The design of dilution loops for continuous-flow analysis

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Introduction

The sensitivity of many colorimetric and other methods of detection used in continuous-flow analysis means that samples frequently need to be diluted to bring them into the range of a particular method. On-line dilution can take the form of a dialyser, in which a proportion of the analyte diffuses through a permeable membrane into a recipient stream, or a dilution loop, where the sample is diluted, usually with water, then led back to the inlet side of the pump (hence the term 'loop'), where a small proportion is resampled. Because the resampled flow is often unsegmented, unlike the original sample flow in which the inter-sample air bubble separates the sample from the wash liquid, the dispersion in a dilution loop can be high, resulting in a lower optimum sample rate.

However, by the correct choice of fittings, and by segmenting the resample stream where appropriate, dispersion in a dilution loop can be reduced to a level where it is not the limiting factor in system performance.

Dilution factor and flow rate

It is usually possible to achieve a given dilution factor by several combinations of flow rate. For example, a dilution of about 11 times can be effected with a sample line of 0.1 ml/min and diluent 1.0 ml/min, 0.23 sample and 2.0 diluent, 0.32 and 3.4, and so on. Considerations applying to the diluent flow rate are the stability of flow and segmentation, delay time, back pressure and diluent consumption. When a roller in a peristaltic pump lifts off the pump tube there is a momentary reduction in flow rate as the tube expands to its original shape. Thus the flow during one cycle of the pump, i.e. roller to roller, is variable, or pulsed. This effect is more marked with large diameter tubes, and the rapid variation in flow during one pump cycle can cause irregular bubble injection: for this reason tubes with a maximum flow of 2 ml/min are used when possible. When higher flow rates are necessary, the diluent can be added in two or more stages, with or without intermediate mixing, to promote regular flow at the air injection point.

It is always desirable to minimize delay time, and this corresponds to high flow rates or narrow diameter tubing. For a given diameter of transmission tubing in the lengths used in a dilution loop it is safe to pass up to eight times the flow developed by a pump tube of similar diameter.

For example, the pump tube nearest in diameter to the 2 mm glass tubing used in AAI1 systems has a flow rate of 2.4 ml/min: thus about 10 ml/min can be pumped through a 2 mm dilution loop without developing excess back pressure, which lowers the effective pumping rate and creates problems with air injection.

The lower limit on sample flow rate is mainly determined by the size of any solids in the sample, and is normally 0.1 ml/min (0.015 in i.d.), with 0.05 ml/min (0.010 in i.d.) occasionally being used.

Thus the optimum range of dilution is up to 20 times (0.1 ml/min sample, 2 ml/min diluent), with up to about 150 times (0.05 sample, 2 × 3.9 diluent) possible. The flow rate of the resample line is generally determined by the method, and it is this flow rate which determines the choice of debubbler for unsegmented resample lines.

Unsegmented resample lines

As with all parts of a manifold, the aim in selecting a debubbler for an unsegmented resample line is to minimize dispersion. This occurs in two parts of the resample fitting: the part connected to the resample pump tube and the part containing the segmented flow. Figure 1 shows three types of debubbler sometimes used as resample fittings. The C3 contains a ramp to remove extra or oversize bubbles – this is useful on AAI1 systems, but unnecessary on AAI1 systems with regular air injection (the intersample air bubble in a dilution system will be necessarily small). The A2 is better in this respect, but the capillary arm of both these fittings is 1.0 mm i.d., the same as that of a 0.6 ml/min pump tube. If a lower flow rate resample pump tube is used this capillary is drained slowly, creating excessive dispersion. For flow rates up to 0.6 ml/min the so-called ‘modified AO’ is used, with 0.025 in i.d. acidflex tubing inserted into the side arm until bubbles just pass over the top without being drawn into the resample line.

Segmented resample lines

If a resample fitting such as the A12 or A16 is used (see figure 1), a small proportion of each bubble in the diluted stream will be drawn into the resample pump tube, segmenting the stream and thereby reducing dispersion. This reduction in dispersion can be quantified, as shown in the next section. Because the AAI1 system uses air injection synchronized with the pump rollers, the small resampled bubbles always occupy the same position in the resample pump tube relative to the rollers, and thus the short-term delivery is constant.
However, the volume of each resampled bubble depends on the velocity at which the main bubble passes across the take-off insert, and this is dependent on not only the total flow rate but also, due to the pulsed flow, on the moment in the pump cycle at which the bubble reaches the insert. The arrival time depends on the number of bubbles and liquid segments between the air injection point and the resample insert, and this is chiefly determined by the flow rate of the diluent pump tube. Because the flow rate of the tube slowly drops with age, the number of segments between these points increases, and the position which a particular bubble (for example the 60th after the injection point) occupies in the dilution loop slowly moves backwards. Thus the volume of resampled air, and therefore the delivery of liquid and the sensitivity of the method, changes with time. More serious is the short-term variation which occurs when the bubble reaches the resample insert just before or after the instant of least velocity in the cycle, as here very short variations in arrival time, such as may be caused by irregular roller spacing or intersample air bubble compression, cause large proportional changes in the resampled bubble volume, and hence a noisy response.

These problems are overcome by using a system whereby the arrival time of a bubble at the resample point does not depend on the number of segments in the loop, but only on the time within each pump cycle. This is achieved by removing the original bubble just before the resample point and immediately introducing another. The new bubble is injected in synchrony with the pump, and, because the distance between its injection point and the resample point is so short, variations in flow rate have a negligible effect on its arrival time. Part No. 170-G063-01 (see figure 1) is such a debubble/rebubble fitting, which can be used with an A12 or A16, and 104-G036-01 is a more recent type which also contains two sapphire inserts for resampling.

**Dispersion**

The dispersion in a dilution loop arises from three sources: the segmented stream from the pump to the resample fitting, the resample fitting itself, and the resample pump tube.
The theoretical contribution of each can be calculated, using the equation described by Hrdina [1] which relates dispersion \( \sigma \) in seconds in unsegmented streams to diameter \( d \) in mm and time \( t \) in s:

\[
4\sigma = 50 \frac{dV't}{d}
\]  

(1)

and the equation derived by Snyder and Adler [2] relating dispersion in segmented streams to flow rate \( F \) in ml/s, internal diameter \( d \) in cm, segmentation frequency \( n \) in bubbles/s, liquid viscosity \( \eta \) in poise, surface tension \( \gamma \) in dynes/cm, a mass transfer coefficient \( D \) that varies with the nature of the sample molecules and the viscosity of the liquid, and dwell time \( t \) in s.

\[
\sigma^2 = \left[ \frac{538 d^{2/3} (F + 0.92 d^3 n)^{5/3} \eta^{7/3}}{\gamma^{2/3} F D} + \frac{1}{n} \right] + \frac{2.25 (F + 0.92 d^3 n)^{5/3} \eta^{2/3}}{\gamma^{2/3} F d^{4/3} t} (2)
\]

A typical value for \( \eta \) is 0.009, for \( \gamma \) 32, and for \( D \) \( 5 \times 10^{-5} \). Further discussions of the applications of this formula can be found in two papers by Snyder [3 and 4].

In this equation the dispersion is expressed as the variance, \( \sigma^2 \), of a Gaussian curve, in s^2.

Table 1 shows the contribution from each part of a dilution loop employing commonly-used flow rates in four configurations. Configurations \((a)\) and \((b)\), shown in figure 2, are both unsegmented, but \((a)\) uses an A2 resample fitting and \((b)\) uses a modified AO.

| Hardware description | Dispersio... | Within fitting | Resample pump tube | Total |
|----------------------|--------------|----------------|--------------------|-------|
| \((a)\) 2 mm glassware A2 unsegmented | 7.7 | 10.1 | 19.8 | 23.5 |
| \((b)\) 2 mm glassware modified AO unsegmented | 7.7 | 4.9 | 19.8 | 21.8 |
| \((c)\) 2 mm glassware 170-G063-01 + A16 segmented | 7.7 | 12.2 | 2.9 | 14.7 |
| \((d)\) 1mm glassware 104-G036-01 segmented | 5.4 | 3.8 | 2.9 | 7.2 |

Choice between segmented and unsegmented resample lines

It is apparent from equation (3) that the part of a system with the greatest dispersion dominates the total dispersion, and that efforts to minimize dispersion must therefore be centred here. Applying this principle to dilution loops, it is clear that the benefits of segmented resample lines are only seen when the dilution loop is one of the major contributors to system dispersion. If this is not the case it is usually preferable to use unsegmented resample lines as these are the simpler type. The figures for dispersion in table 1 can be compared to those for a 1.5 mm flow cell with a pull-through of 1.2 ml/min at 13 s, a 2 mm flow cell with the same pull-through at 23 s, a sample line segmented by the normal inter-sample air bubble at about 15 s, and the segmented stream of a manifold incorporating a heating bath with a flow of 1.5 ml/min and a delay of 5 min at 11 s.
Comparative benefits of dilution loops and dialysers

Table 2 shows a comparison of dilution and dialysis. The principal advantages of dilution loops are their insensitivity to temperature variations and the ease of predetermining the dilution factor. As recovery in a dialyser depends on path-length, membrane type and condition, temperature, flow rate of donor and recipient streams, back pressure on both sides of the membrane, and ionic conditions, it is seldom possible to predict an exact dilution. The primary advantage of dialysis is the separation of solids and high molecular weight solutes from the analyte. The book by Furman [5] is a useful reference for this and other aspects of system design.

Conclusion

The design of dilution loops is governed by simple principles. Applying these principles to the requirements of a method allows systems with low dispersion and robust performance to be constructed. The theoretical dispersion of the individual parts of a system can be predicted, and this information can be used to select the most suitable dilution method for a particular application.

References

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References

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1. MANKS, D. P., Journal of Automatic Chemistry, 3 (1981), 119.
2. MALMSTADT, H. V., in Topics in Automatic Chemistry, Ed. Stockwell, P. B. and Foreman, J. K. (Horwood, Chichester, 1978), p. 68.

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