Determination of total carbon dioxide in beer and soft drinks by gas diffusion and flow injection analysis

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A gas diffusion FIA method for determination of total CO₂ in beer and soft drinks is described. The composition of the acceptor stream for diffused carbon dioxide is critical. Bromocresol purple has been selected among a large number of tested pH indicators and if the detection is made at a wavelength of 430 nm, stable baseline conditions and positive deflections of the resulting FIA peaks are obtained. The selected indicator is combined with a linear pH buffer. A simple and practical graphical method for determining a suitable starting pH of the acceptor stream, as well as the expected dynamic range and the linearity of the calibration graph, is presented.

Introduction

Determination of total carbon dioxide (CO₂) by gas diffusion FIA was first described by Baadenhuijsen and Seuren-Jacobs [1] for plasma samples. In this method, a silicone-rubber membrane in a gas diffusion cell and a combined carrier/acidic reagent stream was used into which 50 μL of the sample was injected. Cresol red was used as indicator and detection was performed at 410 nm. This basic FIA system has been developed further [2, 3] and theoretical considerations of the gas diffusion process have been presented [4, 5].

In a gas diffusion FIA system, the number of variables is large [6], so it is necessary to carry out screening in order to select significant variables. The response function determines the number and character of significant variables. Sensitivity has most frequently been used as a response function [7, 8]. For beer and soft drink samples, however, sensitivity is not a concern since these samples contain large amounts of total CO₂. Instead, the dynamic range becomes an important issue because the indicator can be totally consumed by the diffused gas. Consequently, a pH buffer must be added to the acceptor stream so that the major portion of the diffused gas reacts with the buffer, thereby enhancing the dynamic range.

For high concentrations of total CO₂ in the sample, a manifold, such as the one described by Kuban and Dasgupta [3], is preferred. In this approach, the sample is injected into a separate carrier stream and then merged with the acid reagent which is used to convert carbonate to gaseous carbon dioxide. The concentration of added acid across the entire sample zone is thus maintained at a constant value. This is in contrast to a single line arrangement where the added acid concentration attains a minimum value when the sample concentration is at its maximum value; this occurs in the centre of the sample plug. The acid concentration must be high enough to cause a sufficient decrease in sample pH so that CO₂ is quantitatively formed. A pH value below 4 is required, i.e. at least 2 pKₐ units below the first dissociation constant of the protonized acid. The choice of pH indicator for the acceptor stream influences the sensitivity and the baseline stability. Considering baseline stability alone, a low absorbance value is preferred when the system is in a stand-by mode. Thus, an increase in the absorbance is obtained when samples containing carbonate are injected.

This paper describes a procedure and a gas diffusion FIA system for determination of total CO₂ in beer and soft drinks. A simple evaluation method for different acceptor solutions with respect to expected dynamic range and expected linearity of the calibration graph, as well as results from runs of real samples, is presented.

Experimental

Instruments and apparatus

A commercial flow injection system (FIAstar 5020 and 5023, Tecator) was used. The gas diffusion cell (Chemifold V, Tecator) was provided with a PTFE gas diffusion membrane (homogeneous plumber’s tape); the FIA manifold is shown in figure 1. The sample volume was 100 μL. The mixing coil length was 30 cm, and the inner diameter was 0.7 mm.

Indicator screening experiments entailed collection of wavelength spectra for various indicator solutions in the range 350–800 nm by means of a diode array spectrophotometer (Hewlett-Packard 8452A).

Reagents, solutions and samples

Deionized water was used as carrier and reagent R1 consisted of 0.2 M sulphuric acid. Reagent R2 was the pH indicator solution: bromocresol purple. A stock solution of 0.02 M bromocresol purple was diluted 1:100 before use.

![Figure 1. FIA manifold for determination of total CO₂](image)

Figure 1. FIA manifold for determination of total CO₂; S = sample; C = carrier; R1 = H₂SO₄; R2 = indicator (acceptor) solution; and G.D. = gas diffusion cell.
Table 1. Screening results for all indicators investigated.

| Indicator          | Concentration (g/l) | Isobestic point (nm) | Suitable wavelength (nm) | Corresponding absorbance (A.U.) |
|--------------------|---------------------|----------------------|--------------------------|--------------------------------|
| Bromophenol blue   | 0.07                | 440                  |                          | pH 4: 1.00, pH 9: 0.16         |
| Bromocresol green  | 0.028               | 510                  | 440                      | pH 4: 0.70, pH 9: 0.15         |
| Bromocresol purple | 0.027               | 490                  | 430                      | pH 4: 1.00, pH 9: 0.10         |
| Cresol red         | 0.033               | 485                  | 430                      | pH 4: 1.90, pH 9: 0.40         |

*This indicator had complex spectra.

The solution was prepared by dissolving 0.1 g of bromocresol purple in 18.5 ml 0.01 M NaOH and adding water to a final volume of 250 ml. R2 is prepared by taking a 10 ml portion of the indicator stock solution and mixing with 1 ml of a linear buffer, which was prepared according to Aaström [9]. This buffer consisted of formic acid (0.0025 M), acetic acid (0.0150 M), malonic acid (0.012 M), piperazine (0.0218 M), Bis-tris (0.0144 M), N-methylmorpholine (0.0226 M), N-methyldiethanolamine (0.0223 M) and sodium chloride (1.0 M) and was pH-adjusted by addition of NaOH which corresponded to a concentration of 0.034 M NaOH in the final solution. Carbonate standard solutions were prepared from sodium hydrogen carbonate. Deionized water was used throughout and all chemicals were of reagent grade. The samples were all chilled at +4°C.

When the wavelength screening of different potential indicators for R2 was performed, the following pH buffers were used: pH 4–0.05 M KH-phthalate; pH 7–0.025 M phosphate; pH 9–0.01 M borax. A stock solution of each pH indicator was prepared by taking a certain amount of the indicator and then diluting it 1:15 with addition of the various buffer solutions. Final concentrations for four of the indicators are given in Table 1.

Designing the FIA system

Three variables in the FIA system shown in figure 1 were subjected to a two-level reduced factorial design, namely the sulphuric acid concentration of R1 (0.2/2.0 M), the coil length (30/60 cm), and the coil inner diameter (0.5/0.7 mm). All other variables were kept constant. None of the three studied variables was found to be significant when the peak height and the repeatability were employed as responses. The variable values selected are shown in figure 1.

Results and discussion

Screening of indicators

More than 20 indicators were tested for possible use when composing reagent R2. Spectra were recorded for each indicator at three different pH values: 4, 7, and 9. In figure 2 a typical set of spectra is shown for bromocresol purple. For most indicators an isobestic point is found. On either side of this isobestic point, absorbance maxima appear. At these maxima, the absorbance is either increasing or decreasing as a function of pH. Because carbon dioxide diffusing through the membrane and entering the acceptor stream R2 will cause a decrease in pH, it is desirable that this decrease in pH will result in a corresponding increase in indicator absorbance. This is fulfilled in the 390–470 nm region for bromocresol purple, see figure 2. Thus, a positive deflection of the FIA response curve is registered in this wavelength range. A maximum deflection is found at the absorbance maximum attained by the pH 4 curve, see the vertical full line at 430 nm in figure 2.

The indicators listed in table 1 could be used for composing the reagent R2. However, the choice is dictated by the additional requirement that the background absorbance of the indicator at a starting pH value of 8–9 must be low. If the starting absorbance is too high, the validity of Beer's law might be jeopardized, resulting in non-linear calibration graphs at high analyte concentrations. Bromocresol purple was finally selected because a stable and reliable baseline appeared at 430 nm, in contrast to the baselines obtained for bromocresol green and cresol red. Bromophenol blue might be the second-best choice.

Composing the acceptor solution (R2)

The shape of the calibration graph is governed by the starting pH value and the buffer capacity of the acceptor solution, R2. Figure 3 shows the results of a simple evaluation method for an acceptor solution consisting of bromocresol purple and linear buffer. The starting pH of the acceptor solution was 12. Titration with hydrochloric acid and simultaneous recording of the absorbance values at 430 nm were then performed. The results in figure 3 provide useful information regarding a
suitable starting pH value of the acceptor solution, the dynamic range and, finally, the expected linearity of the calibration graph.

In figure 4, four different types of calibration graphs are presented. The shape of each one of these can be explained by referring to figure 3. For curve (a) in figure 4, a starting pH value which was too high was selected; for curve (b), it appears that appropriate conditions were selected. For curve (c), a starting pH value which was too low was selected. For curve (d), a ‘zero suppression’ was accomplished by starting at a high pH value; which results in poor sensitivity at low concentrations of carbonate.

Real samples

Results for a limited selection of beer and soft drinks with the FIA system shown in figure 1, using bromocresol purple and a linear buffer as an acceptor stream for the diffused carbon dioxide, are shown in table 2. The numbers of respective replicate runs are denoted by ‘n’ in table 2. Standard deviation values are given in the same table. The reported analytical results are in full agreement with results obtained by using the commonly applied titration method.

As expected, the sample pretreatment procedure is crucial. When opening a bottle or a can, carbon dioxide evolves and will be lost if the partial pressure of the gas is not substantially lowered. A direct detection of carbon dioxide in the original sample by, for instance, a CO₂ electrode is, for obvious reasons, impossible. However, by increasing the pH of the sample by adding sodium hydroxide, volatile carbon dioxide is transferred to carbonate and determination of total CO₂ in the original sample can be accomplished. Triethanolamine, in combination with sodium hydroxide, is suitable for adding to the sample, but the exact procedure for this addition must be carefully defined to guarantee reproducible results.

Conclusions

When large concentrations of analyte, such as carbonate, sulphite, and ammonium, are to be determined by gas diffusion in an FIA system, the composition of the acceptor stream is critical. The acid-base indicators usually employed cannot be added to the acceptor stream in sufficient quantities to produce the buffering capacity needed for a wide dynamic range. A separate buffer is required. Theoretically, the dynamic range can be calculated for any given composition of the acceptor stream using pKₐ values. In practice, however, an evaluation method such as the one described and illustrated in figure 3 is to be preferred.

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