

UNCLASSIFIED

AD NUMBER
AD866576
NEW LIMITATION CHANGE
TO Approved for public release, distribution unlimited
FROM Distribution authorized to U.S. Gov't. agencies and their contractors; Administrative/Operational Use; MAR 1970. Other requests shall be referred to Department of the Army, Fort Detrick, Attn: Technical Releases Branch/TID, Frederick, MD 21701.
AUTHORITY
BDRL D/A ltr, 29 Sep 1971

THIS PAGE IS UNCLASSIFIED

AD

AD 866576

TECHNICAL MANUSCRIPT 589

GROWTH OF L CELLS  
IN A CHEMICALLY DEFINED MEDIUM  
IN A CONTROLLED ENVIRONMENT  
CULTURE SYSTEM

Gordon W. Taylor  
John P. Kondig  
Stanley C. Nagle, Jr.  
Kiyoshi Higuchi

MARCH 1970

STATEMENT #2 UNCLASSIFIED

This document is subject to special export controls and each transmittal to foreign governments or foreign nationals may be made only with prior approval of Dept. of Army, Fort Detrick, ATTN: Technical Release Branch/TID, Frederick, Maryland 21701

DEPARTMENT OF THE ARMY  
Fort Detrick  
Frederick, Maryland

D D C  
RECEIVED  
MAR 04 1970  
REGISTERED  
C

Reproduced by the  
CLEARINGHOUSE  
for the Social Sciences & Humanities  
Washington, Springfield Va. 22151

49599

ACCESSION NO.		
CPRTI	WHITE SECTION	<input type="checkbox"/>
DDC	BUFF SECTION	<input checked="" type="checkbox"/>
UNANNOUNCED		<input type="checkbox"/>
STATION		
BY		
DISTRIBUTION AVAILABILITY CODES		
DIST.	AVAIL.	SPECIAL
2		

Reproduction of this publication in whole or in part is prohibited except with permission of the Commanding Officer, Fort Detrick, ATTN: Technical Releases Branch, Technical Information Division, Fort Detrick, Frederick, Maryland, 21701. However, DDC is authorized to reproduce the publication for United States Government purposes.

#### DDC AVAILABILITY NOTICES

Qualified requesters may obtain copies of this publication from DDC.

Foreign announcement and dissemination of this publication by DDC is not authorized.

Release or announcement to the public is not authorized.

#### DISPOSITION INSTRUCTIONS

Destroy this publication when it is no longer needed. Do not return it to the originator.

The findings in this publication are not to be construed as an official Department of the Army position, unless so designated by other authorized documents.

DEPARTMENT OF THE ARMY  
Fort Detrick  
Frederick, Maryland 21701

TECHNICAL MANUSCRIPT

GROWTH OF L CELLS IN A CHEMICALLY DEFINED MEDIUM  
IN A CONTROLLED ENVIRONMENT CULTURE SYSTEM

Gordon W. Taylor

John P. Kondig

Stanley C. Nagle, Jr.

Kiyoshi Higuchi

Medical Bacteriology Division  
BIOLOGICAL SCIENCES LABORATORIES

Project 1B562602AD01

March 1970

GROWTH OF L CELLS IN A CHEMICALLY DEFINED MEDIUM  
IN A CONTROLLED ENVIRONMENT CULTURE SYSTEM\*

ABSTRACT

Equipment has been developed to permit monitoring and automated control of environmental variables such as pH, temperature,  $pO_2$ ,  $pCO_2$ , and redox potential in order to study their effects on the growth and metabolism of cultured mammalian cells. A battery of six water-jacketed 500-ml Bellco spinner flasks was instrumented to provide (by electrode probes) information on pH,  $pO_2$ , and redox potential of each culture during growth. Stepping switches and motorized valves coupled to the sensing probes permitted control of the environment. Studies with automated control of  $pO_2$  levels in L cell cultures showed that dissolved  $O_2$  tensions of about 9% were optimal for cell growth. At  $pO_2$  values of 5 and 20%, peak cell yields as well as growth rates were reduced by approximately 20%. Peak yields of L cell cultures exceeded  $5 \times 10^6$  cells per ml when grown for 4 days without medium renewal from inocula of  $1.0 \pm 0.05 \times 10^6$  cells per ml in a defined medium sparged with 5%  $CO_2$  and adequate  $O_2$  to maintain 9% dissolved  $O_2$  tension. The redox potentials of L cell cultures reflected the  $pO_2$  levels in the medium and ranged from -25 to +150 mv (calomel reference) for  $O_2$  values ranging from 2 to 20% dissolved oxygen tension.

Workers in cell culture research have long recognized that factors such as temperature, pH,  $O_2$  partial pressure,  $CO_2$  partial pressure, and redox potential have significant effects on cell physiology. Cooper,<sup>1</sup> Paul,<sup>2</sup> Daniels,<sup>3</sup> McLimans,<sup>4</sup> Kilburn and Webb,<sup>5</sup> and Moore,<sup>6</sup> as well as many others, have employed various procedures to study the effects of one or more of these factors on the growth and metabolism of cultured mammalian cells. We believe, however, that much more remains to be done in order to understand fully the effects of these variables on cellular physiology.

Equipment developed here permits studies of effects of varied environmental conditions on mammalian cells grown in a chemically defined medium. Results obtained in this system concerning the effects of varied oxygen tension on cell growth and related data on culture pH and redox potential are included here.

The cell employed in these studies was a substrain of the mouse fibroblast cell strain L described and designated L-DR by Daniels. The culture medium was the chemically defined medium of Nagle et al.,<sup>7</sup> modified as indicated by recent studies in our laboratory (Table 1).

---

\* This report should not be used as a literature citation. Material to be published in the open literature. Readers interested in referencing the information contained herein should contact the senior author to ascertain when and where it may appear in citable form.

TABLE 1. A MODIFIED, CHEMICALLY DEFINED MEDIUM FOR THE GROWTH OF L CELLS IN SUSPENSION

Component	Per Liter
USA #1 (2X basal), Nagle <sup>7</sup>	500 ml
Vitamin mixture (100X)	10 ml
Choline chloride	48.0 mg
L-Proline	115.0 mg
L-Serine	105.0 mg
L-Asparagine	150.0 mg
NaH <sub>2</sub> PO <sub>4</sub> ·H <sub>2</sub> O	176.0 mg
Iron (as ferric ammonium citrate·3H <sub>2</sub> O)	3.14 mg
Glucose	2000 mg
Methylcellulose (15 centipoise)	1000 mg
Kanamycin/Penicillin	50 mg/100,000 units
Insulin (zinc insulin) <sup>a/</sup>	50 units
5% Sodium bicarbonate to pH 7.0	

a. May be replaced with  $10^{-6}$  M ZnSO<sub>4</sub>·7H<sub>2</sub>O.

Stock cultures of L cells were maintained at cell populations ranging from  $1 \times 10^6$  to  $10 \times 10^6$  cells per ml in 25 ml of medium in 100-ml serum bottles on the gyrotory shaker at 35 C with daily medium replacement. Inocula for our larger culture vessels were grown in shaken 250-ml Woulff bottles containing 100 ml of culture medium (Fig. 1). These vessels required venting when populations exceeded  $10^6$  cells per ml because of the need for more adequate gas exchange. Excellent pH control was thereby obtained because the escape of metabolic CO<sub>2</sub> prevented excessive drop in pH of the cultures.

The experimental cultures were inoculated with  $1.00 \pm 0.05 \times 10^6$  cells per ml and were grown in a battery of six water-jacketed, 500-ml Bellco spinner flasks, arranged on a table with magnetic stirrer drives. The temperature in the culture vessels was maintained at 35 C with a Haake constant-temperature water circulator. Each spinner flask was constructed with four ports (Fig. 2). A Yellow Springs oxygen probe was inserted in one port, a platinum electrode and a combination pH probe were inserted in the second and third ports. All probes were chemically sterilized in acid-alcohol for 10 minutes, rinsed in sterile distilled water, and aseptically inserted into the flask. The remaining port was used for sampling and gas sparging lines, and to vent exhaust gases.

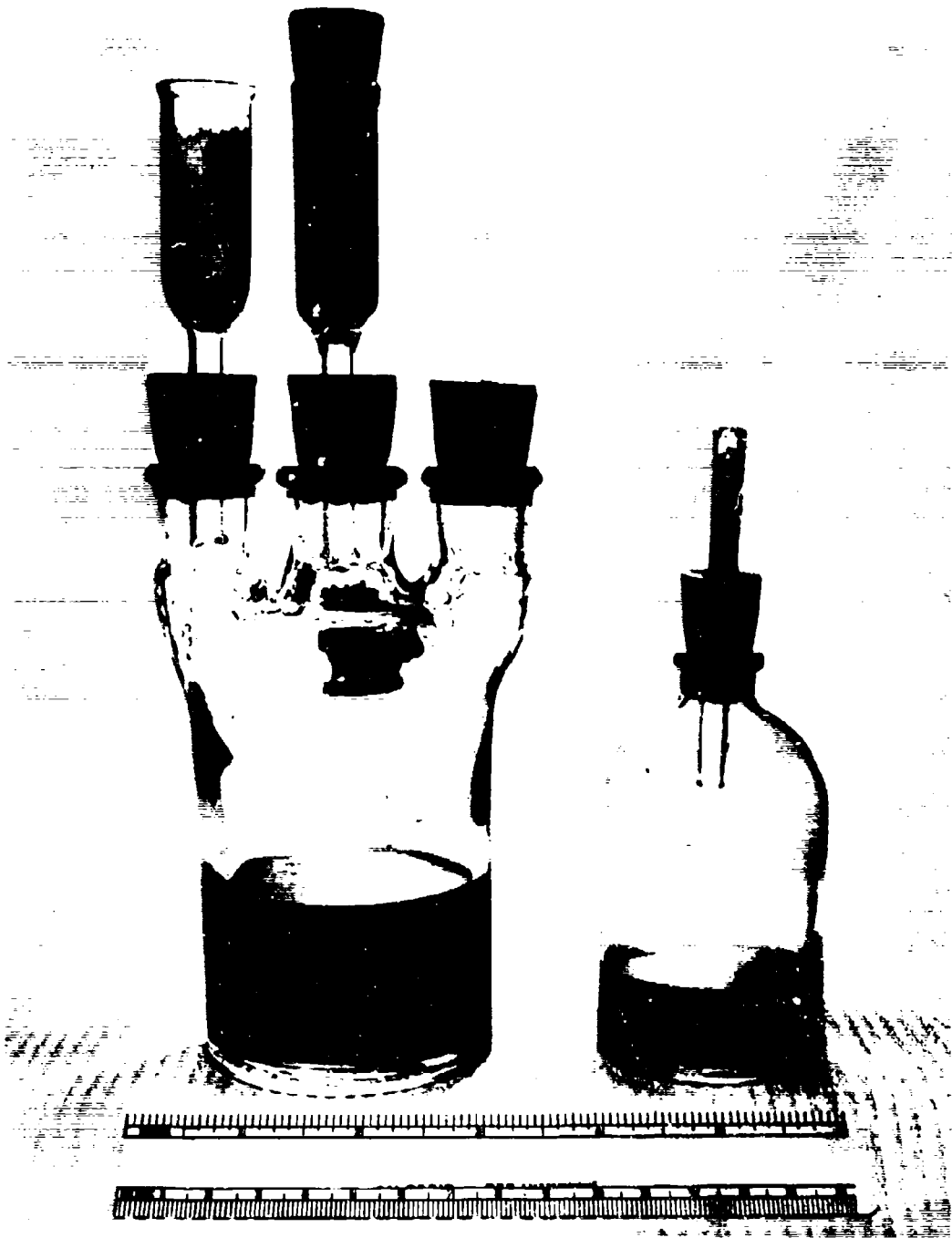


FIGURE 1. L Cell Cultures in Vented Wouff (left) and Serum (right) Bottles.

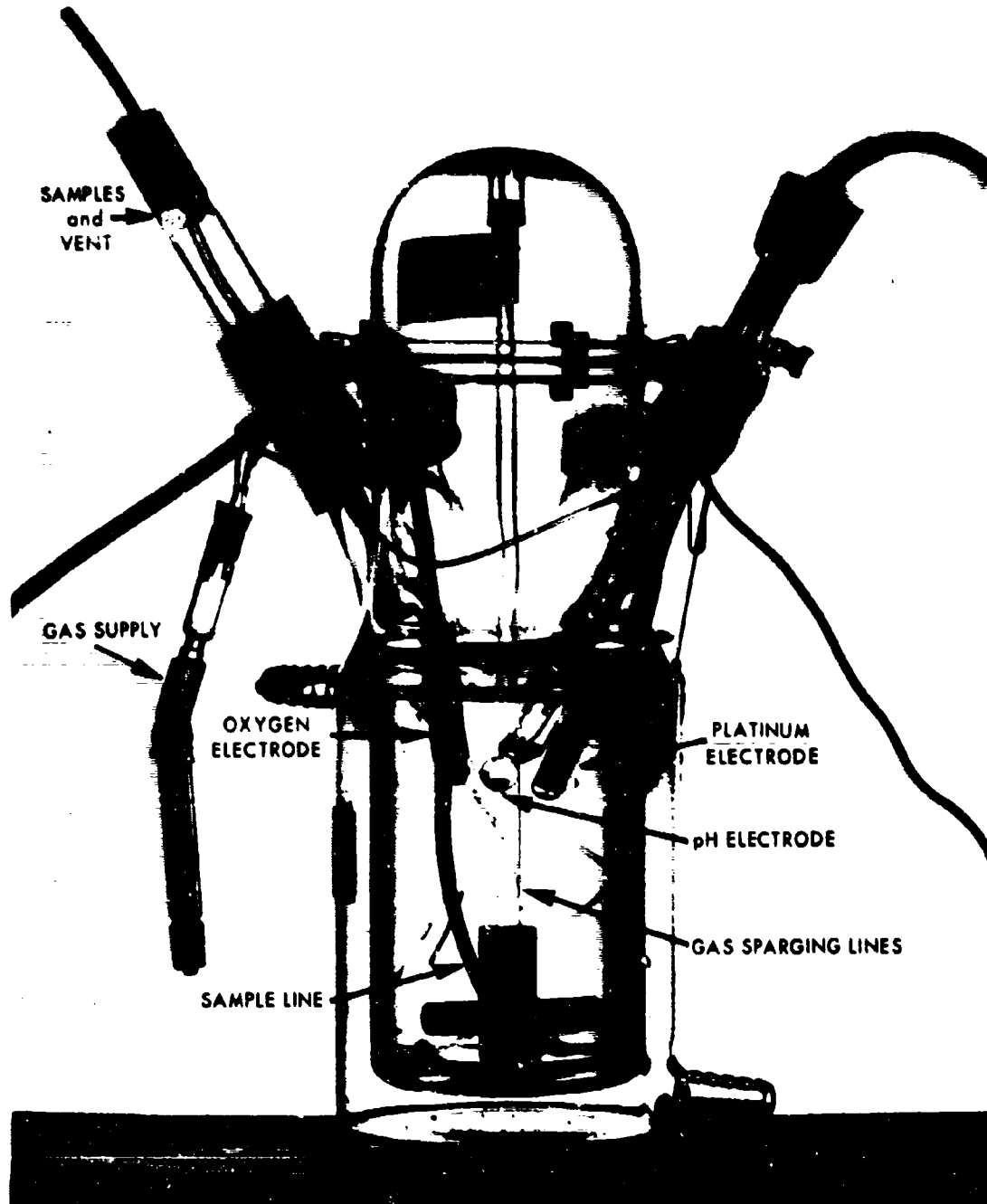


FIGURE 2. Single Instrumented Spinner Flask.

Initial results showed that even with continuous sparging with a gas mixture consisting of 10% oxygen, 5% CO<sub>2</sub>, and 85% N<sub>2</sub>, the dissolved oxygen tension dropped to near zero when cell populations approached  $4 \times 10^6$  cells per ml. Because of this problem of maintaining pO<sub>2</sub> values at preselected levels, a control system was designed that maintained pO<sub>2</sub> at desired levels by varying O<sub>2</sub> input to the sparge gas mixture in response to changes in oxygen electrode readings. This system is shown in Figure 3.

Sequential measurements of oxygen tension in each flask could be recorded with stepping switches. Similarly, the redox potential and pH values of each culture could be determined and recorded as shown in Figure 4. The complete controlled environment culture system is illustrated in Figure 5.

The growth curve of L cells obtained in this system with the pO<sub>2</sub> maintained at 9%, together with continuously recorded pH and redox values, is shown in Figure 6. As shown here, relatively uniform values of medium pH and of redox potential were maintained during the growth cycle under controlled pO<sub>2</sub> condition. In our culture system, pH values generally remained within  $6.9 \pm 0.2$  pH units at varied pO<sub>2</sub> levels without further adjustment as long as the CO<sub>2</sub> content of the sparging gas mixture was maintained at 5%. On the other hand, redox potentials clearly reflected differences in pO<sub>2</sub> values. When growth of L cell cultures was compared over a pO<sub>2</sub> range of 2 to 20%, optimum growth was obtained at a pO<sub>2</sub> value of approximately 9% (Fig. 7). This value agrees with results reported by other workers cited earlier. Redox potentials of cultures at pO<sub>2</sub> values ranging from 2 to 20% showed readings, based on a saturated calomel reference electrode, ranging from -25 to +150 millivolts (Fig. 8). It appears, therefore, that the redox potential for optimal growth of L cells is approximately +90 millivolts. This result is in essential agreement with the results obtained by Daniels<sup>6</sup> and Wiles.<sup>5</sup>

Continuing investigation with this equipment will include studies on the effects of varied CO<sub>2</sub> partial pressures as well as effects of varied pH on cell growth and physiology. The equipment is versatile and has many other applications to studies of effects of environment on cell physiology.

In summary, we have described equipment designed to permit study of the effects of environmental variables on the physiology of cultured mammalian cells. In a totally synthetic culture medium and an automatically controlled system, an oxygen partial pressure of approximately 9% was optimal for growth of L cells. The applicability of this system to studies on effects of such variables as pH, pCO<sub>2</sub>, and redox potential is evident.

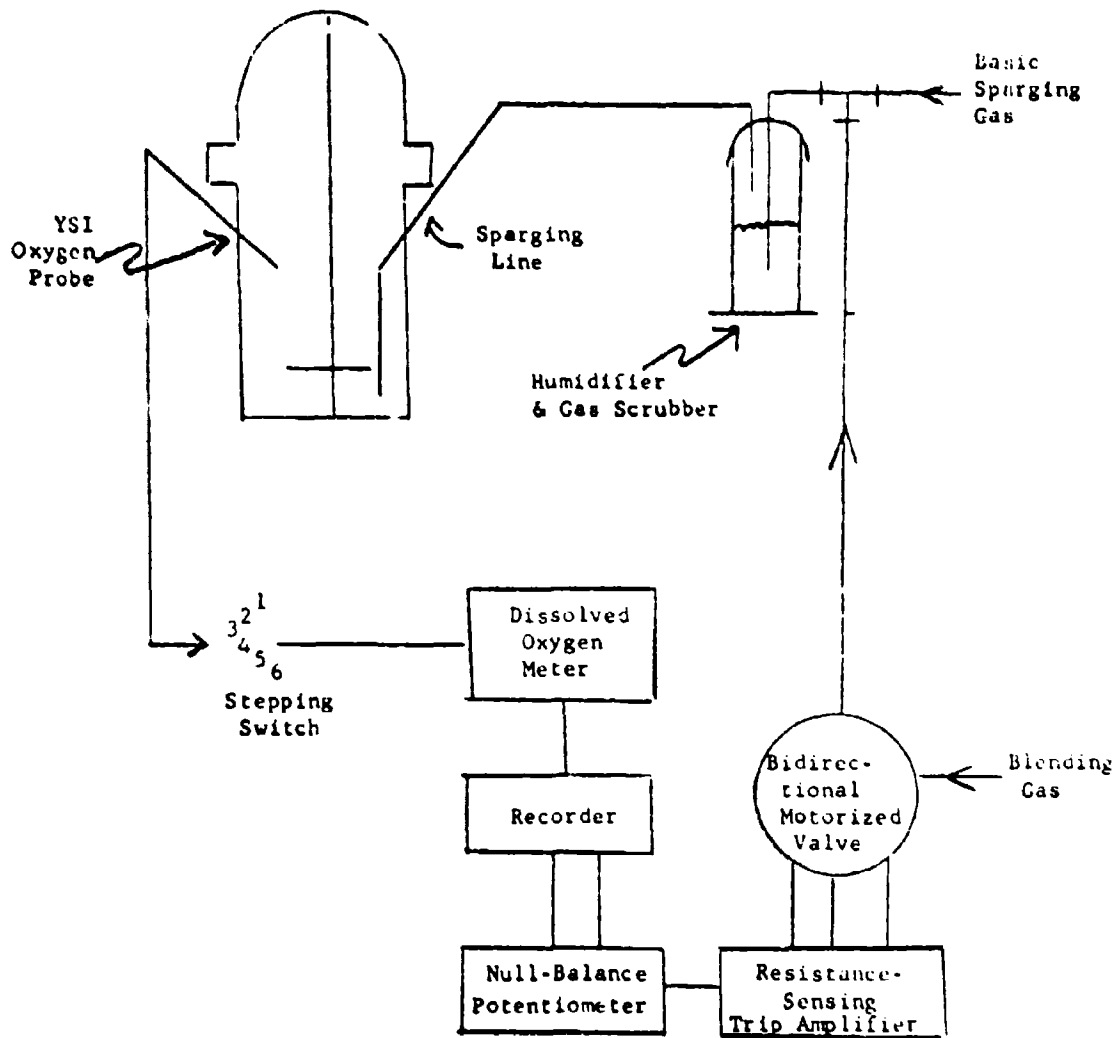


FIGURE 3. Block Diagram of Automated Oxygen Control System for Cell Cultures.

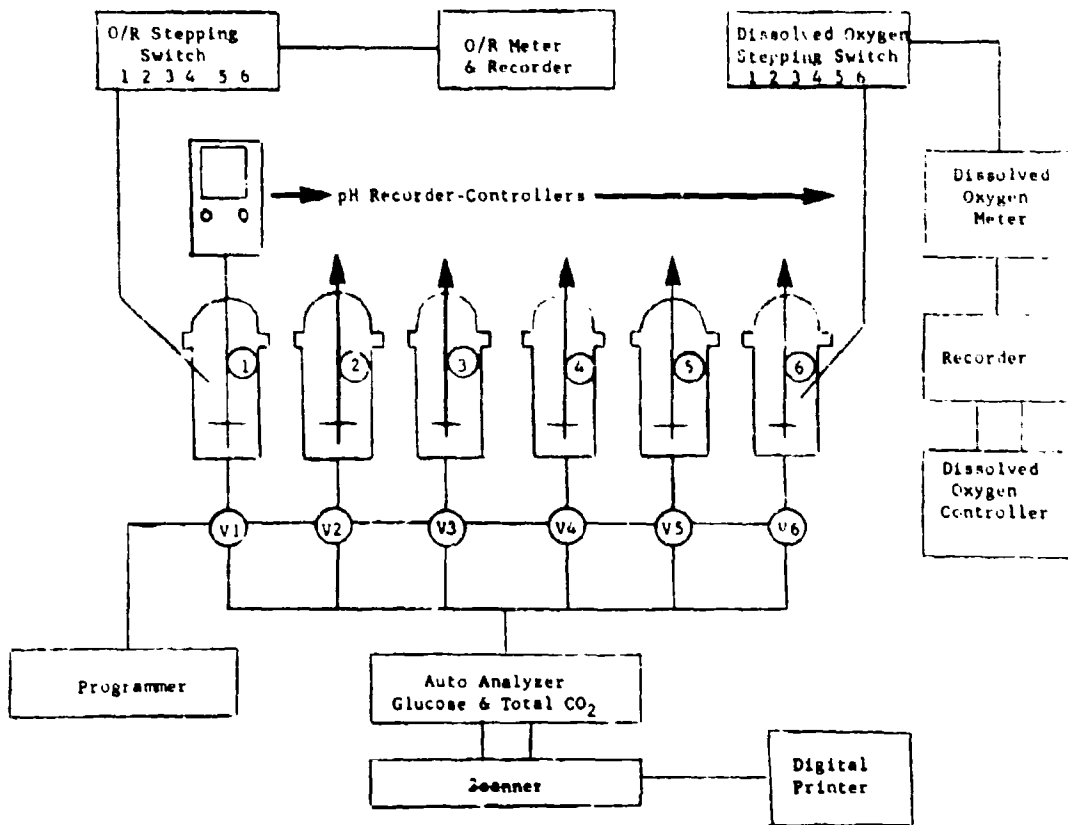


FIGURE 4. Block Diagram of Automated Cell Culture Environmental Control and Monitoring System.

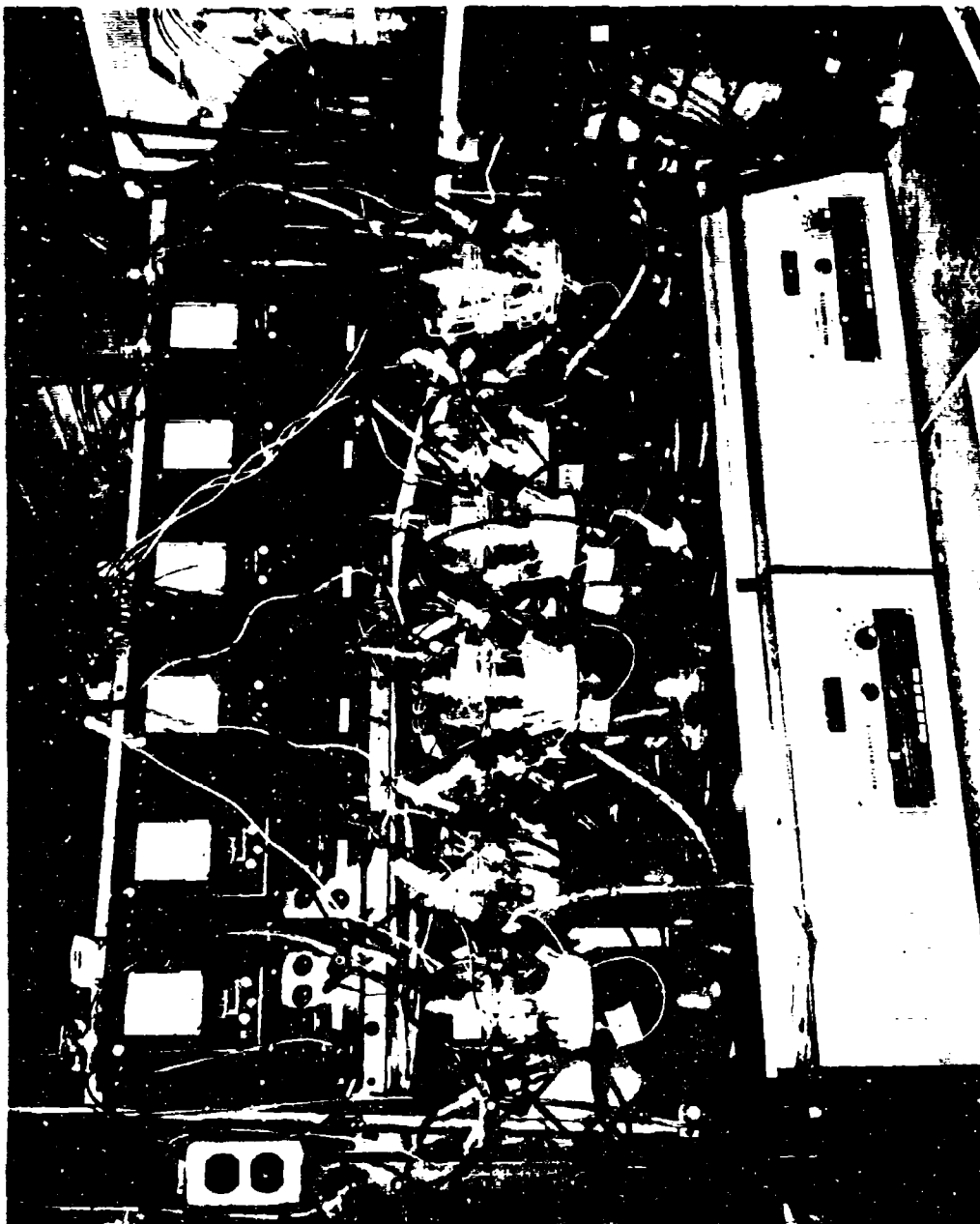


FIGURE 5. Complete Automated Cell Culture Environmental Control and Monitoring System.

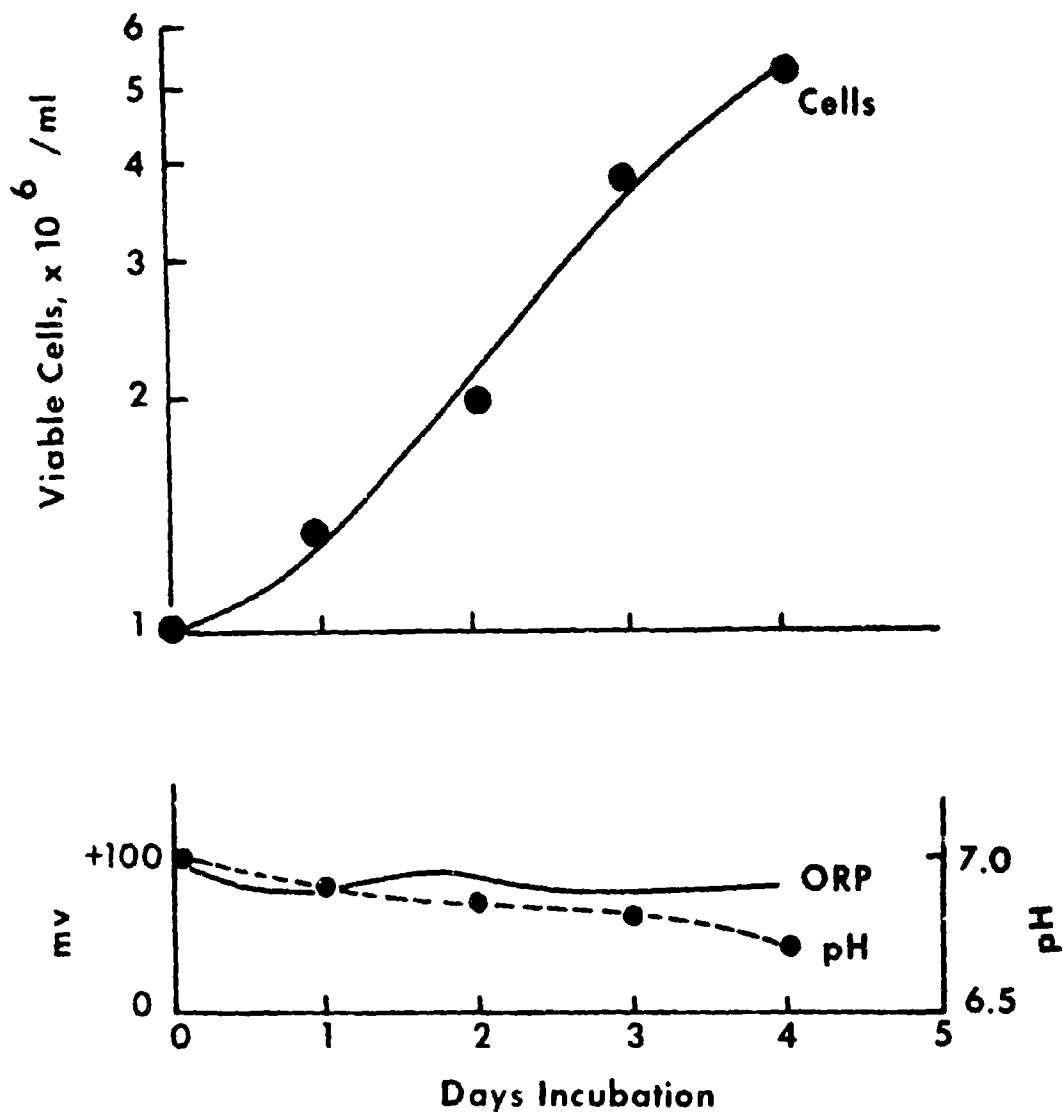


FIGURE 6. Growth, ORP, and pH Values of L Cell Cultures Controlled at 9% Dissolved Oxygen Tension.

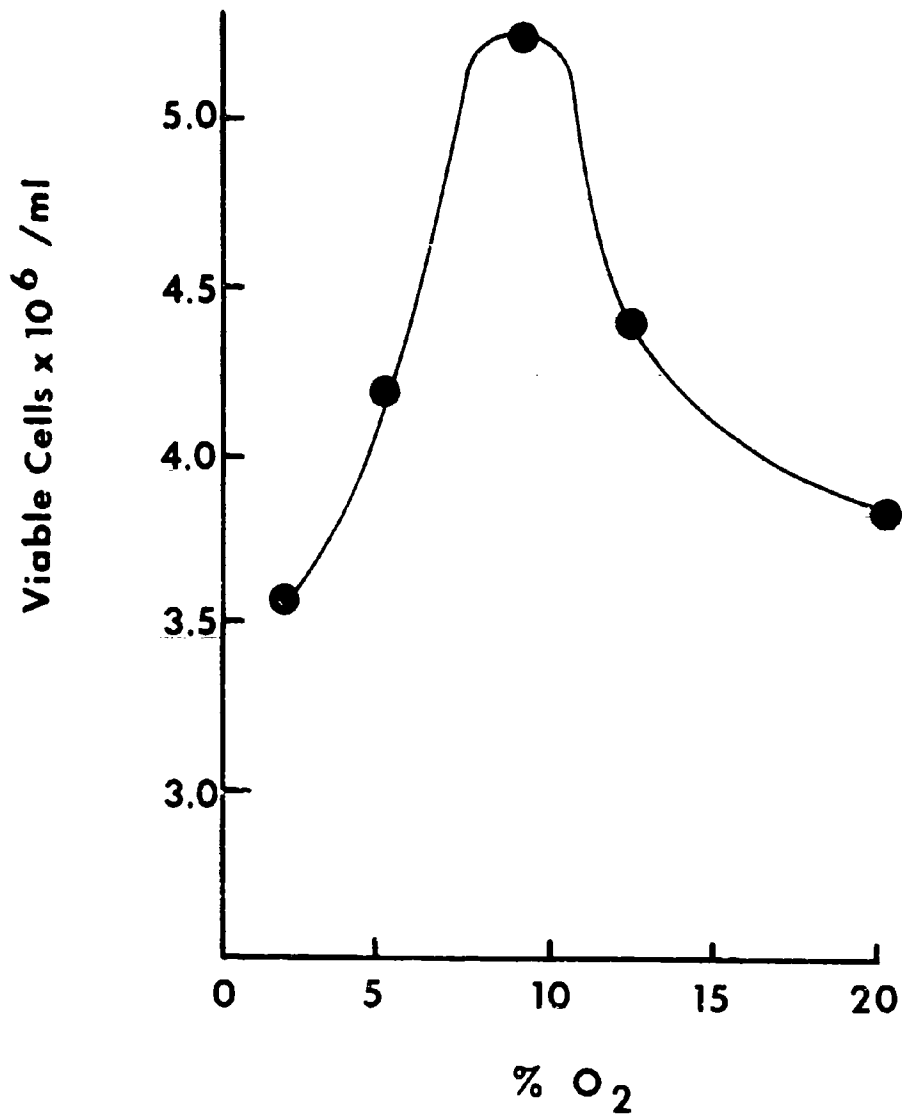


FIGURE 7. Peak Yields of L Cell Cultures Grown At Various Controlled Dissolved Oxygen Tensions.

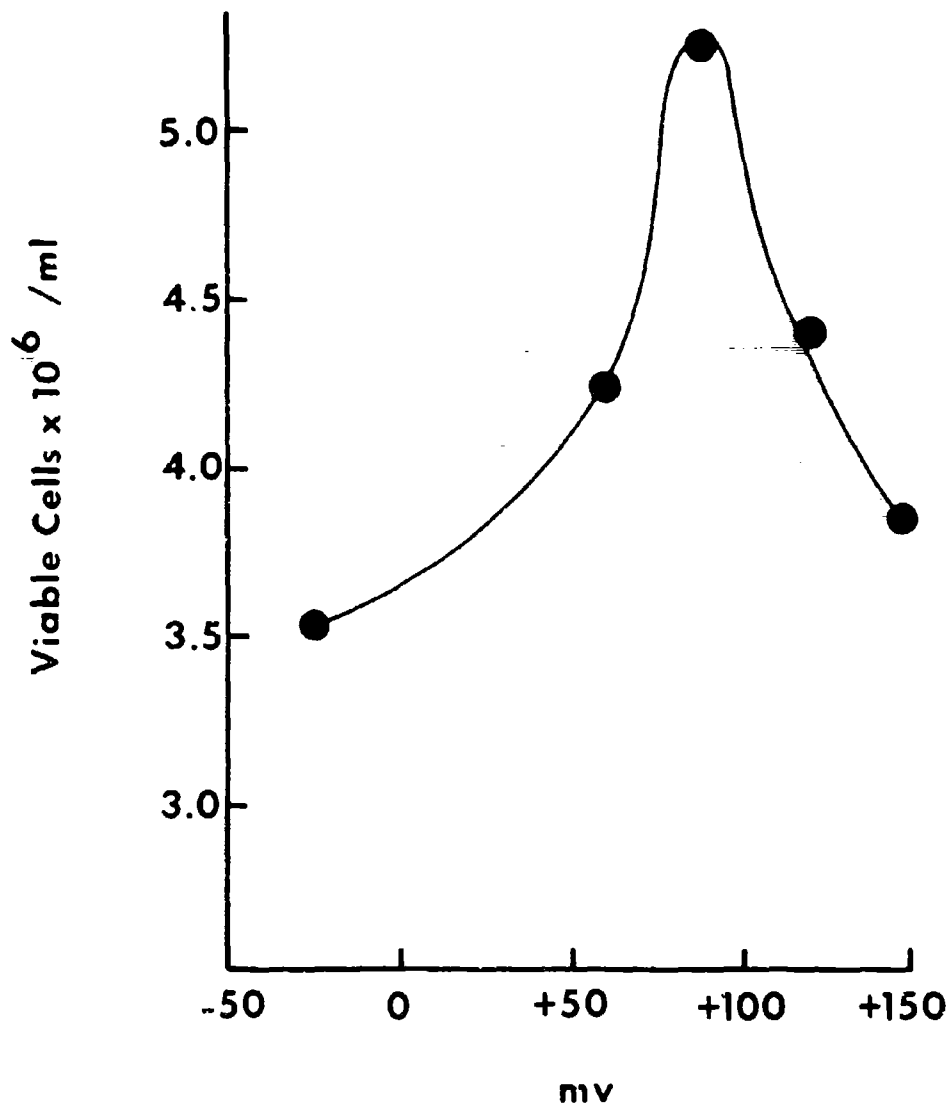


FIGURE 8. Correlation Between Peak Yields and Recorded Redox Potentials of L Cell Cultures.

LITERATURE CITED

1. Cooper, P.D.; Wilson, J.N.; Burt, A.M. 1957. The bulk growth of animal cells in continuous suspension culture. *J. Gen. Microbiol.* 21:702-720.
2. Paul, J. 1959. Environmental influences on the metabolism and composition of cultured cells. *J. Exp. Zool.* 142:475-505.
3. Daniels, W.F.; Parker, D.A.; Johnson, R.W.; Schneider, L.E. 1965. Controlled pH and oxidation-reduction potential with a new glass tissue-culture fermentor. *Biotechnol. Bioeng.* 7:529-553.
4. McLimans, W.F.; Mount, D.T.; Bogitch, S.; Crouse, E.J.; Harris, G.; Moore, G.E. 1966. A controlled environment system for study of mammalian cell physiology. *Ann. N.Y. Acad. Sci.* 139:190-213.
5. Kilburn, D.G.; Webb, F.C. 1968. The cultivation of animal cells at controlled dissolved oxygen partial pressure. *Biotechnol. Bioeng.* 10:801-814.
6. Moore, G.E.; Hasenpusch, P.; Gerner, R.E.; Burns, A.A. 1968. A pilot plant for mammalian cell culture. *Biotechnol. Bioeng.* 10:625-640.
7. Nagle, S.C., Jr.; Tribble, H.R., Jr.; Anderson, R.E.; Gary, N.D. 1963. A chemically defined medium for growth of animal cells in suspension. *Proc. Soc. Exp. Biol. Med.* 112:340-344.
8. Daniels, W.F.; Garcia, L.H.; Rosensteel, J.F. 1969. The importance of oxidation-reduction potential levels in the growth of tissue cell lines using Earle's cells as a prototype. Paper 65A, American Institute of Chemical Engineers Symposium on Bioengineering Techniques, Washington, D.C., November 1969.
9. Wiles, C.C.; Smith, V.C. 1969. Oxidation-reduction potential controlled submerged tissue culture fermentation in pilot-scale fermentors. Paper 65B, American Institute of Chemical Engineers Symposium on Bioengineering Techniques, Washington, D.C., November 1969.

Unclassified

19

Security Classification		
DOCUMENT CONTROL DATA - R & D		
<i>(Security classification of title, body of abstract and indexing annotation must be entered when the overall report is classified)</i>		
1. ORIGINATING ACTIVITY (Corporate author) Department of the Army Fort Detrick, Frederick, Maryland, 21701		20. REPORT SECURITY CLASSIFICATION Unclassified
2. REPORT TITLE GROWTH OF L CELLS IN A CHEMICALLY DEFINED MEDIUM IN A CONTROLLED ENVIRONMENT CULTURE SYSTEM		
3. AUTHOR(S) (First name, middle initial, last name) Gordon W. Taylor      Stanley C. Nagle, Jr. John P. Kondig      Kiyoshi (NMI) Higuchi		
4. REPORT DATE March 1970	7A. TOTAL NO. OF PAGES 20	7B. NO. OF REFS 9
5A. CONTRACT OR GRANT NO. a. PROJECT NO 1B562602AD01 b. Task-Work Unit 01-018 c. DD 1498 Agency Access, DA OL 0236	5B. ORIGINATOR'S REPORT NUMBER(S) Technical Manuscript 589	
5C. OTHER REPORT NO(S) (Any other numbers that may be assigned this report) CMs 6652 SMUFD-AE-T 49599		
10. DISTRIBUTION STATEMENT Qualified requesters may obtain copies of this publication from DDC. Foreign announcement and dissemination of this publication by DDC is not authorized. Release or announcement to the public is not authorized.		
11. SUPPLEMENTARY NOTES Medical Bacteriology Division	12. SPONSORING MILITARY ACTIVITY Department of the Army Fort Detrick, Frederick, Md., 21701	
13. ABSTRACT Equipment has been developed to permit monitoring and automated control of environmental variables such as pH, temperature, pO <sub>2</sub> , pCO <sub>2</sub> , and redox potential in order to study their effects on the growth and metabolism of cultured mammalian cells. A battery of six water-jacketed 500-ml Bellco spinner flasks was instrumented to provide (by electrode probes) information on pH, pO <sub>2</sub> , and redox potential of each culture during growth. Stepping switches and motorized valves coupled to the sensing probes permitted control of the environment. Studies with automated control of pO <sub>2</sub> levels in L cell cultures showed that dissolved O <sub>2</sub> tensions of about 9% were optimal for cell growth. At pO <sub>2</sub> values of 5 and 20%, peak cell yields as well as growth rates were reduced by approximately 20%. Peak yields of L cell cultures exceeded 5 x 10 <sup>6</sup> cells per ml when grown for 4 days without medium renewal from inocula of 1.0 ± 0.05 x 10 <sup>6</sup> cells per ml in a defined medium sparged with 5% CO <sub>2</sub> and adequate O <sub>2</sub> to maintain 9% dissolved O <sub>2</sub> tension. The redox potentials of L cell cultures reflected the pO <sub>2</sub> levels in the medium and ranged from -25 to +150 mv (calomel reference) for O <sub>2</sub> values ranging from 2 to 20% dissolved oxygen tension.		

DD FORM 1473

REPLACES DD FORM 1473, 1 JAN 66, WHICH IS OBSOLETE FOR ARMY USE.

Unclassified  
Security Classification

Unclassified  
Security Classification

14. KEY WORDS	LINK A		LINK B		LINK C	
	ROLE	WT	ROLE	WT	ROLE	WT
Culture system L cells Chemically defined medium Environmental control Controlled environment Suspension cultures Redox and pO <sub>2</sub> relationship ORP and pO <sub>2</sub> relationship Oxygen tension						

Unclassified  
Security Classification