Measurement of Cardiac Output in Man with a Nonrecirculating Indicator

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A B S T R A C T The present investigation was undertaken to evaluate the utility of constant-rate injection of a nonrecirculating indicator (H₃) for the measurement of cardiac output in man. 42 patients were studied during cardiac catheterization and 8 during acute complications of arteriosclerotic heart disease, including acute myocardial infarction. Pulmonary (or systemic) arterial H₃ concentration was measured chromatographically from 2.0 ml blood samples drawn during constant-rate injection of dissolved H₃ into the systemic venous circulation (or left heart). The chromatograph was a thermal conductivity unit housed in a constant-temperature water bath to achieve an improved signal-to-noise ratio. Intrapulmonary H₃ elimination from mixed venous blood was measured directly in 14 patients and averaged 98 ±1.5% (sd). Reproducibility of output measurements was evaluated using triplicate determinations obtained over 45-60 sec in 25 consecutive patients. Coefficients of variation (sd/Mean × 100) averaged 3.4 ±2.0%, making it possible to evaluate relatively small changes in measured output with conventional statistical tests. Individual measurements could be repeated at 10-15 sec intervals. Comparisons of H₃ and direct Fick measurements were made in 44 patients; H₃ outputs averaged 106 ±4% (SEM) of Fick outputs (P > 0.1). Comparisons of H₃ and dye dilution measurements were performed in an additional 24 patients. Seven had angiographically-negligible valvular regurgitation and dye outputs averaged 106 ±3% of H₃ outputs (P > 0.1). 17 had moderate-to-severe regurgitation and dye outputs averaged 91 ±4% of H₃ outputs (P < 0.05), suggesting a small but systematic error due to undetected recirculation of dye. The H₃ technique appears advantageous for rapidly repeated determinations of output, for quantitation of small changes in output, and for situations in which recirculation of conventional indicators is a potentially significant problem.

INTRODUCTION

Conventional indicator-dilution techniques for measuring cardiac output are not optimum in patients with heart disease from at least three points of view: (a) the validity of empirical corrections required to exclude recirculating indicator is uncertain when output is low and/or valvular regurgitation is present; (b) even with repeated measurements, coefficients of variation (sd/ Mean × 100) often exceed ±10% and alterations in output of less than ±20% are difficult to appreciate; (c) practical limitations make it difficult to repeat measurements more frequently than at 1-2 min intervals. In 1958, Chidsey, Fritts, Hardewig, Richards, and Courand (1) suggested that the problem of recirculation could be minimized by employing constant-rate infusion of ⁸⁵Kr, an inert gas which is 90-95% eliminated in a single passage through the lungs. These studies were extended by Rochester, Durand, Parker, Fritts, and Harvey from the same laboratory (2) and illustrative measurements were presented at 1 min intervals during exercise. Certain disadvantages of the ⁸⁵Kr technique were identified: (a) all expired air had to be collected during, and for several minutes after the completion of each measurement to avoid contamination of the test area with radioisotope; (b) intrapulmonary elimination of ⁸⁵Kr was expected to be reduced in patients with ventilation-perfusion abnormalities; (c) recirculation was still sufficiently large to require its measurement, thereby necessitating an additional sampling site and increasing the amount of blood withdrawn for each determination to 10 ml. More recently, studies in our laboratory have validated the use of dissolved hy-

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drogen (H₂) for the constant-rate injection technique (3). Since H₂ is not radioactive and only trace amounts are employed, collection of expired air is unnecessary. Because H₂ is only one-fourth as soluble as krypton, its intrapulmonary elimination is 97–99% complete in normal situations and it is theoretically less susceptible to ventilation-perfusion abnormalities (4). Measurement of recirculation is therefore also unnecessary, as little as 2 ml of blood can be utilized for each measurement and it is practical to repeat measurements at 15-sec intervals. The original studies validating the H₂ technique were performed in experimental animals and showed agreement between the H₂ and conventional Fick and dye dilution techniques, and between the H₂ and a direct volumetric technique. The present study was intended to establish the utility of the H₂ technique in conscious man; to explore its potentially unique advantages for rapidly repeated determinations and for quantitation of small changes in output; and to obtain comparative measurements between the H₂, Fick, and dye dilution techniques, including one situation in which early recirculation is a potential problem, i.e., valvular regurgitation.

METHODS

Basic approach. The constant-rate injection H₂ technique has been described in detail previously (3). 5% dextrose-in-water (D/W)³ containing dissolved H₂ is infused at a constant-rate upstream to the heart and intravascular H₂ concentration is sampled downstream to the heart after complete mixing of incoming indicator and circulating blood. For right heart output, injection is into the right atrium (or ventricle) and sampling from the pulmonary artery. For left heart output, injection is into the left atrium (or ventricle) and sampling from a systemic artery. Using the conventional formula:

Cardiac Output (ml/min) = Rate of infusion (ml/min) × C_H₂_Samp.

where:

C_H₂_Samp. = concentration of H₂ in blood at the sampling site

C_H₂_Samp. = concentration of H₂ in blood at a point just proximal to the infusion site

All H₂ concentrations are expressed in arbitrary chromatographic units per milliliter. In view of data presented below C_H₂_Samp. is assumed to be 2% of C_H₂_St.St. During any individual infusion, the rate of infusion, H₂ concentration of the infusate and the small correction for incomplete H₂ elimination remain constant. Repeated measurements of output may therefore be obtained as rapidly as downstream blood samples are drawn.

H₂ preparation, sampling, and analysis. H₂ was passed from a standard gas cylinder through a sterile disposable filter ⁴ and bubbled through an inverted infusion bottle of 5% D/W for 15–20 min. The H₂ D/W was then poured into a sterile 200 ml syringe ⁵ and any air was carefully expelled. Infusion was accomplished with a gear-driven pump ⁶ which was shown to be accurate within ±1% of its calibration in preliminary experiments. Rates of infusion were usually 12 or 18 ml/min. The large infusion syringe allowed for repeated determinations over a prolonged period. Blood samples were collected in heparinized 2 ml glass syringes with male-Luer tips ⁷ and sealed with mercury-filled metal caps. A “blank” blood sample was obtained before each H₂ infusion, since small amounts of H₂ are sometimes present in mixed venous blood (5). Sampling during H₂ infusion was arbitrarily begun at 45–60 sec. In most patients, three to six samples were obtained during a 1–2 min period (individual samples being drawn over 10–20 sec). In some patients, sampling was less frequent but more prolonged.

The gas chromatographic system for measuring blood H₂ concentration has been refined since its original description (6). The significant changes are illustrated in Fig. 1. Dissolved gases in each blood sample are “vacuum-extracted” for 2–3 min in a volumetric Van Slyke apparatus. Carylic alcohol is used to minimize foaming but hemolyzing agents are unnecessary. The extracted gases are transferred to a 0.5 ml sampling loop attached to what is normally the waste outlet of the Van Slyke apparatus. A switching valve allows the extracted gases to be inserted into the chromatographic-carrier gas stream. The chromatograph is a thermal conductivity unit housed in a constant-temperature water bath to achieve an improved signal-to-noise ratio. Water vapor and CO₂ are removed from the extracted gases by adsorbsents located between the sampling loop and the water bath. Since O₂ is employed as the carrier gas, the large amount of O₂ extracted from each blood sample is not detected. H₂ is separated from the remaining gases (chiefly nitrogen and argon) by packed columns and detected with a sealed thermistor detector. H₂ concentration is quantitated from the height of the H₂ chromatographic peak and expressed as arbitrary chromatographic units per milliliters of blood. H₂ concentration of the injectate is also measured chromatographically, using a 2.0 ml portion sampled anaerobically from the infusion syringe. Previous studies (3) have shown that the peak height response of the chromatograph is linear within 1–3% over the range of H₂ concentrations encountered and that the chromatograph has the same sensitivity for H₂ in D/W as for H₂ in blood.

Quantitation of intrapulmonary H₂ elimination. 14 patients were studied in conjunction with routine cardiac catheterization. Pulmonary arterial- and systemic arterial-blood samples were obtained for H₂ analysis between 2 and 5 min of a constant-rate infusion of H₂ D/W into the systemic venous system. H₂ elimination was calculated by expressing the difference between pulmonary and systemic arterial H₂ concentrations as a percentage of the pulmonary arterial concentration.

Output measurements. 42 patients were studied during routine cardiac catheterization. Right and left heart H₂ measurements were employed interchangeably, the only cri-

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³ Abbreviations used in this paper: D/W, dextrose-in-water.


⁵ When necessary, storage of compressed H₂ can be avoided by use of a portable H₂ generator ("Elhygen" H₂ generator, Milton Roy Co., St. Petersburg, Fla.)

⁶ R-200-YL, Becton-Dickinson & Co., Rutherford, N. J.


⁸ Model 2YP, Becton-Dickinson & Co.

⁹ Model 425-A, Becton-Dickinson & Co.
Fig. 1 Schematic drawing of chromatographic system for $H_2$ analysis. See text for details. FC, flow controller (Model 8843 ELF, Needle Taper 1, Brooks Instrument Co., Hatfield, Pa.); the upper controller is set to deliver between 2 and 3 ml/min and the lower between 4 and 5 ml/min. AC Col. = 3 ft. column, $\frac{1}{4}$ in. O.D., packed with activated charcoal. SG Col. = 7 ft. column, $\frac{1}{4}$ in. O.D., packed with silica gel. Det., thermal conductivity detector (Model 1160, Carle Instruments Inc., Fullerton, Calif.); this is operated at 11 MA. CO$_2$ Ads., CO$_2$ adsorbent (Ascarite, Arthur H. Thomas Co., Philadelphia, Pa.), packed in $\frac{1}{4}$ in. O.D. tubing. H$_2$O Vapor Ads., water vapor adsorbent (4A molecular sieve), also packed in $\frac{1}{4}$ in. O.D. tubing. Switching Valve, model L-206-4, Loenco, Inc., Altadena, Calif. Samp. Loop, sampling loop; this is fused to the waste outlet of the Van Slyke apparatus; the fluid level is adjusted as shown before the switching valve is opened.

terion being the location in which catheters were already in place. Eight additional patients were studied during treatment for acute complications of arteriosclerotic heart disease in a Coronary Care Unit. Right heart measurements were employed in all cases, using a Swan-Ganz flow-directed catheter* to obtain pulmonary-arterial blood samples (7).

14 of the patients undergoing routine catheterization also had nearly simultaneous measurements of cardiac output by the direct Fick method. $O_2$ uptake was obtained using either closed-circuit rebreathing of pure $O_2$ (with absorption of CO$_2$) or collection and analysis of expired air. When using expired air collection, pulmonary and systemic-arterial blood samples were obtained at the midpoint of each determination and analyzed for $O_2$ content with a manometric Van Slyke apparatus. When closed-circuit rebreathing was employed, blood samples were taken just before the rebreathing was begun.

24 patients undergoing routine catheterization also had measurements of cardiac output by dye dilution techniques. For each curve, a 3.0 ml bolus containing 3.75 mg of indocyanine green dye* was injected through an atrial catheter and flushed rapidly with 6-9 ml of isotonic saline. Arterial blood was sampled with a Gilford Model 103-IR cuvette densitometer** using rates of withdrawal of 23 or 46 ml/min. The densitometer was calibrated using the patient's own blood by the method outlined by Sinclair, Sutterer, Fox, and Wood (8). Each calibration curve included a minimum of two points in addition to zero. Since the calibration became alinear at the higher dye concentrations achieved, recorded curves were replotted in terms of actual dye concentrations before semilogarithmic extrapolation. The area was then obtained in the conventional fashion. Finally, each of the 24 patients was classified as having negligible or moderate-to-severe regurgitation on the basis of left ventricular cineangiograms interpreted independently by three observers.

**RESULTS**

Quantitation of intrapulmonary $H_2$ elimination. $H_2$ elimination averaged 98 ±1.5% (SD) and was in no case less than 96%. Mean left atrial pressures ranged from 13 to 28 mm Hg and averaged 19 ±7 mm Hg, but there was no statistical correlation between $H_2$ elimination and left atrial pressure. On the basis of these and similar data in anesthetized animals (3), we have elected to assume a constant elimination of 98% in all subsequent studies.

Output measurements. Reproducibility of output measurements was evaluated using triplicate $H_2$ determinations obtained over 45-60 sec in 25 consecutive patients. Average values of output in individual patients ranged from 2.21 to 8.63 liters/min and averaged 4.18 liters/min.

![Figure 2](image)

**Figure 2** Multiple sequential measurements in three patients. Mean cardiac outputs were 7.81 ±0.52 (SD) liters/min (D. S.), 5.45 ±0.24 liters/min (E. P.) and 2.47 ±0.10 liters/min (J. W.). In patient J. W., $H_2$ samples were drawn as rapidly as possible.

*Cardiogreen, Hynson, Westcott & Dunning, Inc., Baltimore, Md.

**Gilford Instrument Labs, Oberlin, Ohio.****
Coefficients of variation (sd/mean × 100) for triplicate determinations ranged from 1.0 to 7.5% and averaged 3.4 ±2.0% (sd). Fig. 2 illustrates a similar reproducibility for multiple sequential determinations over somewhat longer periods of time.

Fig. 3 illustrates the use of H₂ measurements to evaluate relatively small changes in output at different pacing rates in Coronary Care Unit patients with complete heart block. In patient D. J., a spontaneous heart rate of 64 was accompanied by a cardiac output of 2.27 ±0.03 (SEM) liters/min. Ventricular pacing at rates of 80 and 96 increased output to 2.58 ±0.04 liters/min (P < 0.01) and 2.81 ±0.04 liters/min (P < 0.01). An additional increase in rate to 120 was associated with an output of 2.99 ±0.06 liters/min but this value did not differ significantly from that at a rate of 96. In patient M. B., a cardiac output of 2.18 ±0.06 liters/min at a spontaneous rate of 60 did not differ significantly from outputs of 2.32 ±0.02 and 2.36 ±0.06 at ventricular pacing rates of 80 and 102.

Fig. 4 illustrates the use of repeated H₂ measurements to follow trends in cardiac output during the early stages of acute myocardial infarction. In patient W. O., cardiac index was at the upper limits of normal on admission (3.62 ±0.08 liters/min per m²). During the first 24 hr in the Coronary Care Unit, it dropped to 2.38 ±0.04 liters/min per m² and then returned to a clearly normal value. Pulmonary arterial pressures was elevated during the first 24 hr but then returned to within normal limits. In contrast, patient H. C. exhibited a cardiac index which was consistently in the normal range.

Fig. 5 illustrates comparative H₂ and direct Fick measurements during cardiac catheterization. When each H₂ measurement was expressed as a percentage of the corresponding Fick measurement, H₂ outputs averaged 106 ±4% (SEM) of Fick outputs (P > 0.1, paired "t" test using absolute values for cardiac index). The correlation coefficient was 0.86 (P < 0.01).

Fig. 6 illustrates comparative H₂ and dye measurements during cardiac catheterization. Both dye and H₂ were injected into the left atrium and sampled from a systemic artery. There was no significant difference between dye and H₂ outputs when the group was considered as a whole (P > 0.1, paired "t") or when comparisons were confined to patients with indices <2.5 liters/min per m² (P > 0.1) regardless of whether dye or H₂ is used as a reference. When the total group was divided into patients with and without valvular regurgitation, results differed. In patients without regurgitation, dye indices ranged from 91 to 118% and averaged 106 ±3% (SEM) of H₂ indices (P > 0.1). In patients with regurgitation, dye indices ranged from 61 to 114% and averaged 91 ±4% of H₂ indices (P < 0.05). 10 of the regurgitation patients were also studied with right-atrial dye injection and arterial sampling, and righth-atrial dye indices ranged from 58 to 106% and averaged 88 ±4% of H₂ indices (P < 0.05).

**DISCUSSION**

The most obvious advantage of using dissolved H₂ as an indicator for measuring cardiac output is the absence of recirculation. General principles governing the intrapulmonary elimination of a dissolved gas infused intravenously have been summarized by Farhi and his col-

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**Fig. 3** Use of groups of H₂ outputs to determine effects of different pacing rates in two patients with complete heart block. Numbers in parentheses, number of H₂ determinations used to calculate each mean value. Vertical bars represent ±1 SEM.

**Fig. 4** Use of H₂ cardiac indices for following trends in acute myocardial infarction. Abbreviations, etc. as in Fig. 3. CI, cardiac index.
leagues (4, 9). The crucial steady-state parameters are the solubility of the gas and the ventilation-perfusion ratio of each alveolus. Gas elimination is directly related to ventilation-perfusion ratio and inversely related to solubility, and can be approached from two viewpoints: (a) any intrapulmonary shunt in which mixed venous blood is not exposed to alveolar air will be totally ineffective for elimination of H₂ (and all other gases); (b) since the amount of true intrapulmonary shunting is, at least in normal situations, only 0.2% (10), the alveoli of crucial importance are those with ventilation-perfusion ratios which are abnormally low but greater than zero. These alveoli are notoriously ineffective for O₂ and CO₂ exchange (4, 9) and, from that viewpoint, are “effective shunts.” On the other hand, they remain surprisingly effective for H₂ elimination. For example, an alveolus with a ventilation-perfusion ratio of 0.2 may raise mixed venous O₂ content by only half the amount required to achieve full saturation but still eliminate 93% of incoming H₂. An alveolus with a “normal” ventilation-perfusion ratio of 0.8 is only 5% more effective in eliminating incoming H₂. In the present studies, the relatively brief durations of H₂ infusion probably had the additional advantage of being too short to allow systemic arterial H₂ concentrations to rise to true steady-state levels. Because of H₂’s extremely low solubility (\(a = 0.015 \text{ ml/ml per 760 mm Hg}\)), even a nonventilated alveolus can remove significant amounts of H₂ from mixed venous blood during the first few minutes of an H₂ infusion.

In any event, the measured values for H₂ elimination in the patients in this study do not differ from those previously reported for normal experimental animals (3), despite significant elevations in left atrial pressure. On the basis of these values we have made it a practice to multiply downstream H₂ concentration by 0.98 to correct for the small amount of H₂ not normally eliminated from mixed venous blood. This correction is admitted arbitrary. Since it represents an average value, it will not be precisely correct for all individuals. In addition, since it does not take into account H₂ removal from arterial blood by body tissues during the first few minutes of an H₂ infusion, it may overestimate the amount of recirculating H₂ reaching the sampling site. In view of these considerations and the small size of the correction, some workers may prefer to omit it. It will continue to be important, however, to make direct measurements of H₂ elimination when the H₂ technique is applied to abnormal situations not previously studied.

The ease of obtaining multiple samples with the H₂ technique seems noteworthy. As mentioned above, the standard deviation of three consecutive measurements during a steady-state is 3% of the mean value. When evaluating changes in flow with single sequential samples (e.g., Fig. 2), statistical significance can be ascribed to changes of less than ±10%. In addition, the ability to draw samples at 15-sec intervals (as opposed to 1-2 min intervals) means that changes of relatively short periodicity may be identified. Using sequential groups of H₂ flows (e.g., Figs. 3 and 4), standard errors of mean values are sometimes as small as 2-3% and statistical significance may be associated with means differing by as little as ±5%. The studies in patients with acute complications of arteriosclerotic heart disease are of particular interest in this regard. Trends are easily identified even when absolute changes are small.

There is relatively little information available to make a similar evaluation for dye dilution measurements. Smulyan (11) has analyzed the variability of 67 duplicate indicator dilution studies culled from five laboratories and has stated that reproducibility (i.e., ±2 sd

**Figure 5** Comparison of measurements of cardiac index by constant-rate H₂ injection and direct Fick methods.

**Figure 6** Comparison of measurements of cardiac index by constant-rate H₂ injection and dye dilution. Each dye value represents the mean of two to four consecutive determinations. LA Dye, left atrial dye injection.
of mean difference) is ±24%. In our laboratory, 31-paired dye measurements yielded a reproducibility of the same order of magnitude, i.e., ±18%. Using three dye curves, we were able to improve reproducibility to ±12%. It is possible that sets of more than three dye curves may be used with even smaller differences. We have found it more practical to obtain multiple Hs measurements during a short (1–2 min) H2 infusion. This also allows us to take advantage of the somewhat greater reproducibility of this technique.

The Hs technique is susceptible to the same sampling errors as other indicator-dilution techniques utilizing constant-rate injection (12). However, Bassingthwaighte, Knopp, and Anderson (13) have pointed out that errors related to time-averaged sampling may be significantly less during constant-rate injection than during bolus injection. Even disregarding the possibility of sampling errors, a change in cardiac output is not instantaneously reflected in a change in downstream Hs concentration because of the effects of the mixing chamber interposed between the sites of injection and sampling. Rochester and his colleagues (2) have analyzed the change from one steady-state indicator concentration to another assuming a monoeponential washout of indicator from the right heart. Their approach suggests that 90% of any change in downstream concentration should be achieved in less than 5 sec in normal hearts and in less than 15 sec in abnormally large hearts with a low stroke volume. Since indicator washout is probably not a monoeponential process, a more precise statement is that of Zierler and his colleagues (14), who have emphasized that downstream indicator concentration becomes a valid representation of an altered output only when the last particle of indicator which entered the circulation before the alteration has passed the sampling site. As far as the present technique is concerned, there is no possibility of obtaining beat-to-beat changes. It is also unlikely that the full magnitude of respiratory-induced variations of output can be appreciated.

The present Hs and Fick measurements are felt to show reasonable agreement. Admittedly, ideal validation of a technique for measuring cardiac output requires comparison against an absolute standard. The constant-rate injection Hs technique has been validated in this fashion in experimental animals, using volumetric measurements of cardiac output during right heart by-pass (3). Similar measurements have not been feasible in man.

The comparative Hs and dye measurements require consideration from several points of view. A particularly controversial area is the adequacy of the traditional exponential extrapolation in abnormal hemodynamic states. As Zierler has pointed out (14), any treatment of recirculation is likely to be satisfactory when recirculation occurs late in the course of the primary dilution curve. Uncertainty arises as recirculation begins earlier in the curve. Oriol, Sekelj, and McGregor have clearly demonstrated failure of the extrapolation to detect coronary recirculation in experimental and clinical shock (15, 16). Maseri, Caldini, Permutt, and Zierler (17) have also presented an analogue model indicating how undetected recirculation could cause an incorrectly slow exponential decay to be assumed. The present findings are compatible with a small but systematic error due to undetected recirculation of dye in valvular regurgitation. The magnitude of the error approaches that reported by Rahimtoola and Swan (18) but is somewhat greater than that reported by Samet, Bernstein, and Castillo (19). A more definitive interpretation of the present data is complicated by the many possible reasons other than recirculation for a discrepancy between dye dilution and another technique for measuring flow. Although these factors cannot be reviewed in the present paper, several of them have been summarized in the recent studies of Saunders, Hoffmann, Noble, and Domenech (20). Our over-all feeling is that the magnitude of potential errors related to undetected recirculation and these other factors has not been fully defined, particularly in abnormal states. In our own laboratory, Hs measurements are more easily performed than dye measurements and some of these “uncertainties” are avoided.

Two potential limitations of the constant-rate injection Hs technique should be mentioned specifically. Our usual practice is to infused Hs D/W at rates of 12 or 18 ml/min. Even though the infusate does not contain saline, the volume of fluid delivered must be maintained within acceptable limits. In order to decrease the amount of fluid administration, one may utilize larger downstream samples for Hs analysis—e.g., when 6 ml samples are employed, the rate of infusion can be reduced by a factor of three. The technique also requires a sampling as well as an infusion site on the venous (or arterial) side of the lungs. The advent of the Swan-Ganz catheter has made pulmonary arterial sampling appreciably more practical in the acute care situation.

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