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**Research and Development Technical Report** 

DELET-TR-78-0563-2

## LITHIUM - THIONYL CHLORIDE BATTERY

A.N. DEY W. BOWDEN J. MILLER P. WITALIS

P.R. MALLORY & CO. INC. LABORATORY FOR PHYSICAL SCIENCE BURLINGTON, MA 01803



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designs on the current carrying capability of the spirally wound D cells and the flat cylindrical cells and found that short circuit current densities of 200mA/cm<sup>2</sup> may be realized by proper choice of the current collector design using a conventional cathode matrix. In addition we carried out a parallel research effort in order to gain increased understanding of the SOCl<sub>2</sub> reduction process. We showed, using cyclic voltammetry and coulometry, that the reduction of SOCl<sub>2</sub> does not lead to the immediate formation of S and SO<sub>2</sub> but to some other intermediates which slowly decompose to form SOCl<sub>2</sub>.

During the second quarter, we tested high rate spirally wound D cells of various promising cathode designs on BA5590 and GLLD laser designator duty cycles. We found that  $\text{Li/SOCl}_2$  BA5590 battery ran for 100 hrs at room temperature on the BA5590 duty cycle and weighed approximately 840 gm. This represents a 20% gain in service life and an 11% reduction in battery weight over that of the  $\text{Li/SO}_2$  batteries.

Li/SOCl<sub>2</sub> D cells when tested on pro-rated bases on the GLLD laser designator duty cycle delivered approximately six times the service life of the presently used Ni/Cd batteries.

We have obtained some spectroscopic evidence in favor of the formation of at least two possible meta-stable intermediates resulting from the discharge of  $SOCl_2$  at room temperature. These results have an important bearing on the devising of ways to improve the safety of the cells.

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The Li/SOCl<sub>2</sub> inorganic electrolyte system (1-4), is the highest energy density system known to date. It consists of a Li anode, a carbon cathode and SOCl<sub>2</sub> which acts both as a solvent and as a cathode active material. The electrolyte salt that has been used most extensively is LiAlCl<sub>4</sub>, but salts such as Li<sub>2</sub>B<sub>10</sub>Cl<sub>10</sub> (5) and Li<sub>2</sub>O(AlCl<sub>3</sub>)<sub>2</sub> (6) have also been used successfully in this system for improving the shelf life characteristics.

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This report describes

The main objective of this program is to develop high rate Li/SOCI<sub>2</sub> cells and batteries for portable applications of the U. S. Army. The cells and batteries must deliver higher energy densities than are presently available and must be safe to handle under U. S. Army field conditions.

We carried out a detailed development (7) on the spirally-wound high rate D cells in order to establish their performance capabilities as well as to identify and correct the limitations in their performance and safety under various use and abuse conditions. Substantial progress was made to correct the cell limitations. We found that the state-of-the-art spirally-wound D cells approach the high rate requirements of the various U. S. Army applications more closely than do any other cell designs at the present time. Accordingly we have used this spirally-wound D cell as a starting point and have improved its rate capability to meet the requirements of two specific applications, viz. the BA5590 Battery for Man Pack Radio and the Battery for the GLLD Laser Designator. Our work during the second quarter was focussed primarily on the testing of D cells for the above two applications.

A parallel research effort was continued in order to gain an increased understanding of the cell discharge reactions, particularly with respect to the unstable intermediates. We believe that this knowledge will be useful in providing guidance for improving the safety of the system. The progress in optimizing the high rate Li/SOCl<sub>2</sub> D cells for the BA5590 battery and the GLLD battery are described in this second quarterly report. In addition we have carried out electrochemical and spectroscopic experiments to clarify the mechanism of thionyl chloride reduction and subsequent chemical reactions in the cell. These results are also included in this report.

#### II. Spirally Wound D Cell

We found (21) that the current distribution of the high rate spirally wound D cells having one current collector tab located at one end of the cathode was controlled by the electrical conductivity of the cathode grid. Therefore, we examined the effect of a variety of cathode tab designs on the current carrying capability of the cathode as determined by the short circuit current density of the cathode. The results obtained during the preceding quarter are reproduced in Table 1. Note that by placing the tab in the middle as in design 2, the short circuit current density was increased from 53 mA/cm<sup>2</sup> to 168 mA/cm<sup>2</sup>. In view of the simplicity of this cathode design and the substantial improvement in performance, we chose this cathode design for the spirally wound D cells to be tested for BA5590 manpack radio battery application. In addition, we evaluated spirally wound D cells having the various cathode designs shown in Table 1 for the GLLD Laser Designator application as well. The experimental details and the results obtained to date are presented here.

#### A. BA5590 Manpack Radio Battery

The specifications of the above battery are as follows.

<u>Dimension</u>: 4.4" x 2.45" x 5.00" <u>Voltage:</u> Nominal 12 or 24V Maximum (OCV) 15 or 30V Average 12.5 or 25V End Voltage 10 or 20V Capacity at 70°F 10 A.Hr (30 hr rate) Max. Rate: 2 hr rate

Duty for 30 volt Operation: 8 ~ load for 100 msec followed by 39 ~ load for 1 mm. followed by 560 ~ load for 9 min. The above is repeated. The presently used batteries contain 10  $\text{Li/SO}_2$  D cells in series and parallel to meet the 15 and 30 volt requirements. In view of the higher OCV of the  $\text{Li/SOCl}_2$  D cells (3.6 volt) only eight D cells are needed for the required voltage.

We carried out the initial evaluations using single D cells instead of the 8 cell battery. Therefore, the duty cycle used was prorated for a single cell as:

l\_ load for 100 msec followed by  $5_{1}$  load for 1 min. followed by 70\_ load for 9 min. The cycle is repeated.

One  $\text{Li/SOCl}_2$  D cell with cathode having one tab in the middle (design 2 of Table 1) was tested under the above duty cycle at 25°C. The cell voltage under the 80 L and 5 L loads are plotted as a function of time and is shown in Fig. 1. The cell ran for 95 hours, delivering a capacity of approximately 10 A.Hr.

Another D cell with similar cathode was tested at  $0^{\circ}$ C under the same duty cycle. In this case the cell voltages under all the three loads were monitored as a function of time and are shown in Fig. 2. This cell ran for 93 hours to 2.5 volt cut off (on 1  $\triangle$  load) corresponding to a cell capacity of approximately 9.8 A.hr.

One BA5590 battery was assembled using 8  $\text{Li/SOCl}_2$  D cells in series. The battery was tested at room temperature on the battery duty cycle mentioned earlier. The battery voltage on the three loads and the battery temperature was monitored as a function of time. The results are shown in Fig. 3. The average battery voltages on the 8  $\pounds$ , 39  $\pounds$  and 560  $\pounds$  loads were 25.5, 27 and 28.5 volts respectively. The battery ran for approximately 100 hours to the 20 volt cutoff. The capacity delivered by the battery was approximately 11 A. Hr. The temperature of the battery remained unchanged during the test.

The above BA5590 battery weighed 840 gms and contained 8 D Li/SOC12 cells. The  $\text{Li/SO}_2$  BA5590 batteries, on the other hand, contain 10 D cells and weigh 940 gms with a service life of 80 hours on the same duty cycle.

Thus, the improvement in performance of the  $\text{Li/SOCl}_2$  BA5590 battery over that of the  $\text{Li/SO}_2$  BA5590 battery was significant at the temperatures of these tests.

B. GLLD Laser Designator Battery

The specifications for the above battery are as follows:

Dimension:	2.82" x 3.75" x 9.30"
Voltage:	Nominal 24V
	Maximum (OCV) 32V
	Average 24V
	End voltage 20V
Duty:	17.5A for 35.5 m.sec. followed by 1.8A for 14.5 m.sec.
	and the cycle continues for 3 minutes. This constitutes
	one burst. This 3 minute cycle occurs every 30 minutes.

The presently used Ni/Cd batteries provide 3 bursts per charge.

The voltage requirement of 32 volt necessitates the use of at least 8  $Li/SOCl_2$  D cells in series. The available battery volume can be occupied by 16 D cells. Therefore, the battery will consist of two D cells in parallel and 8 D cells in series, and a single D cell will experience half the specified constant current loads.

#### 1. Testing of D Cells on Modified Duty Cycle:

We tested the Li/SOCl<sub>2</sub> D cells on a modified duty cycle involving an average current of 6.5A (13 A. for two cells in parallel) or 8A (in some cases) for 3 minutes every 27 minutes. This modified test consumes the same cell capacity per 3 minute burst as the above specified duty cycle. We used the modified duty cycle in order to expedite our cell development program since the test circuit for the specified duty cycle was still under development at this stage of the program. These tests were carried out to demonstrate the efficacy of the various current collector designs for the GLLD laser designator application.

One Li/SOCl<sub>2</sub> D cell with a cathode of design 2 (Table 1) having the tab located at the center of the 20 inch long electrode was tested under the modified GLLD duty cycle using a 6.5A current at room temperature. The voltage and the temperature of the cell were monitored during the three minute burst and are plotted as a function of burst number in Fig. 4. During each three minute 6.5A burst the cell voltage showed a slight recovery during the early stages of the discharge and a slight decline towards the end of the discharge. The cell temperature showed a sharp rise during the three minute burst and declined to almost the original level during the 27 minute open circuit stand. The cell delivered 16 bursts above 2.5 volt corresponding to a capacity of 5.2 A.hr. The cell temperature during the burst periods reached a maximum of 30°C which is well within the safe limits of temperature excursion. This cell was not insulated with glass filter paper.

A second D cell of similar construction was tested under the modified duty cycle using 8A and the cell was thermally insulated using glass filter paper. The test results are shown in Fig. 5. Note that the cell temperature rose to a maximum of 40°C towards the end of the discharge during the three minute 8A bursts. The cell delivered 12 bursts corresponding to a capacity of 4.8 A.Hr.

Another D cell of the same design was tested on the modified duty cycle using 6.5A bursts at  $-30^{\circ}$ C. The results are shown in Fig. 6. Although the cell delivered 12 bursts, the cell voltage remained around 2.0 volt during most of the bursts.

One D cell with a 20 inch long cathode having both a horizontal and a vertical tab (design 4 of Table 1) was tested on the modified duty cycle using 6.5A bursts at room temperature. This cell was also thermally insulated with glass filter paper separator. The results are shown in Fig. 7. The cell delivered 10 bursts corresponding to a cell capacity of only 3.25 A. Hr. This low cell capacity is most likely due to the lower effective cathode area because of the presence of the horizontal tab.

One D cell with a 25 inch long cathode having one tab at one end (design 6 of Table 1) was tested on the modified duty cycle using 8A bursts at room temperature. The results are shown in Fig. 8. The cell showed unusually high polarization as well as a sharp rise in the cell temperature during the 3 minute bursts. The cell temperature reached a maximum of 55°C during the last burst. The cell only delivered 4 bursts above 2.0 volt. This poor performance may be attributed to the non-uniform current density distribution as a result of the poor current collection.

Two D cells with the 25 inch long cathode having both a horizontal and a vertical tab as in design 7 of Table 1 were tested on the modified duty cycle using 6.5A bursts at room temperature. The results from these two cells are shown in Fig. 9 and 10 respectively. The cells delivered 10 and 12 bursts corresponding to capacities of 3.25 and 3.9 A. Hr respectively. The maximum cell temperature remained below  $40^{\circ}C$ .

Another D cell of the same design was tested using 8A bursts at room temperature. The results are shown in Fig. 11. The cell delivered only 7 bursts corresponding to a capacity of only 2.8 A. Hr. Although the use of the horizontal tab leads to some improvement over the single vertical tab the improvements are only moderate because of the lowering of the active area by the horizontal tab.

Two D cells were made with 25 inch cathodes having two vertical tabs located equidistant from the cathode ends and the middle. One cell was tested on the modified duty cycle using 6.5A and the other one was tested using 8A at room temperature. The results are shown in Fig. 12 and 13 respectively. The cells delivered 16 bursts at 6.5A and 12 bursts at 8A corresponding to cell capacities of 5.2 and 4.8A. Hr respectively. This represents a significant improvement over the cells of the previous design.

Several D cells were made with 25 inch long cathodes having three tabs as in design 8 of Table 1. One cell was tested at room temperature on the

modified duty cycle using 6.5A. The results are shown in Fig. 14. The average cell voltage on load remained only slightly below 3 volts and was found to be rather stable. The cell temperature rose to a maximum of only 36°C during the 6.5A burst. The cell delivered 16 bursts above 2.5 volt corresponding to the cell capacity of 5.2 A.Hr.

Two D cells of the same design were tested at  $-30^{\circ}$ C on the modified duty cycle using 6.5A current. The results are shown in Fig. 15 and 16 respectively. Note, that the average cell voltage on load was above 2.0 volt. The two cells delivered 10 and 8 bursts corresponding to the cell capacities of 3.25 and 2.6 A. Hr respectively. The performance of these cells was somewhat better than the cells of design 2 at  $-30^{\circ}$ C.

Another D cell of the same design was tested at room temperature on a modified duty cycle using 8.75A. The results are shown in Fig. 17. The cell delivered 11 bursts and went into reversal during the 12th burst. The capacity delivered was 4.8 A.Hr.

This concludes all the testing of D cells on the modified duty cycle. The results (summarized in Table 2), indicate that the cells with 20 inch long cathodes having a tab in the middle provide adequate performance, it is also the simplest one to fabricate. The cells with 25 inch long cathodes with two or three tabs also performed satisfactorily. The use of horizontal tabs is not an efficacious and will not be considered for any further tests.

#### 2. <u>Testing of D Cells on Regular Duty Cycle</u>

We designed and built test equipment capable of testing 8 batteries at one time on the GLLD Laser Designator duty cycle specified earlier. The test equipment was designed to draw constant currents of 17.5A and 1.8A for the desired time intervals from the battery down to a voltage of approximately 0.8 volt at which point the test is automatically terminated thus preventing any force-discharge on the cells. Since the GLLD battery will contain two D cells in parallel, we carried out the initial tests using two D cells in parallel instead of using the full 16 cell battery.

The D cells with 20 inch long cathodes having one tab in the middle (design 2) were tested at room temperature using the regular duty cycle. The results are shown in Fig. 18. The cell voltages on both the 17.5A and 1.8A loads are shown as a function of number of bursts. The two D cells delivered 9 bursts above 2.0 volts. On the 10th burst, the cell voltage on 17.5A dipped to 1.0 volt at which point the cell temperature rose sharply to 50°C. The test results of a second pair of D cells of the same design are shown in Fig. 19. In this case, the cells delivered 12 bursts. These results are approximately similar to the previous test results on the modified duty cycle.

The D cells with 25 inch long cathodes having three tabs (design 8) were tested at room temperature on the regular duty cycle. The results are shown in Fig. 20. The voltages on the first two bursts were not measured because of recorder malfunction. Note that the cell voltage on 17.5A was substantially above 2.5 volt and that the voltage remained quite stable during the 3 minute bursts. The two cells (in parallel) delivered 17 bursts corresponding to a total capacity of 11 A. Hr from the two D cells or 5.5 A. Hr/cell.

The above tests were repeated with another pair of the D cells of the same design. The results are shown in Fig. 21. In this test, the individual cells were covered with a heat shrinkable plastic jacket. The cell wall temperature was monitored during the test and it rost to a maximum of 40°C towards the end of the test. The cells delivered 18 bursts above 2.0 volt corresponding to a capacity of approximately 5.85 A. Hr/cell. This is approximately 50% of the total available cell capacity of 12 A. Hr. These results are extremely encouraging particularly in view of the fact that the presently used Ni/Cd batteries deliver only 3 bursts per charge.

#### C. <u>Conclusions</u>

The rate capability of the Li/SOCl<sub>2</sub> D cells was increased substantially by improving the cathode designs. The cells with the improved cathodes were used to fabricate a BA5590 manpack radio battery containing 8 cells in series. The battery had a service life of 100 hrs at room temperature. This represents a 20% increase in service life with an associated weight saving of 11% relative to the presently used  $\text{Li/SO}_2$  BA5590 batteries which use 10 D cells.

The GLLD laser designator battery can accommodate 16 D cells which will be connected 8 in series and two in parallel.  $\text{Li/SOCl}_2$  D cells of various cathode designs were tested on a prorated basis using single cell and two cells in parallel. The results showed that these batteries are capable of providing six times longer service life than do the currently used Ni/Cd batteries. The temperature of a single cell remained below 40°C during these high current tests.

#### III. Electrochemical and Spectroscopic Studies on SOGI<sub>2</sub> Reduction

Calorimetric (12) and DTA studies (7, 11) of  $\text{Li}/\text{SOCl}_2$  D cells showed that chemical reactions continue to occur liberating heat after the discharge of the cell. Since these reactions do not occur in an undischarged cell, it is reasonable to assume that the reduction products of  $\text{SOCl}_2$  are responsible, at least in part, for these spontaneous exothermal reactions. The reported spontaneous explosions of partially discharged  $\text{Li}/\text{SOCl}_2$  cell on casual storage may also be initiated by the above unknown reactions involving the  $\text{SOCl}_2$  reduction intermediates. Thus, knowledge regarding the nature of the unstable intermediates of  $\text{SOCl}_2$  reduction may be useful as a guide for the improvement of the safety of the  $\text{Li}/\text{SOCl}_2$  cells. We postulated (9, 13) the overall cell reaction to be

 $SOCI_2 + 2Li \longrightarrow SO + 2LiCI [1]$ 

where SO may dimerize and then decompose to S and SO2 or may form polymers. There is substantial evidence (5, 14) in favor of cell stoichiometry [1]. The quantitative formation of LiCl and qualitative formation of S and SO2 are known. Very little has been known regarding intermediate species formed during the reduction of SOCl2. Attempts to use cyclic voltammetry (15, 16) in neat SOCl2- $LiAlCl_4$  solutions to study the discharge reaction were complicated by electrode passivation due to the precipitation of LiCl, insoluble in SOCl2. We have circumvented this electrode passivation problem by using a supporting electrolyte consisting of tetrabutylammonium hexafluorophosphate ( $N(C_4H_9)_4 PF_6$ ) in organic solvents such as dimethyl formamide (DMF) acetonitrile (AN), methylene chloride and dimethyl sulfoxide, for studying the SOCI2 reduction. The electrode passivation is absent in the above solutions since both tetrabutylammonium chloride and S are soluble in the solvents studied. We carried out both cyclic voltammetry and coulometry in the above electrolytes and found that neither S nor SO2 are formed as immediate reduction products. There exists at least one surprisingly long lived intermediate species which slowly decomposes to regenerate SOC12. Our previous experiments have demonstrated a stoichiometry of 2.0 for the reduction of SOCl2.

We have examined the formation of this intermediate species and its reactions by a number of techniques, including high sweep rate cyclic voltammetry, UV-VIS spectroscopy and low temperature electrochemical studies. The results are reported below.

#### A. Experimental

Electrochemical experiments were performed using a PAR 173 potentiostat and PAR 175 function generator with associated ancillary equipment. Pulse polarograms were performed with a PAR 174A polarographic analyzer. The data were collected using conventional x-y and strip chart recorders. The data for fast sweep experiments were taken on a Tektronix model 5115 storage oscilloscope or a PAR 4101 scan recorder. Platinum working electrodes were pretreated by chromic acid followed by a wash with distilled water and air drying. Experimental solutions were routinely degassed with argon before the substrates were added. A silver chloride coated silver wire was used as a reference electrode. This electrode has a potential of 3.30V vs. Li in  $PF_6^-$ -DMF solutions. A Tl·Hg/ TICl<sub>3</sub> reference electrode was used for the low temperature experiments in DMF.

The experiments were performed in cells of conventional design. The coulometric experiments were carried out in a two compartment H-cell using a glass frit to prevent passage of material from the working and auxiliary electrode compartments. The low temperature experiments were carried out using the jacketed cell in Figure 22.

The organic solvents were either Burdick & Jackson "Distilled in Glass" DMF, and CH<sub>2</sub>Cl<sub>2</sub> or Eastman Spectro Grade acetonitrile or Aldrich "Gold Seal" Dimethyl Sulfoxide. The supporting electrolyte was prepared by metathesis of tetrabutylammonium chloride (TBACl) and lithium hexafluorophosphate in acetone/ water and purified by multiple recrystallizations in hot ethanol.

#### B. <u>Results</u>

#### 1. Cyclic Voltammetry

In addition to the solvents described earlier we have utilized dimethylsulfoxide (DMSO) because of its great ability to coordinate with and

thus stabilize highly reactive molecules.

A series of voltammograms for ~10 mM SOCl<sub>2</sub> in  $CH_3CN/0.1$  N TBAPF<sub>6</sub> are shown in Fig. 23. In these voltammograms on a constant current scale of 25µA/cm, the sweep rate, V, was varied from 0.20V/sec to 2.0V/sec. As the sweep rate increases, the peak current increases, but an oxidation process near 0.0V appears at 500 mV/sec and becomes more prominent at 1.0 and 2.0V/sec. The chemical entity which is being oxidized at this potential is not present in the solution originally since there is no anodic current at this potential until some of the thionyl chloride has been reduced. The sweep rate dependence of this oxidation wave shows that the species responsible for the oxidation is reacting fairly rapidly in solution. The oxidation and coupled reduction near +1.0V is due to oxidation of chloride ion to chlorine. The chloride itself is generated by reduction of thionyl chloride.

The effect of concentration on the  $SOCl_2$  reduction has been examined. Cyclic voltammograms of  $SOCl_2$  at various concentrations at a Pt electrode are shown in Fig. 24. For a given sweep rate of 500 mV/sec the oxidation near 0.0V becomes less prominent as concentration increases. From the sweep rate dependence of this oxidation, we can conclude that the species responsible for this oxidation wave is reacting with thionyl chloride in solution or with itself. Change in the reduction behavior was also noted as the wave near -0.65V became less prominent and the one near -1.0V more prominent as the concentration increased.

We have also examined the effect of substrate changes on the thionyl chloride reduction. Ordinary graphite or porous carbon electrodes displayed too high a background current to be usable in these experiments. We have, however, examined the use of nickel, indium, gold, amalgamated gold, copper and silver as electrode materials for observing the SOCl<sub>2</sub> reduction. Cyclic voltammograms on nickel, iridium and gold electrodes were quite similar to those observed with Pt as shown in Fig. 25. The copper electrode showed a very limited area of usable voltage while the results with silver electrode are anomalous and reproduced in Fig. 26.

We have continued our work on the thionyl chloride reduction in DMF (and in AN) and the reduction appears to take place by the same mechanism. The better separation of the two reduction waves in DMF makes it a more attractive solvent for use in strictly electrochemical studies on this system. Experiments in which sweep rate and concentration were systematically varied in DMF/TBAPF<sub>6</sub> gave similar results for the reduction of SOCl<sub>2</sub> to those found in acetonitrile. The second reduction wave remained more prominent in DMF, suggesting that coordination with DMF was stabilizing the species responsible for the reduction process.

To examine the formation of the intermediate species more closely we also examined the reduction at higher sweep rates than usual, recording the current/voltage trace on a storage oscilloscope. At sweep rates as high as 100V/sec no qualitative change in the voltammogram could be observed. Except for uncompensated iR losses the voltammogram was similar to those generated at lower sweep rates. We were also able to use the storage oscilloscope with multiple sweeps of the same voltage range to examine the relationships of the two reduction waves and the oxidation near 0V. Using this method we were able to generate a "steady state" voltammogram of reduced thionyl chloride species. This voltammogram was quite unique (not shown because of the absence of photographic attachment) in that the first thionyl chloride reduction wave was absent while the second reduction wave and the oxidation near OV were still present. This result indicates that the oxidation near 0.0+ does not regenerate thionyl chloride since the first reduction wave corresponding to the reduction of SCCl<sub>2</sub> is absent and as such the reduction of thionyl chloride is irreversible.

For further comparison we ran cyclic voltammograms in dimethylsulfoxide (DMSC), a solvent which is regarded as very good for reduced species because of its ability to stabilize reactive materials by complexation. DMSC is also known to have a rather low activity of residual water because of the sulfoxide's coordinating ability. A cyclic voltammogram for SOCl<sub>2</sub> in DMSO/TBAPF<sub>6</sub> is shown in Fig. 27. The pattern of two successive reduction waves and an oxidation near 0.0V are very similar to that observed in DMF. The oxidation near 0.0V is quite prominent even at 0.05V/sec sweep rate, indicating that the reaction species involved is greatly stabilized by the DMSO solvent. The definition of the two reduction waves is very clear and there is little evidence for the appearance of the third wave near -1.0V which we noted in other solvents.

#### 2. Coulometry

We have examined the coulometric reduction of  $SOCI_2$  in DMF, acetonitrile, and methylene chloride. In all three solvents we have observed the regeneration of thionyl chloride by destruction of an intermediate species and have confirmed the Faradaic current of 2 equivalents of charge for each equivalent of SOCl<sub>2</sub> reduced. In particular, the reduction in acetonitrile is of interest since the UV-VIS solvent cutoff of 190 nm gives a much better spectral window than DMSO (268 nm), DMF (270 nm), or methylene chloride (230 nm). We were able to monitor the course of a coulometric reduction by using both cyclic voltammetry and UV-VIS spectroscopy to monitor changes in the solution. For comparative purposes, spectra of SC2 and S in acetonitrile are reproduced in Fig. 28 and 29, respectively. The wave lengths of maximum absorbance, 279 nm for SC2 and 277 for S are very similar to that of SCC12, 277 nm as shown in Fig. 30, however, there is a second band in the spectrum near 234 nm which is not present in the spectra of SO2 and S. In this particular experiment  $10\mu l$  of SOCl\_2 was used in 75 ml of  $\rm CH_3CN$  to give a concentration of 1.83 mM  $SOCl_2$  in  $CH_3CN$ . This concentration was experimentally determined to be usable for both cyclic voltammetry and UV-VIS spectroscopy. Changes in the cyclic voltammograms and spectra as a function of charge passed are displayed in Figs. 31 and 32. As the electrolysis proceeds, the absorbance maximum in the UV-VIS spectrum shifts from 277 to 291 nm corresponding to a lower energy. The absorbance also increases gradually with the electrolysis and reaches a maximum of 1.94 from 0.63. The absorbance then declines drastically to 0.79 at n = 1.76(Fig. 33). The absorption peak at 290 nm disappears on storing the electrolyzed

solution (n = 2) for one week at room temperature. At the same time, the first reduction wave in the cyclic voltammogram is destroyed and the second reduction wave becomes much more prominent. The UV-VIS spectra of the solutions after the electrolysis did not show the presence of S or SO<sub>2</sub>. When the solution was allowed to stand overnight the first wave became some. at more noticeable and some poorly defined waves at more negative potentials also appeared. The final voltammograms showed two very poorly defined reduction waves with no SOCl<sub>2</sub> present, while the final spectra showed a substantial decline in absorbance.

#### C. Discussion

The coulometric data demonstrate an overall n value of 2.0 for the reduction of thionyl chloride. Reductions for extended times have given green, air-sensitive solutions possibly including polysulfides, but there is no evidence to suggest that this reflects any process occurring in SOCl<sub>2</sub> reduction since the cyclic voltammograms show that no further unreduced SOCl<sub>2</sub> is present.

The spectroscopic evidence shows that S and  $SO_2$  are not formed during the reduction process itself. Instead a different species is formed with an absorption band near 291 nm. This is confirmed by the cyclic voltammograms which show only a slow formation of species reducing at potentials near those for S and  $SO_2$ . This intermediate species is reasonably stable and only slowly decomposes to generate more  $SOCl_2$ .

We can now present a reasonable reduction mechanism for  $SCCl_2$  which is supported by data at each stage. In an earlier report we suggested that the regeneration of  $SOCl_2$  was due to destruction of a dimer formed by the coupling of one electron reduced species. [2]

$$2SOC1_2 + 2e^{-} \longrightarrow 2SOC1 + 2C1^{-}$$

$$2SOC1 \longrightarrow (SOC1)_2 \longrightarrow SO + SOC1_2 [2]$$

Our results from the high speed experiments make this seem less likely, since we were unable to outrun the dimerization process and observe two successive one electron reductions of  $SOCl_2$ . The other mechanism we suggest now

seems the more likely of the two. In the initial reduction process, which gives the large waves in DMF and DMSC, there are two successive one electron processes to generate SC by a net two electron transfer [3].

$$SOCl_2 + e^- \longrightarrow SCCl + Cl^- + e^- \longrightarrow SC + 2 Cl^-$$
 [3]

For reactions such as this, it is not unusual for  $E_0$  for the second reduction process to be positive of  $E_0$  of the first. Early in the reduction process  $SCCl_2$  is present in large concentration compared to SC and reaction with  $SCCl_2$  [4] is favored.

$$SC + SOCl_2 \longrightarrow SO \cdot SCCl_2$$
 [4]

This species can slowly dissociate to regenerate  $SCC1_2$ . As the electrolysis progresses the concentration of SC increases compared to  $SCC1_2$  as reaction with SC is favored [5].

$$SC + SC \longrightarrow (SC)_n$$

The product  $(SC)_n$  is characterized by the absorbance near 291 nm and is reduced near -0.65V and the reduced product oxidized near +0.3V while the  $SC \cdot SCCl_2$ product is oxidized near 0.0V as shown by the potential shift in this wave as the electrolysis continues. This species  $(SC)_n$  appears to be reasonably inert kinetically and only slowly decomposes to S and  $SC_2$ , perhaps by loss of  $SC_2$ to form successively oxygen deficient polymers. Cbservations of delayed pressure rise (19) and thermal activity (12) in discharged thionyl chloride cells are quite consistent with this hypothesis.

#### D. <u>Conclusion</u>:

Our data strongly suggest that there are at least two intermediate species formed in the reduction of  $SOCl_2$  to LiCl, S and  $SC_2$ . The decomposition of these species  $SC \cdot SCCl_2$  and  $(SC)_n$  are probably responsible for the delayed  $SC_2$  generation and thermal activity noted in discharged  $SCcl_2$  cells.

#### IV. Conclusions and Future Work

The high rate spirally wound Li/SOCl<sub>2</sub> D cells showed considerable promise in both the BA5590 and the GLLD laser designator battery applications. Accordingly, we concentrated our effort on this area during the last two quarters. In view of the encouraging results obtained to date with the D cells we see no additional advantage in developing the 1.8 inch diameter cell which is exactly twice in volume of one D cell and as such we decided not to work on this for the time being. We began the design and the procurement of tooling for the fabrication of the 3 inch diameter flat cell (for the GLLD laser designator battery) during the first quarter. We plan to concentrate our effort on the development of this flat cell during the next quarter.

In addition, we plan to continue our work on the chemistry and the electrochemistry of the  $\text{Li/SCCl}_2$  system in order to improve the safety of the system.

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## TABLE 1

Short-Circuit Currents of Hermetic D Cells with Various Cathode Designs

		Maximum Short Circuit	
<u>No.</u>	Cathode Designs	Current (A)	Current Density (mA/cm <sup>2</sup> )
1.	20 "	24	53
2.	10" 10"	76	168
3.	5 10 5	94	208
4.		58	128
5.	5 10 5	98	217
6.	25	35	62
7.	25	65	115
8.	12-1/2 12-1/2	104	184

TABLE 2

Behavior of High Rate Li/SCCl<sub>2</sub> D Cells on GLLD Modified Duty Cycle

Cathode Designs	Current	Duty Cycles above 2.0 volt	Test Temp.	
۵				
20"	6.5A	10	25° C	
ſ				
25"	6.5A	10	25° C	*
	8	7	25° C	
	6.5A	12	25° C	
n				
05.0				
25	8	4 .	25° C	
f)				
20"	6.5A	16	25° C	
L	8	11	25° C	
	6.5"	0 (11)*	-30° C	
6-1/2 12 6-1/2	6.5A	16	25° C	
	8 A	12		
				•
	6.5A	8 (11)*	-30°C	
	6.5	10 (13)*	-30° C	
	0.5	16	25°C	
	8./5A	11	25° C	

\* Number of cycles before the cell entered voltage reversal.










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26

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....



of a 3 minute 6.5A pulse every 27 minutes at -30°C









The second secon







- - -













3 minutes every 27 minutes at room temperature



room temperature





Fig. 22 Schematic drawing of the jacketed cell used for cyclic voltammetry at various temperatures



Fig. 23 Gyclic voltammogram of SOCl<sub>2</sub> in CH<sub>3</sub>CH/0.1NN(C<sub>4</sub>H<sub>9</sub>)<sub>4</sub> PF<sub>6</sub> showing the effect of sweep rate at a Pt wire electrode

a) 0.20 V/sec b) 0.50 V/sec c) 1.0 V/sec d) 2.0 V/sec







Fig. 26 Cyclic voltammogram of  $SCCl_2$  in  $CH_3CN/0.1N N(C_4H_9)PF_6$ at a silver wire electrode (0.05V/sec)













Fig. 30 UV-VIS spectrum of  $1.8 \text{ mM SCCl}_2$  in CH<sub>3</sub>CN/0.1N N(C<sub>4</sub>H<sub>9</sub>)<sub>4</sub> PF<sub>6</sub> vs CH<sub>3</sub>CN/0.1N N(C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>PF<sub>6</sub> in a 1 cm pathlength cell







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