The Development and Use of an Innovative Laboratory Method for Measuring Arsenic in Drinking Water from Western Bangladesh

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The life expectancy in Bangladesh during the mid-1960s was only 46 years. Many premature deaths resulted from drinking surface water that was contaminated with bacteria causing diarrhea, cholera, typhoid, and other life-threatening diseases (Nyrop et al. 1975). Aid agencies, the Bangladesh government, and private individuals began installing 8–12 million tube wells to prevent these deaths by providing access to microbially safe groundwater for drinking (World Health Organization 2003). By 1995, Bangladesh had 120 million people (Monan 1995), approximately 97% of whom drank tube-well water (WHO 2000), and for a variety of reasons the life expectancy had increased to 55 years (Monan 1995).

Regrettably, this new source of drinking water was not tested for toxic metals. In 1993, Dhaka Community Hospital first diagnosed chronic arsenic poisoning caused by drinking Bangladesh’s groundwater [British Geological Survey (BGS) 1999a]. In 1997, our team produced the first national-scale map of As concentration in Bangladesh’s groundwater [Frisbie et al. 1999; USAID 1997]. This map showed that 45% of Bangladesh’s area had groundwater with As concentrations greater than the 50 µg/L national standard (Frisbie et al. 1999; USAID 1997). In 2003, a risk assessment estimated that 28 million Bangladeshis were drinking water with As concentrations greater than this national standard (Yu et al. 2003). As a result of this exposure, skin cancer, melanosis, leukomelanosis, keratosis, hyperkeratosis, and nonpitting edema from chronic As poisoning are common in Bangladesh (BGS 1999b; Frisbie et al. 2002). In addition, the rates of bladder cancer, liver cancer, and lung cancer are expected to increase in Bangladesh based on an analysis of death certificates for As exposures in Taiwan (Morales et al. 2000).

Fortunately, our 1997 study (Frisbie et al. 1999; USAID 1997) also suggested that testing and sharing tube wells could rapidly and inexpensively provide drinking water with As concentrations less than the 50 µg/L national standard to 85% of Bangladesh’s population. That is, 85% of Bangladesh’s neighborhoods have at least one tube well that does not require treatment for As removal before drinking. Therefore, the vast majority of Bangladeshis with unsafe water could potentially get safe drinking water from their neighbors (Frisbie et al. 1999; USAID 1997). As a result of this discovery, groundwater testing has become a major component of an overall strategy for providing safe drinking water to the people of Bangladesh. To date, > 1 million of Bangladesh’s approximately 10 million tube wells have been tested for As with easy-to-use, relatively inexpensive, and semiquantitative field kits [UNICEF (United Nations Children’s Fund) 2004]. Tube wells are considered safe and marked with green paint if the As concentration is ≤ 50 µg/L, the national standard. Conversely, tube wells are considered unsafe and marked with red paint if the As concentration is > 50 µg/L. Those with safe water are asked to share with their less fortunate neighbors. In addition to this initial survey, periodic testing of tube wells has been recommended to ensure that the population has continued access to safe drinking water. This periodic testing is prudent because the As concentration in some of Bangladesh’s tube wells has changed dramatically over time (Frisbie et al. 1999; USAID 1997).

This urgent need to periodically test approximately 10 million tube wells and the limited resources of Bangladesh have led to the use of these semiquantitative field kits for measuring As. However, these field kits are not precise because the user must estimate the concentration of As from a color chart, similar to that used for measuring pH with pH paper. There are two commonly used field kits in Bangladesh. The first kit is not sensitive and...
An innovative method for measuring arsenic

must be modified to detect As at the 50 µg/L national standard. The second kit is inaccurate and must be modified to avoid underestimating the true concentration of As. After modification, this second kit overestimates the true concentration of As (Geen et al. 2005). As a result of these deficiencies, there is an urgent need to supplement and ultimately replace these field kits with a quantitative, accurate, precise, sensitive, inexpensive, and environmentally safe laboratory method for measuring As in Bangladesh's drinking water. The development and evaluation of such a laboratory method are reported here for the first time.

This new method uses relatively inexpensive reagents and equipment that are easily obtained in Bangladesh and can be used to measure a wide variety of analytes. By design, it uses the same equipment as the silver diethyl dithiocarbamate \( \text{[AgSCSN(CH}_2\text{CH}_3)_2] \) method for measuring As [American Public Health Association (APHA) et al. 1989], which is commonly used in Bangladesh. Therefore, the new method can be readily implemented in Bangladesh, and it does not require expensive equipment. Such resources are rare in Bangladesh; for example, in 1997 there was only one atomic absorption spectrometer in the entire country that was used for the routine analysis of As (USAID 1997). Also, it does not require highly specialized equipment, such as devoted As analyzers, which have relatively high cost and limited utility (USAID 1997).

It is very important to realize that the WHO drinking water guideline for As of 10 µg/L is based on a \( 6 \times 10^{-4} \) excess lifetime skin cancer risk for human males, which is 60 times higher than the \( 1 \times 10^{-5} \) factor that is typically used to protect public health (WHO 1996). The WHO (1996) states that the drinking water guideline for As should be 0.17 µg/L based on the risk of death from skin cancer. However, the detection limit for most laboratories is 10 µg/L, which is why the less protective guideline was adopted: “Guideline values are not set at concentrations lower than the detection limit achievable under routine laboratory operating conditions” (WHO 1993, 1998).

Similarly, Bangladesh has limited access to atomic absorption spectrometers or other sophisticated instruments for measuring As and uses a much higher 50 µg/L drinking water standard, largely because of the poor accuracy of the AgSCSN(CH\(_2\)CH\(_3\))\(_2\) method (USAID 1997). However, a 50 µg/L drinking water standard fails to protect against death from not only skin cancer but also bladder, liver, and lung cancers. Drinking water with 50 µg/L As may cause one extra death from skin cancer per 500 people and one extra death from bladder, liver, or lung cancer per 300 people (Morales et al. 2000). Therefore, > 150,000 Bangladeshis are expected to die from skin, bladder, liver, or lung cancer caused by drinking water with > 50 µg/L As. More than 120,000 of these lives could be saved if Bangladesh complied with the more protective WHO drinking water guideline for As of 10 µg/L by sharing safe water within affected neighborhoods. The present study suggests that Bangladesh could adopt this more protective 10 µg/L guideline for As if it used the arsenomolybdate method for As measurement.

Materials and Methods

Sample Collection, Preservation, Shipment, and Analyses

Groundwater samples were collected from four neighborhoods in western Bangladesh (Fulbaria, Bualda, Jamjami, and Komlapur) during 18–21 July 2002 (Figure 1). A total of 71 random samples were collected from 67 tube wells in these four neighborhoods. A total of 18 random samples were collected from 17 tube wells in each of three neighborhoods. In the fourth neighborhood, access was denied at 1 sampling location; therefore, a

![Image](image-url)
total of 17 random samples were collected from 16 tube wells in that neighborhood.

To the extent possible, the sampled tube wells in each neighborhood were distributed at 500-m intervals along perpendicular axes that radiated in four equal lengths from the center (Figure 2). Two samples were collected from the centermost tube well in each neighborhood. One sample was collected from each of the remaining tube wells. The latitude and longitude of these tube wells were determined using a Garmin Global Positioning System 12 Channel Personal Navigator (Garmin International, Inc., Olathe, KS, USA). The accuracy of this instrument was approximately 15 m.

We used established collection, preservation, and storage methodologies to ensure that each sample was representative of groundwater quality (APHA et al. 1995; Stumm and Morgan 1981). Accordingly, all sampled tube wells were purged by pumping vigorously for 10 min immediately before sample collection. All samples were collected directly into polyethylene bottles. These samples were not filtered. Samples were analyzed immediately after collection with pH paper, preserved by acidification to pH < 2 with 5.0 M hydrochloric acid, and stored in ice-packed coolers. The temperature of all stored samples was maintained at 0–4°C until immediately before analysis at laboratories in Dubai and Vermont. In Dubai, all of these samples were analyzed for As by both the arsenomolybdate and AgSCSN(CH2CH3)2 methods. The samples were subsequently shipped to Vermont and analyzed for As by graphite furnace atomic absorption spectroscopy (GFAAS). Contour maps of the As concentration in tube-well water from these four neighborhoods were drawn (Figure 3) as described under “Mapping.”

Quality Control of Laboratory Analyses

All three methods for determining As were calibrated daily. The calibration for the AgSCSN(CH2CH3)2 method used five different standards, including a blank. The calibrations for the arsenomolybdate and GFAAS methods used six different standards, including blanks. All calibration standards were prepared from the same stock As solution. The concentration of the most dilute nonblank standard was between 1 and 10 times the method detection limit. Calibration check standards were analyzed every 20 samples and as the last analysis of each day to assess data quality. An externally supplied standard was used to assess the accuracy of the calibration standards and calibration check standards (APHA et al. 1995).

We used the recovery of a known addition of standard to eight different samples to assess the matrix effects of all three methods of determining As. In addition, three types of precision were measured for each method from the analysis of seven different standards (precision of standards), the duplicate analyses of eight different samples (precision of samples), and the analysis of a known addition of standard to eight different samples (precision of known additions). The same eight samples were used to assess the matrix effects and precision of all three methods of determining As (Table 1; APHA et al. 1995).

Arsenomolybdate Method

**Apparatus.** The apparatus, shown in Figure 4, includes an arsine (AsH3) generator (a 125 mL Erlenmeyer flask), a scrubber (made from a 10 mL volumetric pipette and a rubber stopper), and an absorber (made from a 20 mL volumetric pipette, a rubber stopper, and a polyethylene cap). The cap of the absorber has four grooves cut along its side to vent H2 gas without the loss of liquid during AsH3 generation. The spectrophotometer (product no. 6305; Jenway, Essex, UK) was set at 835 nm and used a 1.0 cm glass cell. All glassware was acid washed in 1.00 M HCl.

**Reagents.** We purchased concentrated sulfuric acid (H2SO4; product no. 102766H) and concentrated HCl (product no. 101256J) from BDH Laboratory Supplies and zinc (20-mesh granules; product no. 24,346-9) from Aldrich Chemical Company (Milwaukee, WI, USA).

**As solutions.** We prepared the As solution by dissolving 4.0 g sodium hydroxide (product no. 131687; Panreac Química S.A., Barcelona, Spain) in 10 mL distilled water; arsenic trioxide (1.320 g; product no. 22,762-5; Aldrich) was added, and the solution was diluted to 1,000 mL with distilled water. An intermediate As solution was prepared by diluting 5.00 mL stock As solution to 500 mL with distilled water. This intermediate solution was then used to prepare a standard As solution (10.00 mL intermediate As solution diluted to 100 mL distilled water, which was used in the experiments.)
Potassium iodide (50.0% wt/vol). Fifty grams of KI (product no. 102123B; BDH Laboratory Supplies, Poole, UK) was diluted to 100 mL with distilled water and stored in the dark until use.

Stannous chloride dihydrate (40.0% wt/vol). Forty grams of stannous chloride dihydrate (SnCl2·2H2O; product no. 102704Q; BDH Laboratory Supplies) was dissolved and diluted to 100 mL with concentrated HCl.

Lead (II) acetate trihydrate (10.0% wt/vol). Pb(COOCH3)2·3H2O (10.0 g; product no. 21,590-2; Aldrich) was dissolved and diluted to 100 mL with distilled water.

Sodium metabisulfite (6.00% wt/vol). 6.9 g (NH4)6Mo7O24·4H2O (product no. 102474V; BDH Laboratory Supplies) was dissolved and diluted to 100 mL with distilled water; the solution was prepared fresh daily.

Sulfuric acid/ammonium molybdate tetrahydrate. We prepared 100 mL of 6.50 M H2SO4 in distilled water. We then added 6.50 M H2SO4 and the 6.9% (wt/vol) SnCl2·2H2O, 5.0 g Zn was added. Each arsenomolybdate sample was prepared by first adding 2.50 mL I2/KI solution to a 20 mL test tube; 0.50 mL 1.00 M NaHCO3 was then added. This solution was poured into the absorber and the cap was placed on the absorber (Figure 4).

Absorber preparation. The absorber was prepared by placing 0.17 ± 0.03 g glass wool into a 20 mL test tube and mixed. The Na2S2O5 solution was added to the test tube and mixed. The Na2S2O5 was mixed with the treated sample or treated standard in the 125-mL Erlenmeyer flask (the AsH3 generator); after adding and mixing 10.0 mL concentrated HCl and 1.0 mL 40.0% (wt/vol) SnCl2·2H2O, 5.0 g Zn was added. The scrubber and absorber were then connected to the AsH3 generator (Figure 4). We allowed 30 min for the AsH3 to completely evolve from the AsH3 generator to the absorber.

The liquid from the absorber was poured into a test tube calibrated to receive 5.00 mL. 0.50 mL of distilled water was then used to rinse the residual liquid from the absorber to the test tube. One milliliter of H2SO4/ (NH4)6Mo7O24·4H2O solution was added to the test tube and mixed. Then 0.50 mL of 6.00% (wt/vol) Na2S2O3 solution was added to the test tube and mixed. The Na2S2O3 changed the mixture from deep reddish brown to faint yellow (the brown color must be eliminated). When necessary, distilled water was added to until the total volume of liquid was 5.00 mL. Then 0.50 mL 0.20% (wt/vol) SnCl2·2H2O was added and mixed. After waiting 30 min for the bluish green arsenomolybdate color to develop, we measured the absorbance at 835 nm.

Calibration. For calibration, we added 0, 0.50, 1.00, 2.00, 4.00, and 8.00 mL of standard As solution into six separate 125-mL Erlenmeyer flasks and added distilled water until the total volume of liquid in each flask equaled 35.0 mL (Figure 4). The resulting standards contained 0, 0.50, 1.00, 2.00, 4.00, and 8.00 µg of As or 0, 14, 28.6, 57.1, 114, and 229 µg/L, respectively. We used the sample treatment, absorber preparation, absorber preparation, AsH3 generation, color development, and spectrophotometry procedures described above to analyze these standards. We confirmed that the calibration results follow Beer’s law; that is, we tested for a higher order polynomial relationship to confirm linearity, and we tested the null hypothesis that the y-intercept goes through the origin (Neter et al. 1985).

Ultraviolet/Visible Spectrum of Arsenomolybdate

In order to determine the absorption maximum of arsenomolybdate we measured the absorption spectra of three arsenomolybdate samples using an Agilent 8453 ultraviolet/visible spectroscopy system (Agilent Technologies, Palo Alto, CA, USA). The wavelengths of these spectra ranged from 190 to 1,100 nm. Each arsenomolybdate sample was prepared from a 229 µg/L As standard solution using the sample treatment, absorber preparation, absorber preparation, AsH3 generation, color development, and spectrophotometry procedures described above under “Arsenomolybdate Method.” The spectrum of each arsenomolybdate sample was measured relative to the 229 µg/L As standard solution using the same UV/vis spectrometer.

Table 1. Quality control results, equipment costs, and solvents for the determination of As by arsenomolybdate, AgSCSN(CH2CH3)2, and GFAAS methods.

| Solvent                  | Arsenomolybdate | AgSCSN(CH2CH3)2 | GFAAS |
|-------------------------|-----------------|-----------------|-------|
| Method detection limit  | 7 µg/L          | 9 µg/L          | 0.7 µg/L |
| Equipment cost          | $6,700          | $6,700          | $37,000    |
| Recovery of known additions | 101.5 ± 3.6% | 103 ± 1.8%     | 103.3 ± 3.1% |
| Precision of standards  | 2.1 µg/L         | 2.5 µg/L        | 0.22 µg/L |
| Precision of samples    | 4.9 µg/L         | 4.5 µg/L        | 1.7 µg/L |
| Precision of known additions | 7.4 µg/L | 24 µg/L         | 2.0 µg/L |
| Solvent                  | H2O             | CH3OH or C2H5OH | H2O     |

*Includes a spectrophotometer, distillation unit for purifying laboratory water, analytical balance, top-loading balance, hot plate with stirrer, and glassware. *Includes an atomic absorption spectrometer, distillation unit for purifying laboratory water, analytical balance, top-loading balance, and glassware. *95% confidence interval.
to a 0 µg/L As standard solution using these procedures. Each spectrum was measured after 30 min of color development.

**AgSCSN(CH2CH3)2 Method**

We analyzed all samples for As following the AgSCSN(CH2CH3)2 method of APHA et al. (1989). In this method, a 125-mL specimen jar was used for the AsH3 generator. The scrubber and absorber were identical to those used for the “Arsenomolybdate Method” shown in Figure 4. A 35.0 mL aliquot of sample or standard was placed in the AsH3 generator and treated with 5.0 mL concentrated HCl, 2.00 mL 15.0% (wt/vol) KI in distilled H2O, and 0.40 mL 40.0% (wt/vol) SnCl2·2H2O in concentrated HCl to reduce As(V) to As(III). This reduction was allowed 15 min for completion at room temperature. The scrubber was prepared as described above for the “Arsenomolybdate Method.” The absorber received 4.00 mL 0.50% (wt/vol) AgSCSN(CH2CH3)2 in pyridine (C3H6N). Then, 3.0 g Zn was added to the sample or standard to standardize the AsH3. After 30 min of AsH3 generation, the absorbate was measured spectrophotometrically at 535 nm.

**GFAAS Method**

All samples were analyzed for As by GFAAS with a Buck Scientific 220AS autosampler, 220GF graphite furnace, and 210VGP atomic absorption spectrometer (Buck Scientific, East Norwalk, CT, USA). A 1.00 mL aliquot of standard As solution, sample, or diluted sample was loaded onto the autosampler. The six standards contained 0, 1.0, 5.0, 15.0, 30.0, and 50.0 µg/L As, respectively. The matrix of each 1.00 mL aliquot was modified with 50.0 µL 10.0% (wt/vol) ammonium nitrate (NH4NO3; product no. A1216; Spectrum Chemicals and Laboratory Products, New Brunswick, NJ, USA) in deionized water. 50.0 µL 0.2% (wt/vol) palladium nitrate [Pd(NO3)2] in 2% (wt/vol) HNO3 (product no. K, Buck Scientific), and 50.0 µL 1.79% (wt/vol) magnesium nitrate hexahydrate [Mg(NO3)2·6H2O; product no. 5855-1; EM Science, Gibbstown, NJ, USA] in deionized water. The autosampler delivered a 20.0 µL aliquot of this mixture to the graphite furnace. The furnace tube was made from nonplutographic graphite (product no. BS300-1253; Buck Scientific). The furnace initialized at 100°C for 10 sec, heated to 250°C for 20 sec, drier the mixture at 250°C for 15 sec, heated to 750°C for 25 sec, ashed the mixture at 750°C for 10 sec, heated to 2,200°C for 1.5 sec, and atomized the mixture at 2,200°C for 3 sec. The sheath and internal flows of argon gas were 1,200 and 200 mL/min, respectively. The absorbance from a hollow-cathode lamp was read at 193.7 nm through a 0.7 nm slit and after deuterium (D2) background correction. This absorbance was measured for 2.4 sec during atomization. Finally, this absorbance over time was used to calculate As concentration (Buck Scientific Inc. 2002; Harris 1999).

**Statistics**

We measured all 71 samples from this study for As by the arsenomolybdate, AgSCSN(CH2CH3)2, and GFAAS methods. We used a paired t-test of the As concentrations from these samples to determine if the arsenomolybdate and AgSCSN(CH2CH3)2 methods gave equivalent or different results (Table 2); a second paired t-test to determine if the arsenomolybdate and GFAAS methods gave equivalent or different results (Table 3); and a third paired t-test to determine if the AgSCSN(CH2CH3)2 and GFAAS methods gave equivalent or different results (Table 4). Each of these three paired t-tests was evaluated at the 95% confidence level (Snedecor and Cochran 1982).

We used an F-test to determine if two precision values were equivalent or different. The standard deviation indicates precision (Table 1; APHA et al. 1995), and a variance is indicated by precision squared (Snedecor and Cochran 1982). A ratio of these variances in an F-test show the equality of the corresponding precisions (Snedecor and Cochran 1982). Each F-test was evaluated at the 95% confidence level.

**Mapping**

We used the As concentration by the arsenomolybdate method, the sample location by the Global Positioning System, and the Surfer Surface Mapping System (version 7; Golden Software Inc., Golden, CO, USA) to draw one contour map for each of the four neighborhoods (Figure 3). To map the contours shown in Figure 3, we used a variogram to select the equation that best matched the spatial continuity of the actual As concentrations in each neighborhood (Figure 2). We used inverse-distance weighted least-squares equations (Shepard’s method) for Fulbaria and Buulda and a logarithmic equation for Jamjami. We did not use an equation for Komlapur because all the samples in this neighborhood had As concentrations ≤ 10 µg/L, the WHO drinking water guideline (WHO 1996).

**Societal Evaluation**

**Initial interview during sampling.** One principal user of each tube well was interviewed during groundwater sampling during 18–21 July 2002. Each interview was conducted in Bangla from a list of standard questions. Each interviewee was asked whether alternative drinking water sources were available (rain, ponds, rivers, or canals), and the following information was recorded: a) the number of users per tube well; b) the depth and age of each tube well; c) the results from previous As tests, if any; and d) whether any users were known to have melanosis, keratois, gangrene, skin cancer, conjunctivitis, respiratory distress, or enlarged liver. Finally, the willingness of each interviewee to get safe drinking water from their neighbors or to give safe drinking water to their neighbors was evaluated.

**Distributing As results.** Our field staff gave a Bangla-language form letter summarizing the neighborhood’s As results to each interviewee 6 months after their groundwater was sampled. In addition, the contents of each letter were explained in Bangla by our field staff. Therefore, each interviewee knew if his tube well had a safe or unsafe concentration of As. Furthermore, each interviewee knew which neighbor’s tube well had a safe or unsafe concentration of As.

Interviewees with As concentrations ≤ 10 µg/L were informed that their tube wells were safe with respect to this element. In addition,
they were informed that their tube wells should be tested for As at least once a year because the concentration of As can change over time (Frisbie et al. 1999; USAID 1997). Finally, these interviewees were asked to share their drinking water with those who could not get safe water from their own tube wells.

Interviewees with As concentrations > 10 µg/L (WHO drinking water guideline) were informed that their tube wells were unsafe with respect to this element, and the imminent risk of serious health problems from continuing to drink this water was thoroughly explained. Finally, they were asked to get safe drinking water from their neighbors.

**Final interview 1 year after sampling.** Each original interviewee, if available, was revisited 1 year after their groundwater was sampled and 6 months after they were given the As results for their neighborhood. If the original interviewee was not available, then a surrogate interviewee was identified. These people were interviewed in Bangla from a list of standard questions. The purposes of this final interview were to determine (a) if the people with safe tube wells actually gave drinking water to their neighbors who did not have safe drinking water; (b) whether the people with unsafe tube wells actually got safe drinking water from their neighbors; (c) whether the people with safe water followed our instructions and retested their tube wells for As; and (d) the health status of all tube-well users.

**Results and Discussion**

**Rationale for sampling drinking water in western Bangladesh.** We chose western Bangladesh for this study because it has some of the widest ranges of groundwater As concentrations in the country, according to our two national-scale surveys (Frisbie et al. 1999, 2002; USAID 1997). This variability is associated with the complex mixture of flood plain deposits that form western Bangladesh (Yu et al. 2003). As a result, the neighborhoods in western Bangladesh usually have at least one tube well that does not require treatment for As removal before drinking (Frisbie et al. 1999; USAID 1997). For example, the As concentrations in our four randomly selected western Bangladeshi neighborhoods ranged from < 0.7 to 590 µg/L, with 30% of samples > 10 µg/L, the WHO drinking water guideline (WHO 1996). This makes western Bangladesh well suited for evaluating our laboratory method of measuring As in drinking water. It is also well suited for determining whether people will use the results from this method to share safe drinking water with their neighbors.

**Independent evaluation of the arsenomolybdate method.** The arsenomolybdate calibration obeys Beer’s law; that is, the plot of absorbance versus As concentration is linear, and the y-intercept goes through the origin. A test for a higher order polynomial relationship was done for each daily calibration and routinely confirmed this linearity at the 95% confidence level (Neter et al. 1985). Similarly, a test of the null hypothesis that the y-intercept goes through the origin was performed for each daily calibration and routinely confirmed that the y-intercept was equivalent to 0, at the 95% confidence level (Neter et al. 1985). The calibration equation from the arsenomolybdate method detection limit study is absorbance = 0.00169 × (micrograms As per liter), and the correlation coefficient (r) = 1.00. The slope of the equation and the 7 µg/L method detection limit shown in Table 1 suggest that the colorimetric measurement of As by arsenomolybdate is extremely sensitive. The 101.5 ± 3.6% recovery of known additions shown in Table 1 suggests that the results from the arsenomolybdate method are accurate. The respective 2.1, 4.7, and 7.1 µg/L precisions of standards, samples, and known additions shown in Table 1 suggest that the results from the arsenomolybdate method are reproducible.

In this method, As is removed from the sample by reduction to AsH3 gas. This gas is collected in an absorber by oxidation to arsenic acid (H3AsO4). All previous attempts at this separation have suffered from poor reproducibility because of the incomplete recovery of AsH3 (Sandell 1942, 1959). The sources of this incomplete recovery were insufficient concentrations of oxidant in the absorbate, the deterioration of absorbate with time, and improper absorber designs. This problem of incomplete recovery has been solved.

We discovered and corrected an error that has not been reported by previous researchers. This error was from drift caused by the arsenomolybdate absorption spectrum changing with time. Our method requires that the arsenomolybdate color develop for 30 min before its absorbance is measured. This color is relatively stable after 30 min. In addition, a loss of sensitivity from using polychromatic light to measure absorbance has been corrected. Our method requires that the absorbance be measured at 835 nm, the absorption maximum of arsenomolybdate. In addition, previous arsenomolybdate methods were not compared with established methods for measuring As (Milton and Duffield 1942; Sandell 1959). Comparisons of our method with the AgSCSN(CH2CH3)2 and GFAAS methods for measuring As are shown in Tables 1–4.

Finally, all routine analytical methods use a rigorous quality control plan to implement these methods are comparable; the precisions of standards for the methods are equivalent at the 95% confidence level; and the precision of samples for the two methods are equivalent at the 95% confidence level (Table 1).

However, the precision values of known additions for the arsenomolybdate and AgSCSN(CH2CH3)2 methods are different at the 95% confidence level (Table 1). These two precision values were measured using the same eight samples. This suggests that the AgSCSN(CH2CH3)2 method is imprecise at relatively high As concentrations because of the sample matrix. Additional evidence of this imprecision is the 16.6% relative percent difference of the mean As concentrations from all 71 samples measured by the arsenomolybdate and AgSCSN(CH2CH3)2 methods (Table 2). One source of this imprecision may be the incomplete reduction of As(V) to As(III) in Bangladesh’s tube-well water by the relatively mild sample treatment procedure of the AgSCSN(CH2CH3)2 method. More specifically, the arsenomolybdate method uses KI and SnCl2·2H2O at 100°C for this reduction, whereas the AgSCSN(CH2CH3)2 method uses KI and SnCl2·2H2O at room temperature. Another source of this imprecision may be the incomplete generation of AsH3 in Bangladesh’s tube-well water by the relatively mild AsH3 generation procedure of the AgSCSN(CH2CH3)2 method. More specifically, the arsenomolybdate method uses 0.18 g KI, 0.540 g SnCl2·2H2O, 2.0 mL concentrated H2SO4, 10.0 mL concentrated HCl, and 5.0 g Zn per 35.0 mL of sample for AsH3 generation. In contrast, the AgSCSN(CH2CH3)2 method uses 0.300 g KI, 0.16 g SnCl2·2H2O, 5.0 mL concentrated HCl, and 3.0 g Zn per 35.0 mL of sample for AsH3 generation.

Another important difference between the arsenomolybdate and AgSCSN(CH2CH3)2 methods is related to worker health and environmental protection. More specifically, the arsenomolybdate method uses H2O as a solvent, which is nontoxic, nonflammable, and easy to dispose of. In contrast, the AgSCSN(CH2CH3)2 method uses either chloroform (CHCl3) or C2H5N as a solvent.
In summary, the arsenomolybdate method is more accurate and affordable than the GFAAS method (Tables 1 and 3).

**AgSCSN(CH₂CH₃)₂ versus GFAAS.** The AgSCSN(CH₂CH₃)₂ and GFAAS methods for measuring As are different at the 95% confidence level, according to a paired *t*-test of all 71 samples (*p*-value = 0.24; Table 3). As a result, these two methods are highly correlated (*r* = 0.996; Table 3). Both methods can quantify As to less than the 10 µg/L WHO drinking water guideline (WHO 1996; Table 1). However, the equipment cost for the GFAAS method is much greater than that for the arsenomolybdate method (Table 1).

Despite its far lower cost, the arsenomolybdate method is more accurate than the GFAAS method. The recovery of known additions by the arsenomolybdate method is equivalent to 100% (101.5 ± 3.6%; Table 1). In contrast, the recovery of known additions by the GFAAS method is > 100% (103.3 ± 3.6%; Table 1). This suggests that the GFAAS method overestimates the true concentration of As in this matrix by approximately 3.3% (Table 1). This estimate of 3.3% bias by the GFAAS method is based on the recovery of known additions from eight samples. Additional evidence of this bias is the 3.6% relative percent difference of the mean As concentrations from all 71 samples measured by the arsenomolybdate and GFAAS methods (Table 3). This overestimation by the GFAAS method was likely caused by scattered light from sodium chloride or similar matrix salts that remained in the furnace during atomization (Frisbie et al. 1999, 2002; Harris 1999; USAID 1997). If so, the NH₄NO₃, Pd(NO₃)₂, and Mg(NO₃)₂·6H₂O matrix modifiers and D₂ background correction did not fully resolve this interference.

In contrast, the GFAAS method is more precise than the arsenomolybdate method. The standards, samples, and known additions are measured with greater precision by the GFAAS method than by the arsenomolybdate method (Table 1). Each of these three *t*-tests was evaluated at the 95% confidence level.
Education programs concerning the dangers of As-contaminated water have achieved some measure of success, because only one owner of an unsafe tube well in our study appeared not to have understood the implications of his tube well’s As results. This owner, a 70-year-old male, professed during the follow-up interview that his water was safe, and he told us that he had continued to use it and share it freely with others.

Many of the tube-well owners (85%) followed our recommendation and had their tube wells retested. This retesting was not performed by our research team. Only one respondent told us that he thought retesting was not necessary (the one who believed his water was safe when it was not); the others who did not retest their water gave cost as the reason for not retesting. People were especially willing to retest their water if this could be done at no cost to them. Of the respondents who had their water retested, 96% reported that the retesting was done with no charge to them; however, two respondents paid to have their water retested.

Our survey suggests that the number of tube wells, especially private tube wells, is rapidly increasing in Bangladesh as more families acquire the economic resources to build them. Only 14% of tube wells were reported to have been constructed before 1983 (56% private); 27% were reported to have been constructed from 1983 to 1992 (83% private), and 59% from 1993 to 2002 (92% private), making the tube wells in our random sample from western Bangladesh only 9 years old on average. In addition, we asked whether respondents were aware of any As patients in the area. The number of nearby As patients was not statistically related to the concentration of As during our sampling event (p = 0.44). Similarly, the concentration of As during our sampling event was not statistically related to the age of the tube well (p = 0.51). In contrast, the number of nearby As patients was statistically related to the age of the tube well and hence the duration of exposure to tube-well water that was potentially contaminated with As. That is, the oldest tube wells were associated with the highest number of nearby As patients (p = 0.03). This stresses the fact that duration of exposure to As must be considered in addition to the concentration of As at any given time, because this can explain why the tube wells of some As patients are found to contain very low levels of As but have been used over many years (WHO 2001). For example, Figure 5 shows a female As patient with keratosis of the palms and blackfoot disease. When tested in 2002, the tube well used by this patient had an As concentration of 1.4 µg/L, but she had been drinking from this tube well for 34 years. The As concentration in some of Bangladesh’s tube wells has changed dramatically over time (Frisbie et al. 1999; USAID 1997). These results demonstrate the need to periodically test all drinking water tube wells so that the total lifetime exposure to As can be reduced. Because most of the tube wells in the country have been installed quite recently, the numbers of As patients may begin to increase dramatically as more people develop a history of using tube wells for longer than the 5–10 years it may take to develop symptoms of chronic As poisoning. Finally, as the ages of the tube wells and the length of exposure to As increase, it may become even more vital to adopt a drinking-water standard for As lower than the current Bangladesh standard of 50 µg/L, such as the 10 µg/L WHO guideline (WHO 1996).

Conclusions
A valid method for the determination of As by arsenomolybdate is now available. This method is more accurate, precise, and environmentally safer than the AgSCSN(CH₂CH₃)₂ method (Tables 1 and 2), and it is more accurate and affordable than the GFAAS method (Tables 1 and 3).

Most important, the arsenomolybdate method is the only accurate, precise, and safe way to quantify As to less than the 10 µg/L WHO drinking water guideline (WHO 1996) without expensive or highly specialized laboratory equipment (Table 1). This suggests that developing countries such as Argentina, Bangladesh, Bolivia, Chile, China, India, Mexico, Nepal, Pakistan, Peru, and Thailand, with limited access per capita to atomic absorption spectrometers or other sophisticated instruments for measuring As, could lower their 50 µg/L drinking water standards to the more protective 10 µg/L guideline if they use the arsenomolybdate method (Bhattacharya et al. 2002; Ng et al. 2003; United Nations 2004). More than 32 million people from these countries drink water with As concentrations greater than their national standards of 50 µg/L (Bhattacharya et al. 2002; Ng et al. 2003; United Nations 2004; Yu et al. 2003). This suggests that > 170,000 people from these countries will die from skin, bladder, liver, or lung cancer caused by drinking water with > 50 µg/L As. If these countries complied with a 10 µg/L drinking water standard for As, perhaps by sharing safe water as was done in the four neighborhoods from this study, > 140,000 of these 170,000 lives could potentially be saved.

In particular, it is very important that the Government of Bangladesh lower its 50 µg/L drinking water standard for As. The rapidly increasing number of tube wells in Bangladesh suggests that the mortality rate from chronic As poisoning is also rapidly increasing. Lowering this standard will reduce exposure and save lives by encouraging the use of safer drinking water.

Finally, our surveys suggest that Bangladeshis will readily test and share their drinking water to meet the more protective WHO guideline for As of 10 µg/L. More specifically, 85% of tube-well owners were concerned enough to retest their water for As within 1 year, and 90% of the tube-well owners that had tube-well As concentrations > 10 µg/L actually gathered water from their neighbors.

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Figure 5. Photograph of a Bangladeshi female with keratosis of the palms and blackfoot disease. Her well contained 1.4 µg/L As on 21 July 2002; she had been drinking from this well for 34 years.
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