DEVELOPMENTS IN NON-DISPERSIVE INFRA-RED ANALYSERS

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Non-dispersive infra-red analysers have long been used as plant instruments. The best-known form is the "IRGA" instrument [4] based on the original detector developed by Luft. In recent years an alternative form has been developed [1, 2] in which the spectral discrimination is given by narrow-band interference filters rather than by the absorption characteristics of the detector. The instruments described in this paper, for which patent applications have been made [3], have important advantages over previous conventional analysers. In particular, while possessing the same freedom from source and electronic drifts as existing double-beam instruments, they are also insensitive to changes in sample-tube transmission. Dirty or corrosive samples can therefore be accommodated by the instruments, whereas conventional analysers would not be free from error.

Interference-filter instruments are of greatest potential where the absorption spectra are such that all the wanted information can be unambiguously derived from a single absorption line at a relatively short wavelength, say less than 6 μ. In this case the high degree of spectral resolution offered by the filter can be combined with the high radiation sensitivity of a photoconductive detector to give an arrangement capable of detecting very small quantities of trace impurities. The single optical path also results in a substantially simpler instrument with possibly greater reliability and reduced cost.

One such instrument for the analysis of hydrogen fluoride in process-gas streams has been described by Harrison and Reid [1]. This device, called for convenience the Mark Ia analyser, has been redesigned as the Mark Ib to give greatly improved performance for analysis of hydrogen fluoride, and a further design (the Mark Ic) can clearly be the basis for instruments to deal with a variety of other problems.

THE MARK Ia ANALYSER

Details have already been published [1], but a brief description will demonstrate the principles of these interference-filter instruments. Fig. 1 shows the layout of the instrument.
Fig. 1.—General arrangement of Mark Ia analyser.

The instrument works by comparing the intensity of light transmitted at a wavelength absorbed by the sample with that of an adjacent absorption-free wavelength (Fig. 2). The necessary wavelength discrimination is obtained by means of an appropriate interference filter; in the hydrogen fluoride detector this is a mica-spaced filter of comparatively high resolution. The shift from the "sample" wavelength to the "reference" wavelength is achieved by tilting the filter through an angle sufficient to change the wavelength of its pass band by the required amount.

A secondary filter isolates the desired transmission bands of the high-resolution mica filter and also allows the "sample" and "reference" beams to be balanced in intensity with no sample in the absorption cell. The high-resolution filter is tilted back and forth at 10 c/s, so that a 10 c/s output signal, proportional to the difference in intensity between sample and reference beams, is given by the lead sulphide detector. If the two beams are initially balanced, this signal is a direct measure of the absorption at the sample wavelength.

If the sample tube suffers a loss in transmission, both sample and reference beams are equally affected, provided that the loss is uniform over the narrow wavelength band concerned; thus, an error is introduced proportional to the extent of the loss in transmission. To correct for these changes and for any changes in photocell sensitivity and amplifier gain, a standardizing chopper was used in the Mark Ia instrument. This introduced a fixed interruption of about 20 per cent in the light path, so that the gain of the amplifier can be adjusted to give a standard reading from the instrument.

The Mark Ia has no built-in compensation for relative changes in lamp
emission at the two wavelengths caused by changes in lamp temperature; it
was obviated by using a photoelectrically controlled lamp power supply.

Two instruments of this type have been in use for nearly a year, detecting
hydrogen fluoride in the process-gas stream of the Diffusion Plant of the
U.K.A.E.A. at Capenhurst. At the end of its period of service one was
found to have suffered a loss of 95 per cent of its sample-tube optical trans­
mission. It could still be standardized and its calibration remained un­
affected. A conventional form of analyser would have been heavily handi­
capped in such circumstances, and it is doubtful whether any of the alter­
native forms of instrument would have given satisfactory service.

THE MARK I b ANALYSER

The instrument

The layout of this instrument is shown in Fig. 3. The optical parts have
been changed in some particulars: a shutter which interrupts the light beam
at about 1000 c/s has been introduced and the standardizing chopper re­
moved. The primary filter relay now operates at a lower frequency, about
2 c/s. The light falling on the detector is thus interrupted at 1000 c/s, with
an extra modulation at 2 c/s imparted by the tilting primary filter.

The detector output signal is amplified by two amplifiers, in cascade.
The first amplifier is tubed at 1000 c/s; this signal is then rectified and the
2 c/s sample signal modulation extracted and fed forward to the second
amplifier. Since the 2 c/s modulation is only a few percent of the total
intensity, the D.C. component of the rectified 1000 c/s signal is a fairly
accurate measure of the total intensity passing through the system. This
D.C. signal is used in a servo arrangement to control the voltage applied
to the lead sulphide cell, and hence to change its sensitivity so that a con­
stant D.C. output is obtained regardless of changes in sampletube trans­
mission, detector sensitivity and first amplifier gain. The second amplifier brings the 2 c/s signal up to a measurable level before it is phase-sensitively rectified to give the final output. This output is proportional to the absorption in the sample.

**Simple analysis of method of compensation**

A simple analysis makes this clear. The quantities independent of wavelength may be defined as: $G$, 1000 c/s amplifier gain; $g$, 2 c/s amplifier gain; $T$, sample-tube transmission; and $R$, response of photocell in volts per incident photon.

The quantities depending on wavelength are: $Q_1$, $Q_2$ = the photons/second entering the sample tube in the appropriate wavelength intervals; and $\alpha_1$, $\alpha_2$ = the corresponding absorption in the sample. Here suffix 1 indicates the reference wavelength, and suffix 2 the sample wavelength. By definition $\alpha_1 = 0$.

If $V_1$ and $V_2$ are the 1000 c/s amplifier output voltages (see Fig. 3), then

$$V_1 = GRT \cdot Q_1 \quad \text{and} \quad V_2 = GRT \cdot Q_2 (1 - \alpha_2).$$

The action of the servo-system is to arrange that $(V_1 + V_2)/2$ is a constant, $K$, so that we can write

$$GRT[Q_1 + Q_2 (1 - \alpha_2)] = 2K.$$ 

The 2 c/s output signal,

$$S = g(V_1 - V_2),$$

is then given by

$$S = g \cdot 2K \frac{[Q_1 - Q_2 (1 - \alpha_2)]}{[Q_1 + Q_2 (1 - \alpha_2)]}.$$

The secondary filter is so arranged that $Q_1 = Q_2$; hence

$$S = g \cdot 2K \frac{\alpha_2}{(1 - \alpha_2)}.$$

Since $\alpha_2$ is very small for low sample levels, $S = g \cdot 2K \alpha_2$ to a good approximation.

Thus the only instrumental parameter left uncompensated is the gain of the 2 c/s amplifier, $g$, which is low and practically constant.

If the lamp temperature changes so that $Q_1 \neq Q_2$, then

$$S = g \cdot 2K \frac{[Q_1/Q_2] - (1 - \alpha_2)}{[Q_1/Q_2] + 1}$$

and it is clear that, for small changes of source temperature, a degree of compensation is afforded, since $Q_1/Q_2$ will change less rapidly than either
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Fig. 4.—Typical calibration of hydrogen fluoride meter.

$Q_1$ or $Q_2$. It is therefore possible to dispense with any special lamp power supply: a constant voltage transformer is all that is needed.

The chopping at 1000 c/s effects further improvement over the Mark Ia analyser, since the signal-to-noise ratio of the lead sulphide cell is much greater at this frequency than at 2 c/s.

**Performance of the Mark Ib analysers**

Although obviously suitable for other applications, the Mark Ib has been used primarily as a hydrogen fluoride detector.

The limit of performance for hydrogen fluoride could not be assessed directly. Hydrogen fluoride is a difficult gas to handle; at high pressure it polymerizes easily, and moreover it is highly reactive and adsorbs strongly. In consequence, stable concentrations of known value appear practically impossible to achieve. The detection limit for the instrument had, therefore, to be estimated from signal-to-noise ratios. It appears that the instrument should detect a partial pressure of less than 2 μ of hydrogen fluoride in a total pressure of 1 atm. A full-scale deflection for 200 μ partial pressure at a noise level of ± 1 per cent is therefore achievable. A typical calibration curve is given in Fig. 4; the law of the calibration is as previously described for the Mark Ia analyser [1].

**THE MARK Ic ANALYSER**

**The instrument**

Although the Mark Ib analysers are satisfactory for the application for which they were designed, further improvements are possible. The main disadvantages of the Mark Ib are: firstly, because the high-resolution filter isolates a single rotational fine-structure line, a small part only of the available spectral information is used; and, secondly, because a high-resolution filter is used, the nominal aperture of the system must be kept low (f/8 to f/10).
The Mark Ic has been designed to avoid these defects. In this instrument, the primary filter is made with two areas of different transmission wavelengths. This is moved by the relay so that the light beam passes alternately through each half of the filter, thus achieving the wavelength stepping action. This system allows a larger spectral signal to be obtained, since it is now possible to use a complete band or branch instead of a single line.

Because the filter is of comparatively low resolution, the restriction on nominal aperture can be relaxed, and apertures of f/4 are practicable. The increase in aperture can be achieved by using the sample tube as a collimator. In this form, the lamp is brought close to one window of the sample tube, and the primary filter is arranged close to the other window. A slit is placed immediately in front of the secondary filter, and the detector is as close as possible behind it, as shown in Fig. 5. The light path in free air is thus reduced to less than an inch, which is an advantage if atmospheric gases such as water vapour or carbon dioxide are to be detected.

With a fixed secondary filter, another method is required for balancing the instrument at zero sample. This is achieved by using another set of the relay contacts to switch the gain of the amplifier, so that balancing is now done electronically and not optically.

In other respects, i.e. the use of a 1000 c/s chopper, two-section amplifier and stabilizing feedback, the instrument is similar to the Mark Ib.

**Performance of the Mark Ic analyser**

This instrument has been made as a water-vapour detector, operating at the water band at 2.7 μ. With a nominal aperture of f/4, and a sample tube 30 cm long, the ultimate sensitivity (signal-to-noise ratio of one) of the instrument is 1.5 μ water vapour in a total pressure of one atmosphere.
FURTHER DEVELOPMENTS

These instruments are new and there appears to be no reason why they cannot be made to detect substances having absorption spectra within at least the wavelength range 0.4-7 μ, i.e. the range for which detectors of short response time are available. This range would allow carbon monoxide, carbon dioxide, nitrous oxide, ammonia, methane, hydrogen chloride, hydrogen bromide and other compounds to be detected. It also appears possible to construct liquid absorptiometers, working on the same principles.

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REFERENCES