A Constant-Feed Direct-Current Arc

GEOLOGICAL SURVEY BULLETIN 1084-J
A Constant-Feed Direct-Current Arc

By C. S. ANNELL and A. W. HELZ

CONTRIBUTIONS TO GEOCHEMISTRY

GEOLOGICAL SURVEY BULLETIN 1084-J

A study of methods for introducing samples into a d-c arc, accompanied by low fractional distillation

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A CONSTANT-FEED DIRECT-CURRENT ARC

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ABSTRACT

The development of a constant-feed method for the continuous vaporization of successive increments of powdered rock and mineral samples into a 10-ampere direct-current arc is described. Long, deep-bore graphite electrodes having a 0.092-inch outer diameter and a 1.5-inch bore depth of 0.046-inch diameter are used as sample anodes. Concomitant vaporization and excitation of such elements as titanium, aluminum, silicon, copper, germanium, arsenic, and lead are obtained by gradually moving the electrode into the arc through a channel in a brass, water-cooled collar. The collar is placed on an electrode holder which was designed to permit controlled atmospheres to pass upward around the electrodes. A gas mixture flowing at the rate of 14 cubic feet per hour argon and 7 cubic feet per hour oxygen was selected to suppress cyanogen band interference in the spectra. Samples mixed with graphite powder burn more evenly in the arc and show less selective volatilization than untreated samples. The addition of lithium borates to the sample and graphite generally enhances atom lines but tends to suppress ion lines. Tests of matrix influence are made in which intensity ratios of atom lines of silicon, aluminum, magnesium, and iron are compared with ion lines of iron, titanium, and calcium (in NaCl and ZrO₂ bases) for sample-graphite mixtures of 1:1, and sample lithium tetraborate graphite mixtures of 1:2:3. The ranges of percent deviation from a unity intensity ratio, which can be considered a measure of the matrix effect of NaCl and ZrO₂, are 13–32 percent for atom lines in the graphite admixture, and 13–33 percent for the same lines in the lithium-buffered mixture. The ion lines show 23–55 percent and 16–27 percent deviations, respectively. Comparative results with the same sample compositions are given for more conventional ¼-inch cupped graphite electrodes, for which an electrode holder is described. The above atom line intensity ratios for these electrodes deviate from unity by 30–65 percent and 12–36 percent, respectively. Comparable deviations of the ion line ratios are 159–286 percent, and 46–139 percent. Marked reductions in line broadening and self reversal are noted in the constant-feed direct-current arc.

INTRODUCTION

The study of a constant-feed method for introducing samples into the arc for spectrochemical analysis was the primary purpose of this investigation. Such a method is envisioned as one in which the sample is brought into the arc in many successive increments, each unit of sample undergoing the same processes in the arc independently. The
attendant decrease in selective volatilization would possess several advantages (Ahrens, 1950, p. 68, 85, 115):

1. The variations in average electron energies during the course of an exposure would be diminished.
2. Fewer internal standards would be suitable for a greater number of elements to be determined.
3. A proper buffer could be added to the sample and be present at a relatively constant rate throughout the arcing period.

In the spectrochemical analysis of rocks and minerals large diversities of chemical compositions and physical forms are handled. Accurate analyses require standards similar to the samples analyzed. The work reported here is directed towards reducing the number of standards required and generalizing procedures to handle a larger variety of materials.

Methods directed toward accomplishing the above objectives and applicable to d-c arc excitation have been reported. Semenov (1956) describes the use of a “carbonized straw” of 0.9 mm outside diameter, 0.8 mm inside diameter, and 80 mm length, which contains the sample. This “straw” is fed into the arc through a hollow, water-cooled anode by a motor-driven mechanism, the speed of which can be varied depending on the nature of the sample. A modification of this method is described by Noar (1957). A pellet is made from a mixture of sample, Li₂CO₃, and cellulose powder. This pellet is fed into the arc by a motor-driven push rod, through a hollow carbon anode. Heat is conducted away from the electrodes by brass finned collars.

A method for introducing powdered ore samples between two carbon electrodes inclined 45° from the horizontal is described by Toishi (1940). Another method for introducing powdered samples directly into an a-c arc is described by Rusanov and Khitrov (1958). The work of Danielsson and Sundkvist (1959) illustrates the excellent precision obtainable when dissolution of the sample is combined with a constant-feed principle. In this report our investigations were confined to the use of powdered rock samples.

**ELECTRODE DESIGN**

Semenov’s idea for introducing the sample into the arc seemed attractive but too complex for the routine handling of a large variety of samples. It was attempted in this study to determine whether an effective constant-feed action could be obtained with a long, thin electrode that would be relatively easy to load. Electrodes of the following dimensions (in inches) were cut from graphite rods 0.125 inch in diameter:
Burning-behavior and moving-plate tests of different rock samples with and without admixtures were made with these electrodes and compared with conventional 0.25-inch-diameter, shallow-crater electrodes similar to ASTM, S-8 (Am. Soc. Testing Materials, 1957, p. 38), herein designated as "cupped."

Electrode No. 1 showed progress toward the constant-feed principle, but selective volatilization was still very apparent. The thinner walls of electrode No. 2 provided further improvement. The deeper bore electrode No. 3, as may be seen from figure 32, provided a longer period of uniform volatilization than electrode No. 2. Lines of a few elements having dissimilar excitation and vaporization properties are illustrated in figure 32.

Self reversal and line broadening (for example, Mg 2852.1) have been significantly reduced with the use of the deep-bore electrode. This gives further evidence of the benefit of the constant-feed process, whereby the concentration of any one atom species is more uniformly distributed throughout the arcing period.

**SAMPLE ADMIXTURES**

A few admixed buffers and (or) fluxes were tested with different samples to observe their effect on volatility, line intensity, and general burning qualities. The choice of an admixture was guided by its (a) ability to maintain steadier excitation energies, (b) properties of reducing differences in the physical and chemical forms of materials, (c) convenience in handling, and (d) availability. A d-c arc of 9–10 amperes was used throughout these tests.

**UNTREATED SAMPLES**

When the sample was arced in either the cupped or the deep-bore electrodes, without any additive, the behavior of the arc appeared steady with most silicate rocks. When a sample of limestone (NBS standard sample 1a) was arced in a deep-bore electrode, diffusion of calcium through the electrode walls was noted. The anode burning spot would extend, periodically, a few millimeters below the crater tip along the side of the electrode.

If samples were arced in the electrodes without any additives, their moving-plate spectra showed extensive fractional distillation. This would occur with both erratically and smoothly burning samples.
FIGURE 32.—Time intensity plots of Si, Ge, Ti II, and Cu lines using 0.1-inch-diameter deep-bore electrodes. A, with electrode No. 2, bore depth 0.75 inch; B, with electrode No. 3, bore depth 1.50 inches. Electrodes contain a sample of W–1, LiBO$_2$ + 2 percent Ge (added as GeO$_2$), and graphite = 1 : 4 : 6. Direct-current arc of 10 amperes; controlled atmosphere, argon-oxygen mixture with a rate of flow of 14 cfm Ar and 7 cfm O$_2$. Lines measured: Si 2435.2, Ge 2533.2, Ti II 3251.0, Cu 3247.0.

SAMPLE-GRAPHITE MIXTURES

The addition of powdered graphite to a sample generally improves its burning properties in the arc (Ricard, 1954; Ahrens, 1950, p. 40). The silicate and limestone samples burned smoothly with a 1 : 1 sample-graphite mixture, although diffusion of the charge through the electrode walls was still apparent. The diffusion problem was remedied when a gas jet was used in the arcing procedure, discussed below. The graphite aided in a more concomitant distillation of elements such as copper, titanium, sodium, aluminum, and silicon; but zirconium appeared only toward the end of the period. Graphs illustrating the improvement of uniform volatilization with the addition of graphite are given in figure 33. The graphite powder, in addition to altering the physical characteristics of the sample charge, is responsible for a host of other chemical reactions during the arcing period (Leuchs, 1950).
Moving-plate spectra of the granite standard G–1 (Fairbairn and others, 1951) with powdered graphite in a 1:1 mixture showed more fractional distillation when the cupped electrodes were used than when the deep-bore electrodes were used. However, the differences in the fractional distillation observed with the two types of electrodes decreased when larger proportions of graphite (G–1: graphite = 1:10) were mixed with the G–1.

**ADDITION OF BUFFERS AND FLUXES**

<table>
<thead>
<tr>
<th>Compound tested</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li&lt;sub&gt;2&lt;/sub&gt;CO&lt;sub&gt;3&lt;/sub&gt;</td>
<td>Harvey, 1947, p. 22; Weaver and Brattain, 1949</td>
</tr>
<tr>
<td>LiBO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Hasler, 1953</td>
</tr>
<tr>
<td>Li&lt;sub&gt;2&lt;/sub&gt;B&lt;sub&gt;4&lt;/sub&gt;O&lt;sub&gt;7&lt;/sub&gt;</td>
<td>Tingle and Matocha, 1958</td>
</tr>
<tr>
<td>Li&lt;sub&gt;2&lt;/sub&gt;GeO&lt;sub&gt;3&lt;/sub&gt;</td>
<td>None</td>
</tr>
<tr>
<td>GeO&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Frisque, 1957; Jaycox, 1958</td>
</tr>
<tr>
<td>B&lt;sub&gt;2&lt;/sub&gt;O&lt;sub&gt;3&lt;/sub&gt;</td>
<td>Gillette and others, 1954</td>
</tr>
</tbody>
</table>

The above materials were tested for effectiveness as buffers by mixing in various concentrations with powdered rock samples both with and without graphite. The d-c arc was used in a controlled atmosphere of carbon dioxide flowing at the rate of 20 cfh (cubic feet per hour). References show their use in other analytical methods.

The Li<sub>2</sub>CO<sub>3</sub> gave unsteady burning with occasional bursts of particles into the arc, which was caused by the carbon dioxide generated from its decomposition. The GeO<sub>2</sub> and Li<sub>2</sub>GeO<sub>3</sub>, in mixtures with the sample and powdered graphite, generally gave enhancement of spectral lines as compared to sample-graphite mixtures. The last two columns of table 1 give the results for a mixture of G–1:Li<sub>2</sub>GeO<sub>3</sub>: graphite = 2:6:14 relative to a mixture of G–1:graphite = 2:4. A G–1:Li<sub>2</sub>GeO<sub>3</sub>: graphite = 2:14:6 mixture, not shown in the table, gave less intense spectra than the 2:6:14 mixture when arced in either type of electrode. The entire spectrum, in general, was very weak when both deep-bore electrodes and cupped electrodes were used with Li<sub>2</sub>GeO<sub>3</sub> or GeO<sub>2</sub> mixtures not containing graphite. An exception was noted in the LiBO<sub>2</sub>–sample mixtures, where most lines were enhanced when arced with the cupped electrodes.

The Li<sub>2</sub>GeO<sub>3</sub> mixtures had a tendency to cause spattering of particles, possibly due to water inclusions or adsorbed moisture, and the GeO<sub>2</sub> mixtures resulted in rather heavy background. The B<sub>2</sub>O<sub>3</sub> mixtures burned very steadily, but line enhancement was only slightly greater than with plain graphite mixtures. In addition, a rather heavy background was created, partly caused by BO band emission.

---

1 Prepared by fusing 1 part Li<sub>2</sub>CO<sub>3</sub> and 1.5 parts GeO<sub>2</sub> with an oxygen–natural gas flame in a platinum crucible. The material was crushed and ground in a boron carbide mortar.
The more promising burning and buffering properties of the lithium borates lead to extended testing. The relative intensities for some selected lines are listed in table 1, in which results are reported for G–1 in lithium borate–graphite mixtures with both the cupped electrode and 0.75-inch deep-bore electrode No. 2.

Table 1.—Comparison of line intensities obtained with deep-bore electrode and cupped electrode using various admixtures

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.75-in. deep-bore electrode</td>
<td>Cupped electrode</td>
<td>0.75-in. deep-bore electrode</td>
</tr>
<tr>
<td>Si 2453.2</td>
<td>0.78–6.65</td>
<td>1.10</td>
<td>2.42</td>
<td>0.93</td>
<td>1.78</td>
</tr>
<tr>
<td>Al II 3055.5</td>
<td>0.51–0.81</td>
<td>1.30</td>
<td>5.48</td>
<td>2.96</td>
<td>1.43</td>
</tr>
<tr>
<td>Fe II 2676.1</td>
<td>6.86–0.60</td>
<td>0.89</td>
<td>8.68</td>
<td>1.12</td>
<td>0.84</td>
</tr>
<tr>
<td>Mg II 2802.7</td>
<td>7.61–0.40</td>
<td>0.80</td>
<td>8.40</td>
<td>0.75</td>
<td>0.56</td>
</tr>
<tr>
<td>Mg 3085.1</td>
<td>0–4.33</td>
<td>0.80</td>
<td>8.80</td>
<td>0.75</td>
<td>0.56</td>
</tr>
<tr>
<td>Fe 3047.0</td>
<td>0.96–4.14</td>
<td>0.90</td>
<td>8.58</td>
<td>1.45</td>
<td>1.38</td>
</tr>
<tr>
<td>Ti II 3065.0</td>
<td>6.81–4.05</td>
<td>0.80</td>
<td>8.40</td>
<td>0.95</td>
<td>0.85</td>
</tr>
<tr>
<td>Cu II 3178.3</td>
<td>6.09–7.02</td>
<td>1.20</td>
<td>1.00</td>
<td>1.18</td>
<td>1.19</td>
</tr>
<tr>
<td>Cu 3247.5</td>
<td>6.09–7.02</td>
<td>1.94</td>
<td>2.14</td>
<td>Bands</td>
<td>2.22</td>
</tr>
</tbody>
</table>

2 Too dark, slightly denser in graphite.
3 Too dark, considerably denser in LiBO₂.

Moving-plate spectrograms of several sample, buffer, and graphite mixtures emphasize the role graphite plays in reducing the tendency for fractional distillation when using the cupped electrode (fig. 33). Similar tests with deep-bore electrode No. 2 showed less differentiation owing to the better control of volatilization (fig. 34). The use of the deep-bore electrode No. 3 with a water-cooled jacket around the electrode further minimized volatility differences (figs. 38 and 39).

Subsequent tests with commercially available Li₂B₄O₇ gave results similar to LiBO₂ and showed less tendency to absorb moisture from the air. This provided greater ease in handling the materials during the weighing and mixing operations.

**CONTROLLED ATMOSPHERES**

References to the use of controlled atmospheres have been made in the above tests. The use of an open jet, which directed flowing gases upward along the sides of the long-bore electrodes, had contributed to a stabler arc and reduction in diffusion of the charge through the walls of the electrodes.

Earlier work with controlled atmospheres had led to the use of carbon dioxide with an open-end cylindrical tube. In addition to

*Lithium borate was prepared by fusing 1 part Li₂CO₃ and 1.7 parts H₂BO₃ using a platinum crucible in a muffle furnace at 1000°C. The fused material was ground in a boron carbide mortar.*
Figure 83.—Time-intensity plots of Si, Fe, Ti II, Cu, and Li lines using 0.25-inch cupped electrodes containing different proportions of sample and admixtures. Direct-current arc of 10 amperes; controlled atmosphere, CO₂ with a flow of 20 cfh. Lines measured: Si 2438.8, Fe 3008.1, Ti II 3242.0, Cu 3247.0, Li 2741.3 (Li lines too intense for measurement to include in two of the above plots).
FIGURE 34.—Time-intensity plots of Si, Fe, Ti II, Cu, and Li lines using deep-bore electrode No. 2 (bore depth 0.75 inch) containing different proportions of sample and admixtures. Direct-current arc of 10 amperes; controlled atmosphere, CO₂ with a flow of 20 cfh. Lines measured: Si 2438.8, Fe 3008.1, Ti II 3242.0, Cu 3247.0, Li 2741.3 (Li lines too intense for measurement to include in two of the above plots).
stabilizing the arc and cooling the sides of the electrodes, the exclusion of nitrogen from the arc plasma suppresses cyanogen band interference. The cyanogen region from approximately 3500 A to 4200 A contains numerous sensitive atom and ion lines that could be used for analytical purposes.

The jets used here and described below are different from the Stallwood design (1954) in that, for the work reported here, the gas is not directed as much against the electrode for cooling. They are not suggested as a substitute for the very promising Stallwood principle. The initial purpose of the jets used in this report was to eliminate the cyanogen band spectra in as simple a way as possible. Subsequent developments indicated, however, that the jet is indispensable for the long electrode.

Gas jets made for use with cupped electrodes and deep-bore electrodes are shown in figures 35-37. The orifice in each gas jet consists of a commercially available ceramic nozzle which is used in electric arc welding with inert atmospheres. The electrode-holding mechanism was designed and built by C. J. Massoni. The gas jet replaces the lower electrode holder of a standard open arc-spark stand. Gas enters the jet through the curved copper tubing. These gas jets are effective in eliminating cyanogen band interference. The gas flow is directed upward along the sides of the electrode and air is thereby excluded from the arc plasma. The use of these jets eliminates possible contamination and cleaning problems associated with enclosed gas chambers.

The selection of carbon dioxide as the controlled atmosphere for preliminary work was based on intensity and precision tests. Carbon dioxide afforded more intense atomic spectra than He–O\textsubscript{2} mixtures and more intense ionic spectra than air with the spectrographic conditions used at that time. Although the intensities of many lines may be enhanced more effectively by other ambient gases (Vallee and others, 1950; Ellenberg and Owen, 1951; Thiers, 1953), carbon dioxide appeared to be a good compromise for the diverse materials analyzed in geological work.

The most pronounced effect observed when using carbon dioxide in these tests was the even volatilization obtained from the long-bore electrodes. However, the presence of C\textsubscript{2} bands of the Swan system (Pearse and Gaydon, 1950, p. 75) portended difficulties in using the 4500–4700 A region for analyses. Consequently, tests were made with other gases to test their efficacy in rendering a spectrum having line intensities comparable to air or carbon dioxide without attendant C\textsubscript{2} band emission.
Preliminary tests upon silicate rock samples with admixtures of LiBO$_2$ and (or) graphite, using the deep-bore electrodes, demonstrated the necessity of O$_2$ for sample consumption. Subsequent tests with Ar+O$_2$ and He+O$_2$ were made (Shaw and others, 1958). The spectra in figure 38 show the relative line and band intensities when air, CO$_2$, He–O$_2$ and Ar–O$_2$ atmospheres were used. The Ar–O$_2$ mixtures seemed to serve the purpose most satisfactorily. The He–O$_2$ mixtures gave lowest background and line intensities, whereas CO$_2$ engendered the heaviest background with line intensities comparable to those from Ar–O$_2$ mixtures. The C$_2$ and CN band intensities decreased from CO$_2$ to Ar–O$_2$ atmospheres without a similar decrease in line intensities. Variations in the argon: oxygen ratio had little influence on line or background. As a result, the 14 cfh Ar+7 cfh O$_2$ mixture was selected for a more thorough study. Although this relatively high O$_2$ concentration, as compared to air, is high, the sample consumption was not too rapid. No sample loss was detected with silicate rocks. In addition, there could be a little variation of Ar:O$_2$ ratios without noticeably affecting line:background intensity ratios.

Moving-plate studies of different sample–LiBO$_2$–graphite mixtures were made with the deep-bore electrode using a gas flow of 14 cfh Ar+7 cfh O$_2$. Fractional distillation still persisted for extreme examples of the very volatile elements, such as arsenic and mercury, and refractory elements, such as zirconium and tungsten. The effect of increasing the gas flow so as to increase the cooling of the electrode was tested. Previously the total gas flow of 21 cfh was adequate to stabilize the arc, but under more extreme circumstances presented by arcing zirconium minerals, or alkali salts, volatilization differences arose. When gas flows of 21 cfh Ar–7 cfh O$_2$ and 24 cfh Ar–4 cfh O$_2$ were used, no improvement in volatilization was found and less arc stability resulted. The need for water cooling as described by Semenov (1956) was indicated. This was accomplished by replacing the ceramic nozzle of the gas jet (figs. 36, 37) by a brass water jacket (fig. 39) having eight holes for the passage of gas. The water jacket is lowered over the electrode after the electrode has been clamped in place.
Figure 35.—Gas jet for 0.25-inch-diameter cupped electrode shown in position on arc stand. A, Photograph showing electrode, which is clamped in a steel spring chuck by rotating the lever shown projecting to the left in the photograph; B, diagram showing jet parts in exploded view, a, ceramic nozzle.
Figure 36.—Photograph of gas jet for 0.125-inch-diameter deep-bore constant-feed electrode shown in position on arc stand. The knob to the left of the cylindrical support is for clamping the electrode in a steel spring chuck. The long insulated arm to the right is for feeding the electrode into the arc as it burns; upward motion of 1% inches is provided. The counter electrode is adjusted independently.
FIGURE 37.—Diagram of gas jet for 0.125-inch-diameter deep-bore constant-feed electrode shown in figure 36, mounted directly on the optical bench.  

a, Ceramic nozzle;  
b, clamp for 1/4-inch-diameter electrode;  
c, rack and pinion height adjustment for moving the electrode into the arc region;  
d, insulated handle actuating mechanism for tightening electrode in clamp b;  
e, flexible wire, electrical connection;  
f, copper gas inlet;  
g, insulated base.
Figure 38.—Spectra obtained from a G-1: Li$_2$B$_2$O$_7$: graphite mixture in a 1:4:6 proportion. Samples were arced in 1.5-inch deep-bore electrodes at 10 amperes direct current, using the gas jet of figures 36 and 37. The following gas compositions (cfh) were used:

<table>
<thead>
<tr>
<th>a.</th>
<th>Air 20 cfh</th>
<th>d.</th>
<th>14 Ar+7 O$_2$</th>
<th>g.</th>
<th>14 He+7 O$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>b.</td>
<td>CO$_2$ 20 cfh</td>
<td>e.</td>
<td>10 Ar+10 O$_2$</td>
<td>h.</td>
<td>10 He+10 O$_2$</td>
</tr>
<tr>
<td>c.</td>
<td>16 Ar+4 O$_2$</td>
<td>f.</td>
<td>16 He+4 O$_2$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
A test sample was prepared in which oxides of silicon, lead, and arsenic were added in minor quantities to a zirconium oxide base. This sample was used to demonstrate the advantage of the water-cooled jet in approaching the desired constant-feed effect (fig. 40).

EXCITATION

In addition to the conventional 10-ampere, resistance-controlled, d-c power source for the arc, a high-voltage a-c arc and a constant-current d-c arc (Green and Kyper, 1940) were tested. The a-c arc was not studied further when moving-plate results indicated volatilization behavior similar to the higher current d-c arc. Complete sample consumption with the a-c arc also required much longer exposures. It is believed, however, that the potentialities of this type of source have not been fully exploited.

Table 2 is a summary of the precision obtained with the two d-c arc controls, using two kinds of sample electrodes. Twenty replicate exposures were made with each electrode, using each power source at 10 amperes direct current. Spectrographic, photographic, and densitometric conditions are the same as given on p. 248. The coefficients of variation and standard deviations for Si, Al, and Ti in diabase W-1 (Fairbairn and others, 1951) were determined using germanium as an internal standard. A 2-percent concentration of Ge in LiBO₂ was prepared by fusing GeO₂ with Li₂CO₃ and H₃BO₃. (See footnote 1, p. 235.) No advantage in using either source with the deep-bore electrode was found. However, regardless of the power source used, greater precision for titanium was obtained with the deep-bore electrode than with the cupped electrode. Similar conclusions are indicated by matrix tests (table 3).

TESTS OF MATRIX INFLUENCE ON LINE INTENSITY

The tests to this point have been attempts to select the best of a few possible spectrographic variables that contribute toward the constant-feed principle. Qualitative studies of volatilization behavior and burning characteristics have been used for selection of the long-bore electrode, LiBO₂ buffer (Li₂B₂O₇ used later), graphite admixture, and controlled atmosphere. The tests in this section serve to demonstrate the extent to which these variables contribute in minimizing the ever-present matrix influences.
Figure 39.—Water-cooled gas jet used with deep-bore constant-feed electrode No. 3. A. Photograph showing ceramic cup of figure 36 replaced with a brass collar; B. diagrams of top and side view showing: a, cylindrical brass case; b, copper insert soldered to case; c, 0.125-inch copper tubing for water circulation.
A method for testing the effect of buffers in reducing matrix influences was described by Mohan and Fry (1958). In their work, matrices of ZrO₂ and NaCl, each containing 5 percent Ba, were used as test samples. These samples were mixed with various diluents which had been reported as effective buffers by different authors. The intensity ratio of Ba 3071.59 to a diluent line was obtained for each matrix. The ratio of these ratios then served as a measure of the matrix effect, the lower values indicating a smaller effect.

The matrix tests, using Ba 3071.59 as the indicator, were made with the cupped and deep-bore electrodes. Several different sets of sample admixtures and arc atmospheres were used. The most definitive result observed from these tests was the large reduction in matrix influence when lithium borate was admixed with samples arced in the cupped electrodes. The deep-bore electrodes showed relatively low matrix effect in all these tests.
CONTRIBUTIONS TO GEOCHEMISTRY

Table 2.—Precision test with d-c constant-

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration in W-1 (percent)</th>
<th>Line pair</th>
<th>1.5-inch deep-bore electrode</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Standard deviation</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Coefficient of variation</td>
</tr>
<tr>
<td>Si</td>
<td>24.4</td>
<td>Si 2987.6</td>
<td>Ge 3124.8</td>
<td>0.78</td>
</tr>
<tr>
<td>Al</td>
<td>8.3</td>
<td>Al 2588.0</td>
<td>Ge 3124.8</td>
<td>0.83</td>
</tr>
<tr>
<td>Ti</td>
<td>0.67</td>
<td>Ti II 3072.1</td>
<td>Ge 3124.8</td>
<td>0.037</td>
</tr>
</tbody>
</table>

1 Fairbairn and others, 1951.
2 Standard deviation, \( \sigma \), determined as

\[
\sigma = \sqrt{\frac{\Sigma d^2}{n}}
\]

where \( d \) = deviation from chemical value, \( n \) = number of determinations.

A variation of the above tests was sought that would be more representative of the analytical problems at hand. It was also hoped that greater contrast in the properties of the two electrodes could be demonstrated. For this purpose NaCl and ZrO₂ matrices were each prepared to contain 5 percent W-1, the diabase standard (Fairbairn and others, 1951).

Six elements commonly determined in rock analyses were studied. No internal standard line was used. The sample:graphite ratio was 1:1 and a sample:Li₂B₄O₇:graphite proportion of 1:2:3 was used. Relative intensities of the following lines were measured:

<table>
<thead>
<tr>
<th>Ionization potential, ev</th>
<th>Excitation potential, ev</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si 2524.1</td>
<td>0.01–4.90</td>
</tr>
<tr>
<td>Al 2575.1</td>
<td>0.01–4.81</td>
</tr>
<tr>
<td>Fe 3059.1</td>
<td>0.01–4.09</td>
</tr>
<tr>
<td>Fe II 2598.4</td>
<td>0.05–4.80</td>
</tr>
<tr>
<td>Mg 2779.8</td>
<td>7.86</td>
</tr>
<tr>
<td>Ti II 3075.2</td>
<td>2.70–7.14</td>
</tr>
<tr>
<td>Ca II 3179.3</td>
<td>6.81</td>
</tr>
<tr>
<td></td>
<td>0.01–4.02</td>
</tr>
<tr>
<td></td>
<td>3.14–7.02</td>
</tr>
</tbody>
</table>


SPECTROGRAPHIC CONDITIONS AND APPARATUS

Spectograph: Eagle-mounting, 15,000-line grating, 25 μ slits, reciprocal linear dispersion in first order=5.6 A per mm. Range used, 2460–3850 A, first order. Anode sample electrodes: Pure graphite, 0.125-inch-diameter electrode with 1.5-inch deep bore; and 0.25-inch cupped electrode.
A CONSTANT-FEED DIRECT-CURRENT ARC

current and d-c constant voltage power source

1:4:5. 20 replicate determinations

<table>
<thead>
<tr>
<th>1.5-inch deep-bore electrode—Con.</th>
<th>0.25-inch-diameter cupped electrode</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Constant voltage</strong></td>
<td><strong>Constant current</strong></td>
</tr>
<tr>
<td>Standard deviation  (^\text{a})</td>
<td>Coefficient of variation  (^\text{a}) (percent)</td>
</tr>
<tr>
<td>.62</td>
<td>2.53</td>
</tr>
<tr>
<td>.97</td>
<td>11.6</td>
</tr>
<tr>
<td>.042</td>
<td>6.2</td>
</tr>
</tbody>
</table>

\(^{a}\) Coefficient of variation, \(v\), determined as

\[ v = \frac{100}{C} \sigma, \]

where \(C\) = chemical concentration.

**Table 3.**—Matrix effects on Si, Al, Fe, Mg, Ti, and Ca in NaCl and \(\text{ZrO}_2\) matrices.

Percent deviation from unity for 9 replicate determinations, \(\Delta\)

(The line intensity ratios were determined after burning samples in a 10 amp. d-c arc)

<table>
<thead>
<tr>
<th>Lines measured</th>
<th>1.5-inch deep-bore electrodes, water cooled, 14 cfm (\text{Ar}) + 7 cfm (\text{O}_2)</th>
<th>0.25-inch cupped electrodes, 14 cfm (\text{Ar}) + 7 cfm (\text{O}_2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample: graphite = 1:1</td>
<td>Sample: (\text{Li}_2\text{BO}_3): graphite = 1:3</td>
<td>Sample: graphite = 1:1</td>
</tr>
<tr>
<td>Si 2524.1</td>
<td>13</td>
<td>33</td>
</tr>
<tr>
<td>Al 2575.1</td>
<td>32</td>
<td>19</td>
</tr>
<tr>
<td>Mg 2779.8</td>
<td>19</td>
<td>19</td>
</tr>
<tr>
<td>Fe 3059.1</td>
<td>20</td>
<td>13</td>
</tr>
<tr>
<td>Fe II 2598.4</td>
<td>23</td>
<td>27</td>
</tr>
<tr>
<td>Ti II 3075.2</td>
<td>30</td>
<td>16</td>
</tr>
<tr>
<td>Ca II 3179.3</td>
<td>55</td>
<td>24</td>
</tr>
</tbody>
</table>

Upper cathode: Pure graphite, 0.25 inch, 35° tip with a hemispherical 0.05-inch radius.


Controlled atmosphere: 14 cfm \(\text{Ar}\) + 7 cfm \(\text{O}_2\), using gas jet (fig. 35) for cupped electrode and water-cooled gas jet (fig. 39) for deep-bore electrode.

Exposure: Step sector having 25, 12.5, 6.25, and 3.13 percent transmission. Source focused on grating using 25 mm diameter spherical lens, focal length = 100 mm. Upper and lower edges of grating masked for 50 percent of grating. Electrode images masked off grating.

Processing: SA-1 plates developed in D-19 at 18°C for 5 minutes, fixed for 5 minutes, and air dried. Rocking-tray developing and fixing.
Sample preparation: Sample-Li₂B₄O₇-graphite mixture ground in boron carbide mortar until homogeneous (about 5 minutes).

Electrode loading: Sample mixture of 44 mg loaded in deep-bore electrodes with the aid of a polyethylene funnel and a 0.040-inch tungsten carbide drill rod, used for tamping. The cupped electrode was loaded with 22 mg of sample mixture, using stainless steel funnel, and tamped with a graphite rod. A layer of powdered graphite was added on top of samples in all electrodes in order to minimize sample losses when igniting the arc.

Densitometry: Calibration curve is obtained by plotting galvanometer deflections as ordinate against relative log intensity of step-sectored spectrum, having an integral intensity ratio of 2.0, as abscissa.

RESULTS

The spectra of NaCl and ZrO₂ matrices were exposed alternately on each plate, and a given line intensity ratio was calculated from successive pairs of spectra. The average percent deviation from unity (\(\Delta\)) of nine ratio determinations are listed in table 3.

The data in table 3 clearly demonstrate the relative advantages of the deep-bore electrode for reducing matrix effects. The addition of lithium borate to the sample mixture appears superfluous with this electrode. Ion lines, which are more sensitive to excitation variations, react similarly to atom lines in either NaCl or ZrO₂ matrices when the deep-bore electrode is used. On the other hand, a large effect is noted when the more conventional cupped electrode is used, especially without the addition of Li₂B₄O₇. In the precision tests, reported in table 2, a parallel comparison between atom and ion lines in the two types of electrodes is found. Even the larger proportion of buffer and graphite to sample used in the precision test failed to contribute sufficient uniformity of excitation for sample mixtures arced in the cupped electrodes.

REFERENCES


