E. 1955. A circuit for measuring the resistance of energized A-C windings. *AIEE Transactions*, 214.
- Shafer, M.R. 1961. Performance characteristics of turbine flowmeters. Proceedings of the Winter Annual Meeting, Paper No. 61-WA-25. American Society of Mechanical Engineers, New York.
- Till, C.E. and G.E. Handegord. 1960. Proposed humidity standard. *ASHRAE Transactions* 66:288.
- Tobiasson, W. and C. Korhonen. 1985. Roofing moisture surveys: Yesterday, today, and tomorrow. Proceedings of the Second International Symposium on Roofing Technology, Gaithersburg, MD.
- Tveit, A. 1966. Measurement of moisture sorption and moisture permeability of porous materials. Report No. 45. Norwegian Building Research Institute, Oslo.
- Vernon, H.M. 1932. The globe thermometer. *Proceedings of the Institution of Heating and Ventilating Engineers* 39:100.
- Wentzel, J.D. 1961. An instrument for measurement of the humidity of air. *ASHRAE Journal* 11:67.
- Wile, D.D. 1947. Air flow measurement in the laboratory. *Refrigerating Engineering* 6:515.
- Woodring, E.D. 1969. Magnetic turbine flowmeters. *Instruments and Control Systems* 6:133.
- Worrall, R.W. 1965. Psychrometric determination of relative humidities in air with dry-bulb temperatures exceeding 212°F. In *Humidity and Moisture I:105*. Reinhold Publishing Corporation, New York.

BIBLIOGRAPHY

- Beranek, L.L. 1988. *Acoustical measurements*. Published for the Acoustical Society of America by the American Institute of Physics, New York.
- Beranek, L.L. 1989. *Noise and vibration control*. Institute of Noise Control Engineering, Poughkeepsie, NY.
- EPA. 1991. Introduction to indoor air quality: A self-paced learning module. EPA/400/3-91/002, U.S. Environmental Protection Agency, Washington.
- Harris, C.M. 1987. *Shock and vibration handbook*, 3rd ed. McGraw-Hill, New York.
- IEEE. 1987. Standard digital interface for programmable instrumentation. ANSI/IEEE Standard 488.1-87 (R 1994).
- IESNA. 1999. *Lighting handbook*, 9th ed. Illuminating Engineering Society of North America, New York.
- Lord, H.W. et al. 1987. *Noise control for engineers*. Krieger Publishing, Melbourne, FL.
- Morrison, R. 1986. *Grounding and shielding techniques in instrumentation*, 3rd ed. John Wiley and Sons, New York.
- Spitzer, D.W., ed. 1991. *Flow measurement*. Instrumentation Society of America, Research Triangle Park, NC.
- Steele, W.G., R.A. Ferguson, R.P. Taylor, and H.W. Coleman. 1994. Comparison of ANSI/ASME and ISO models for calculation of uncertainty. *ISA Transactions* 33:339-52.
- Tilford, C.R. 1992. Pressure and vacuum measurements. In *Physical methods of chemistry*, 2nd ed., Vol. 6, pp. 106-73. John Wiley and Sons, New York.