TECHNICAL NOTE

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Preparation of Carboxyhemoglobin Standards and Calculation of Spectrophotometric Quantitation Constants*

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ABSTRACT: A method was developed for the preparation of carboxyhemoglobin (COHB) standards, which were stable for more than four months with the prepared control remaining within acceptable limits during this time. A mathematical equation was developed to more accurately determine the constants A and B used in the equation COHB% = 100[(C - B)/(A - B)], where B = 0%COHB peak ratio at 540 nm and 579 nm; A = 100% COHB peak ratio at 540 nm and 579 nm; and C = the peak ratio at 540 nm and 579 nm for the blood being analyzed. The following equations were developed to calculate A and B: $B = P_{avg} - (P) [(P_{avg} - N_{avg})/(P - N)]; A = B + (P_{avg} - N_{avg})/(P - N), P_{avg} = average$ peak ratio 540/579 for the positive standard run on the spectrophotometer; P = average decimal concentration measured on the CO-OXIMETER for the positive standard; N_{avg} = average peak ratio 540/579 for the negative standard; N = average decimal concentration measured on the CO-OXIMETER for the negative standard. The new equations provided results consistent with those obtained from a CO-OXIMETER.

KEYWORDS: forensic science, forensic toxicology, carboxyhemoglobin, analysis, standards

Determination of the *A* and *B* ultraviolet/visual spectrophotometry (UV/VIS) constants in the calculation of carboxyhemoglobin (COHB) concentration is complicated by the fact that the *A* and *B* constants are based on an assumed 100% carbon monoxide saturated solution of hemoglobin and a 0% carboxyhemoglobin solution. Research has shown that this assumption is not always true and could be problematic when attempting to interpret results (1). In late 1995 commercially available whole blood carboxyhemoglobin standards, used to calibrate a UV spectrophotometer and to serve as controls, were discontinued. This required that a stable and reliable standard and calibration control be developed for carboxyhemoglobin analysis in postmortem whole blood by UV/VIS spectrophotometry. The samples received for analysis by the laboratory do not lend themselves to analysis by CO-OXIMETER due to the presence of clots and occasionally to interfering absorbance (Fig. 1). Procedures found in the literature (1-7) for the preparation of controls were difficult to use and the assumed concentrations could not easily be confirmed, leading to questionable instrument calibration when compared to commercially available controls.

Methods

In this experiment, fresh whole blood (non-smoker) was drawn into four 10 mL gray top vacutainer tubes (VENOJECT, Terumo Medical, Elkton, MD) containing 100 mg sodium fluoride and 20 mg potassium oxalate on the day the carboxyhemoglobin standards were to be prepared. The gray top tubes are used by forensic toxicology laboratories in actual cases to collect postmortem blood. The collection of whole blood in gray top tubes adequately simulates the biological matrix received by the laboratory for analysis. The gray top tubes are used by most forensic toxicology laboratories to prevent clotting and preserve the specimen. Attempts were made to use blood obtained from a local blood bank; however, the blood contained fibrils which resulted in problems with reproducibility of carboxyhemoglobin concentration on the CO-OXI-METER 282 manufactured by Instrumentation Laboratory, Lexington, MA. The fresh whole blood collected should be treated immediately after collection. The blood collected was placed on a tube rocker and rocked for 30 min after collection to make sure the samples were well mixed with the anticoagulant and the antibacterial agents. All of the samples were then placed into a 25 by 150 mm tube and mixed for an additional 30 min.

A "NEGATIVE CO" standard was prepared by placing the fresh blood sample into a 200 mL volumetric flask. The flask was placed in a horizontal position and rotated, while < 1.0 mL/min nitrogen flow was used to purge the flask for 2 to 3 min. A portion of this negative control was pipetted into small plastic 1.5 mL standard micro centrifuge tubes made by Elkay Products, Inc. Each tube was filled to capacity and sealed with an attached plastic cap. These "NEGATIVE CO" standards were stored overnight at 4°C and tested the next day on a CO-OXIMETER to obtain the average concentration for the "NEGATIVE CO" control.

The "POSITIVE CO" standard was prepared under a biological ventilated hood by placing the remaining "Negative CO" blood into a 200 mL volumetric flask. The flask was placed in a horizontal position and rotated, while < 1.0 mL/min carbon monoxide flow was used to purge the flask for 20 min. This approximately 100%

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FIG. 1-Interfering UV spectra.

carboxyhemoglobin standard was diluted to approximately 45% COHB with a portion of the negative control prepared earlier. A portion of this approximately 45% positive control was pipetted into small plastic 1.5 mL standard micro centrifuge tubes. Each cup was filled to capacity and sealed with an attached plastic cap. These "POSITIVE CO" standards were stored overnight at 4°C and tested the next day on a CO-OXIMETER to obtain the average concentration for the "POSITIVE CO" control. It is important to allow the specimen to equilibrate overnight in order to reach complete equilibrium between bond and unbound CO. Kinetic experiments made during this research indicate that it takes several hours after the preparation of the standard for the CO saturated blood to reach equilibrium.

In the preparation of the test reagent, bottles of ammonium hydroxide and sodium dithionite were allowed to equilibrate at room temperature for more than 15 min. In a 1000 mL volumetric flask, 500 mL of deionized water was added along with 5 g of sodium dithionite. The solution was mixed and allowed to completely dissolve before filling the flask to the 1000 mL mark with deionized water. The solution was then placed under a ventilated hood and 4 mL of concentrated ammonium hydroxide was pipetted into the solution. The solution was then poured into a brown bottle pipettor and permitted to equilibrate at room temperature for more than 15 min. The solution has a shelf life of no more than 4 h.

Each NEGATIVE and POSITIVE standard was run on the CO-OXIMETER ten times to obtain an average carboxyhemoglobin level and a standard deviation. These levels were then used to calibrate the HP 8453 UV Visible Spectrophotometer, equipped with an HP 89052B Sipper System, an HP 89072A Autosampler, and controlled by the HP UV-Visible Chemstation Revision A.02.05, which were supplied by Hewlett-Packard, Palo Alto, CA. This method uses the equation COHB% = 100*[C - B)/(A - B)] and the isosbestic points at 540 nm and 579 nm to calculate the concentration of carboxyhemoglobin in the blood. The following equations were developed by the laboratory to calculate the constant values of A and B:

$$B = P_{avg} - (P)[(P_{avg} - N_{avg})/(P - N]]$$
$$A = B + (P_{avg} - N_{avg})/(P - N)$$

where P_{avg} is the average peak ratio 540/579 for the positive standard run on the spectrophotometer, *P* is the average decimal concentration measured on the CO-OXIMETER for the positive standard, N_{avg} is the average peak ratio 540/579 for the negative standard, and *N* is the average decimal concentration measured on the CO-OXIMETER for the negative standard.

Once the instrument was calibrated, the whole blood samples received for analysis were prepared in duplicate. For each specimen, there were two samples prepared and placed into two 16 by 100 mm tubes. Each tube contained 10 mL of the prepared reagent (sodium dithionite/ammonium hydroxide) along with 100 μ L of the unknown blood sample (clot free). Parafilm was placed over the top of each tube and properly mixed. NEGATIVE and POSITIVE standards were prepared in the same way as samples received for analysis.

The samples were then placed into the Hewlett Packard 89072A Autosampler on the Hewlett Packard 8453 Diode Array Spectrophotometer for analysis in the following order: Reference Blank, Positive, Reference Blank, Negative, Reference Blank, Unknown (1a), Reference Blank, Unknown (1b), Reference Blank.

The instrument was started and data were collected on samples. Absorbance was obtained at 540 nm and 579 nm for all specimens (Fig. 2) and used to calculate the % COHB.

A comparison of the prior method for calculating the *A* and *B* constants used in the quantitation of CO were made by taking a portion of the "NEGATIVE CO" standard and placing it into a 200 mL volumetric flask. The flask was placed in a horizontal position and rotated for 20 min, while < 1.0 mL/min flow of carbon monoxide was pumped into the flask to prepare a theoretical 100% carboxyhemoglobin. Nitrogen was used to purge the flask for 2 to 3 min to remove unbound CO. The 0% carboxyhemoglobin was prepared by using oxygen instead of CO in the method provided for preparing the 100% carboxyhemoglobin standard. This blood was used to determine the *A* and *B* constants and the results from these values were compared with values from the new way of calculating *A* and *B*.

Results

The visible spectra for the positive control can be seen in Fig. 2 and the negative control in Fig. 3. In comparing the two instru-



TABLE 1—Statistical comparison of CO-OXIMETER and spectrophotometer and stability of COHB control for 5 months.

Description	Ν	Mean	Variance	SD	CV
Positive control on CO-OXIMETER	10	45.6	0.8	0.9	2.0
Positive control on spectrophotometer	10	45.8	0.1	0.3	0.7
Negative control on CO-OXIMETER	10	1.2	0.1	0.3	24.8
Negative control on spectrophotometer	10	1.3	0.0	0.2	16.5
Positive 49.2% control run for 5 months	44	48.1	19.7	4.4	9.2
Negative 1.6% control run for 5 months	29	1.5	3.3	1.8	125.8

ments used in the experiment, the CO-OXIMETER (Table 1), and UV spectrophotometer (Table 1, UV/VIS), clearly show that the calibrations were close in value for both the CO-OXIMETER and the UV spectrophotometer with standard deviations of 0.9 and 0.3, respectively. Table 1 contains the results of a "negative control" run on the CO-OXIMETER and the UV/VIS. The results of tests run on the same control for over five months can be seen in Table 1.

The new equation gave A = 1.543 and B = 1.128 for the UV constants and the old procedure gave an A = 1.516 and B = 1.142. The *A* and *B* values calculated using the new equation gave results that were within 3% of the carboxyhemoglobin values measured with the CO-OXIMETER for a theoretical 100%, 50%, 25%, 12.5%, 6.25%, 3.13%, and 0%. The *A* and *B* values obtained using the old method gave results that were off by as much as 100% from the CO-OXIMETER values at low concentrations (Table 2).

412 JOURNAL OF FORENSIC SCIENCES

 TABLE 2—Comparison of assumed COHB levels and actual measured levels using the CO-OXIMETER, old method, and new method.

Assumed	CO-OX	Old Way	New Way	
100	92.4	99.2	92.8	
100	92.8	99.7	93.2	
100	92.4	99.3	92.9	
100	92.5	100.3	93.8	
100	91.8	100.9	94.3	
50	50.5	53.2	51.4	
50	51.0	52.3	50.5	
50	51.0	52.6	50.7	
50	51.1	52.9	51.0	
50	51.2	53.0	51.2	
25	25.0	23.4	24.4	
25	24.9	23.1	24.2	
25	25.1	23.3	24.4	
25	24.3	22.8	23.9	
25	24.8	23.3	24.4	
12.5	12.1	9.1	11.6	
12.5	12.5	9.2	11.6	
12.5	12.0	9.5	12.0	
12.5	12.3	9.3	11.8	
12.5	12.0	9.7	12.1	
6.25	5.8	2.6	5.7	
6.25	5.5	2.6	5.7	
6.25	5.7	2.9	6.0	
6.25	5.6	2.5	5.6	
6.25	5.5	2.3	5.4	
3.13	4.0	0.5	3.8	
3.13	3.8	0.5	3.8	
3.13	4.0	0.3	3.7	
3.13	3.7	0.3	3.6	
3.13	3.6	0.4	3.8	
0	3.2	0.0	3.3	
0	3.4	0.0	3.2	
0	3.3	0.0	3.1	
0	3.6	0.0	3.0	
0	3.5	0.0	3.4	

Conclusion

The new method provided reliable whole blood carboxyhemoglobin standards stable for more than four months when stored at 4° C. This is a substantial improvement over the prior commercially available controls, which were stable for only one month. The necessary constants for use in calculating COHB by spectrophotometry are reliably and efficiently provided by the present derived equations for *A* and *B*. The calculated *A* and *B* constants gave values which were a substantial improvement over values measured using the A and B constants from the old method (Table 2).

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