

AD-A047 271

TEXAS A AND M UNIV COLLEGE STATION DEPT OF CHEMISTRY

F/G 7/3

AN ELECTROCHEMICAL INVESTIGATION OF SOME RHENIUM CARBONYL PORPH--ETC(U)

NOV 77 D D AXTELL, G R MILLER, T H RIDGWAY

N00014-75-C-0417

UNCLASSIFIED

TR-18

NL

| OF |
AD
A047 271



END

DATE
FILMED

1 - 78

DDC

AD A047271

12
B.S.

OFFICE OF NAVAL RESEARCH

Contract N00014-75-C-0417

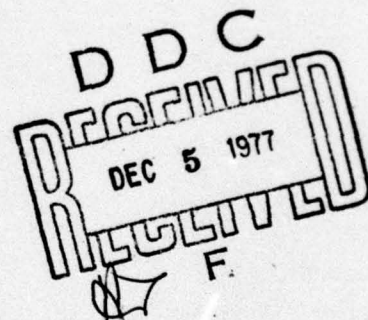
Task No. NR 053-559
Technical Report No. 18

An Electrochemical Investigation of Some Rhenium
Carbonyl Porphyrins

Darrell D. Axtell, G.R. Miller, Tom H. Ridgway, and
Minoru Tsutsui

Prepared for Publication
in the
Journal of American Chemical Society

Texas A&M University
Department of Chemistry
College Station, Texas 77843



November 18, 1977

Reproduction in whole or in part is permitted for
any purpose of the United States Government

Approved for Public Release; Distribution Unlimited.

AD No. —
DDC FILE COPY

REPORT DOCUMENTATION PAGE		READ INSTRUCTIONS BEFORE COMPLETING FORM
1. REPORT NUMBER 9 Technical Report ✓	2. GOVT ACCESSION NO.	3. RECIPIENT'S CATALOG NUMBER
4. TITLE (and Subtitle) 6 AN ELECTROCHEMICAL INVESTIGATION OF SOME RHENIUM CARBONYL PORPHYRINS.	5. TYPE OF REPORT & PERIOD COVERED Interim	
7. AUTHOR(s) 10 DARRELL D. Axtell, G.R. Miller, Tom H. Ridgway Minoru Tsutsui	6. PERFORMING ORG. REPORT NUMBER 14 TR-18	
9. PERFORMING ORGANIZATION NAME AND ADDRESS Department of Chemistry ✓ Texas A&M University College Station, Texas 77843	8. CONTRACT OR GRANT NUMBER(s) 15 N00014-75-C-0417	
11. CONTROLLING OFFICE NAME AND ADDRESS Office of Naval Research Department of the Navy Arlington, Virginia 22217	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS NR 053-559	
14. MONITORING AGENCY NAME & ADDRESS (if different from Controlling Office)	12. REPORT DATE 11-18-77	
	13. NUMBER OF PAGES 8 12 10p.	
	15. SECURITY CLASS. (of this report) unclassified	
	15a. DECLASSIFICATION/DOWNGRADING SCHEDULE	
16. DISTRIBUTION STATEMENT (of this Report) Approved for Public Release, Distribution Unlimited		
17. DISTRIBUTION STATEMENT (of the abstract entered in Block 20, if different from Report)		
18. SUPPLEMENTARY NOTES Submitted to the Journal of American Chemical Society		
19. KEY WORDS (Continue on reverse side if necessary and identify by block number) Rhenium Carbonyl Porphyrins meso-tetraphenylporphine Cyclic Voltmetric Studies [meso-tetraphenylporphinato]bis[tricar- Re-Re Bond bonylrhenium(I)] (meso-porphinato)tricarboxylrhenium(I) meso-tetraphenylporphinato cobalt(II)		
20. ABSTRACT (Continue on reverse side if necessary and identify by block number) Cyclic voltammetric studies of $[\text{Re}(\text{C})_3]_2\text{TPP}$ and $[\text{Re}(\text{CO})_3]\text{H-TPP}$ were carried out in an attempt to verify the previously proposed formation of an Re-Re bond by chemical oxidation. This study indicates that the first oxidation step of both complexes is similar, but the next oxidation step is very different. However, the final product species are very similar. We conclude that electrochemical oxidation of the $[\text{Re}(\text{CO})_3]\text{H-TPP}$ is accompanied by a chemical rearrangement leading to formation of a species containing two Re atoms connected by at least a partial Re-Re bond.		

DD FORM 1 JAN 73 1473

EDITION OF 1 NOV 65 IS OBSOLETE
S/N 0102-LF-014-6601

SECURITY CLASSIFICATION OF THIS PAGE (When Data Entered)

407997

DDC
RECEIVED
DEC 5 1977
F. B. I.

AN ELECTROCHEMICAL INVESTIGATION
OF SOME RHENIUM CARBONYL PORPHYRINS

D.D. Axtell, G.R. Miller⁸, T.H. Ridgway⁸, and M. Tsutsui*

Department of Chemistry
Texas A & M University
College Station, Texas 77843

ABSTRACT:

Cyclic voltammetric studies of $[\text{Re}(\text{CO})_3]_2\text{TPP}$ and $[\text{Re}(\text{CO})_3]\text{H-TPP}$ were carried out in an attempt to verify the previously proposed formation of an Re-Re bond by chemical oxidation. This study indicates that the first oxidation step of both complexes is similar, but the next oxidation step is very different. However, the final products are very similar. We conclude that electrochemical oxidation of the $[\text{Re}(\text{CO})_3]\text{H-TPP}$ is accompanied by a chemical rearrangement leading to formation of a species containing two Re atoms connected by at least a partial Re-Re bond.

ACCESSION for		White Section ☒
NTIS		Buff Section ☐
DDC		
UNANNOUNCED		
JUSTIFICATION		
BY DISTRIBUTION/AVAILABILITY CODES		
DI		SPECIAL
A		

1

AN ELECTROCHEMICAL INVESTIGATION
OF SOME RHENIUM CARBONYL PORPHYRINS

Sir:

Earlier reports^{1,2} described the synthesis and x-ray crystallographic characterization of mono- and di-rhenium carbonyl porphyrins. The x-ray data indicated that in some cases an inter-molecular rhenium-rhenium bond was formed as a result of chemical oxidation. We wish to report electrochemical evidence which tends to confirm the formation of such a metal-metal bond.

The use of cyclic voltammetry to probe the energy levels of metal complexes and to investigate the stability of the resulting oxidation and reduction states is well established^{3,4}. The cyclic voltammetric studies were carried out on a locally constructed instrument, the details of which will appear elsewhere. The three-electrode electrochemical cell is similar to the form developed by VanDuyne⁵, having a Lugen probe for the reference electrode. This allows placement of the probe tip within 1mm of the working electrode, thus minimizing solution resistance effects. The auxiliary electrode is a platinum wire concentrically wound around a glass rod containing an axially mounted 4mm platinum wire used to form the working electrode. The rod was end polished and the electrode buffed between experiments with Whatman #42 filter

— | —

paper to eliminate film formation. The reference electrode was a silver-silver perchlorate in acetonitrile system with an asbestos fiber wick junction to the working solution. Vacuum-dried electrochemical grade tetraethylammonium perchlorate (TEAP) was used as the supporting electrolyte. The complexes were prepared as previously described⁶. The solvent was spectroscopic grade dichloromethane. All solutions were 0.1 M in TEAP and 1-2 millimolar in complex or H_2TPP . The solutions were deoxygenated by saturation with purified nitrogen. The gas was passed through a three stage drying train to remove water and then through a scrubbing tower filled with supporting electrolyte solution. This final step was necessary to presaturate the gas stream and minimize concentration changes caused by evaporation of solvent from the samples.

The electrochemical results obtained for free meso-tetraphenylporphine, H_2TPP , [monohydrogen meso-tetraphenylporphinato]tricarbonylrhenium(I), $(Re(CO)_3HTPP)$, 1, [meso-tetraphenylporphinato]bis[tricarbonylrhenium(I)], $(Re(CO)_3)_2TPP$, 2, and meso-tetraphenylporphinato cobalt(II), $CoTPP$, are shown in Figure 1 as curves A, B, C and D respectively. $CoTPP$ is included as a reference compound because its electrochemistry has been studied previously in this media.⁷

The free H_2TPP , mono- and di-rhenium compounds all exhibit two consecutive, chemically stable, one electron

reductions with peak potentials of -1.52 and -1.83 V for H_2TPP (curve A), -1.41 and -1.74 V for (1), and -1.20 and -1.54 V versus reference for (2). $CoTPP$ shows a chemically stable, reversible one electron transfer process at -1.00 V, but a second reduction (not shown) near -1.60 V leads to a chemically unstable product under these conditions. The chemical kinetics involved can be outrun by using sweep rates in excess of 10 V/s. The reductive electrochemical behavior is unremarkable, exhibiting the expected lowering of energy levels of the first available orbitals as one proceeds from H_2TPP to the mono- and di-rhenium species.

The oxidative electrochemistry of these species is somewhat more complex. Oxidation of H_2TPP occurs in two consecutive one-electron steps at 1.04 and 0.80 V and both product species are stable in solution on the time scale of the experiment (curve A). The di-rhenium complex (2) (curve C) also shows consecutive one-electron oxidations at 0.81 and 1.22 V versus reference. Oxidation of $CoTPP$ is in accord with the results of other workers⁷, with the $Co(II) \longrightarrow Co(III)$ oxidation occurring at 0.57 V and the two ligand oxidations occurring at 0.96 and 1.14 V (curve D). Oxidation of the mono-rhenium species (2) shows an oxidation process whose peak potential occurs at 0.68 V at a sweep rate of 0.1 V/s. The resulting oxidation product is chemically unstable on this time scale although some indication of a

re-reduction can be seen.

A second oxidation occurs at about 0.90 V and the product is even less stable in solution. Both oxidations exhibit a peak current which is essentially identical to that found for the reductions indicating that both are probably one-electron processes. At more positive potentials a further oxidation occurs at 1.22 V versus reference. This product is stable on the time scale of the experiment, while the amount of current obtained is consistent with a two-electron oxidation. Increasing the sweep rate shifts the first wave more anodically and increases the re-reduction current although a shape characteristic of a stable chemical species was not obtained at sweep rates up to 50 V/s where a peak potential of about 0.74 V was observed. The second charge-transfer process is affected very little by the increased sweep rate. Because it was not possible to outrun the chemical kinetic processes involved in the first two oxidations, one can not state the corresponding oxidation potentials precisely, but one can set lower limits of 0.74 and 0.95 V versus reference respectively on the basis of the high sweep rate studies.

The correspondence of the first oxidation potential for H_2TPP and 2 would seem to indicate that the orbitals involved both come from the ligand moiety and contain very little metal character. The second oxidation of 2 involves an orbital having a fair degree of metal character as evidenced by the 0.18 V shift in potential. There are several interpretations for the observed electrochemical behavior of 1 . One can argue

that the first oxidation corresponds to the first process observed in 2 but that the oxidized species then decomposes due to a localized charge defect which is not found in either 2 or H_2TPP . This charge defect could be on the rhenium atom or on the porphyrin ligand. If the site of the defect was interior to the ring structure, it would be shielded by the rhenium atoms in 2 . In conjunction with the x-ray crystallographic data, it seems more likely that the charge defect is on the rhenium atom since 2 shows evidence of an existing metal-metal bond which would tend to stabilize the oxidation product of 2 relative to 1 . The correspondence between the final oxidation potentials of both 1 and 2 indicates that the electrons involved come from orbitals of very nearly the same energy, while the relative stability of the two oxidized species indicates that the moieties have similar chemical properties in the experimental environment. These observations, in conjunction with the two electron transfer involved in the final oxidation of 1 make it tempting to speculate that the chemical rearrangements involved lead to a dimerization of 1 with the formation of at least a partial rhenium-rhenium bond.

This hypothesis is merely speculative at this time but it is consistent with the available data. Further studies involving bulk electrolysis and spectro-electrochemical techniques are in progress in an attempt to clarify the situation.

ACKNOWLEDGMENTS: We wish to acknowledge support of this research project by the Office of Naval Research.

D.D. Axtell, G.R. Miller⁸, T.H. Ridgway⁸, M. Tsutsui*

Department of Chemistry
Texas A & M University
College Station, Texas 77843

REFERENCES AND NOTES:

1. C.P. Hsung, M. Tsutsui, D.L. Cullen and E.F. Meyer, Jr., J. Amer. Chem. Soc., 98, 7878 (1976).
2. S. Kato, M. Tsutsui, D.L. Cullen and E.F. Meyer, Jr., J. Amer. Chem. Soc., 99, 620 (1977).
3. E.R. Brown and R.F. Large in "Physical Methods of Chemistry", Part IIA, "Electrochemical Methods", A. Weissberger and B.W. Rossiter, Eds., Wiley-Interscience, New York, 1971, 423-530.
4. J. Piekarski and R.N. Adams, Ibid., pp. 531-590.
5. R.P. VanDuyne and C.N. Reilly, Anal. Chem., 44, 142, 153, 158 (1972).
6. (a) D.L. Cullen, E.F. Meyer, Jr., S.T. Srivastava and M. Tsutsui, J. Amer. Chem. Soc., 94, 7603 (1972).
(b) M. Tsutsui and C.P. Hsung, Chem. Letters, 941, (1973).
7. K.M. Kadish and D.G. Davis, Annals N.Y. Acad. Sci., 206, 495 (1973).
8. Present address: Department of Chemistry, University of Cincinnati, Cincinnati, Ohio 45221.

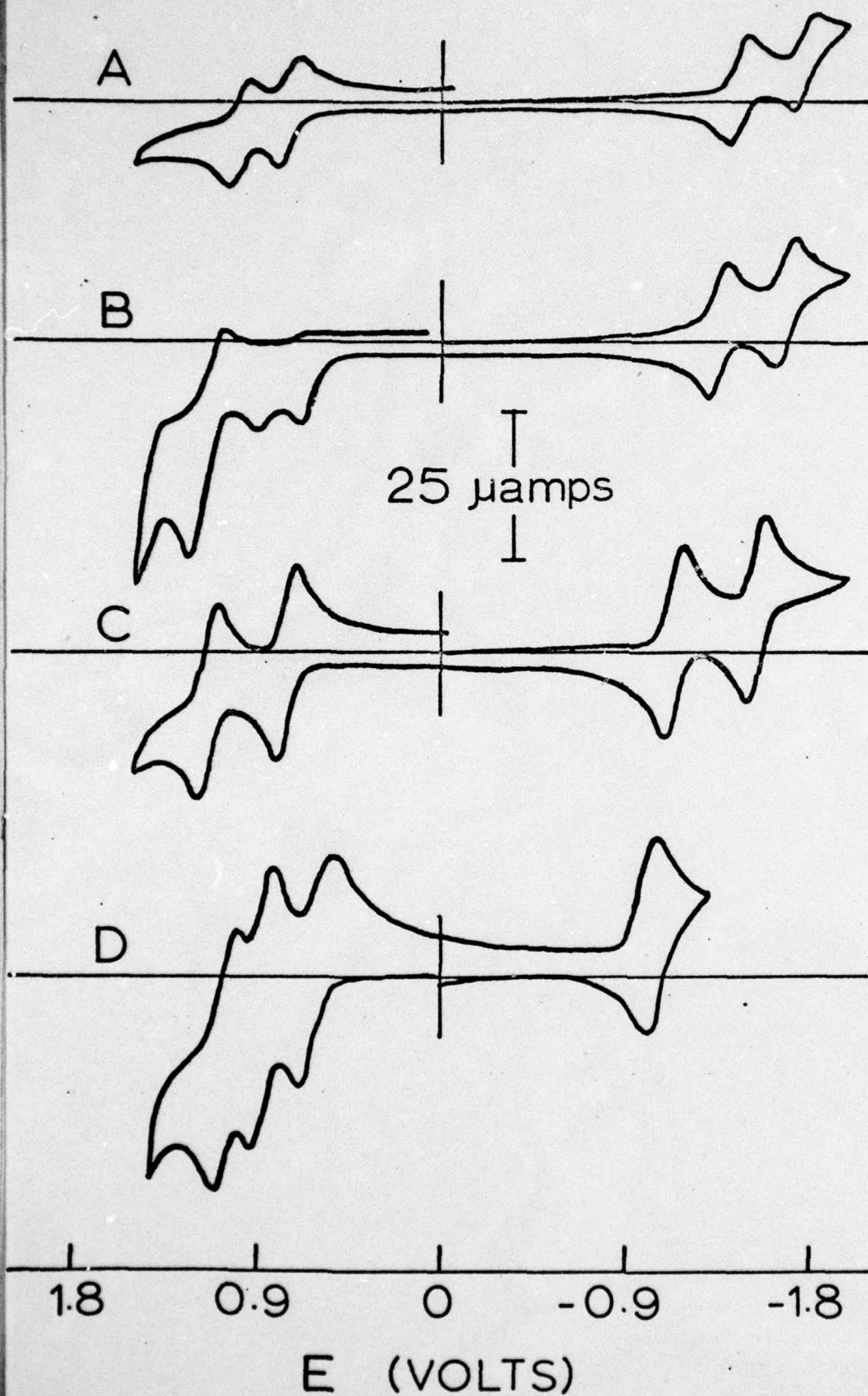


Figure 1. Cyclic voltammograms in 0.1M TEAP at a platinum electrode of (a) H_2TPP , (b) $[\text{Re}(\text{CO})_3]\text{HTPP}$, (1), (c) $[\text{Re}(\text{CO})_3]_2\text{TPP}$, (2), and (d) CoTPP . Scan rate, 0.1 V/S.

TECHNICAL REPORT DISTRIBUTION LIST

<u>No. Copies</u>		<u>No. Copies</u>	
2	Office of Naval Research Arlington, Virginia 22217 Attn: Code 472	12	Defense Documentation Center Building 5, Cameron Station Alexandria, Virginia 22314
6	Office of Naval Research Arlington, Virginia 22217 Attn: Code 102IP 1	1	U.S. Army Research Office P.O. Box 12211 Research Triangle Park, N.C. 27709 Attn: CRD-AA-IP
1	ONR Branch Office 536 S. Clark Street Chicago, Illinois 60605 Attn: Dr. Jerry Smith	1	Naval Ocean Systems Center San Diego, California 92152 Attn: Mr. Joe McCartney
1	ONR Branch Office 715 Broadway New York, New York 10003 Attn: Scientific Dept.	1	Naval Weapons Center China Lake, California 93555 Attn: Head, Chemistry Division
1	ONR Branch Office 1030 East Green Street Pasadena, California 91106 Attn: Dr. R. J. Marcus	1	Naval Civil Engineering Laboratory Port Hueneme, California 93041 Attn: Mr. W. S. Haynes
1	ONR Branch Office 760 Market Street, Rm. 447 San Francisco, California 94102 Attn: Dr. P. A. Miller	1	Professor O. Heinz Department of Physics & Chemistry Naval Postgraduate School Monterey, California 93940
1	ONR Branch Office 495 Summer Street Boston, Massachusetts 02210 Attn: Dr. L. H. Peebles	1	Dr. A. L. Slafkosky Scientific Advisor Commandant of the Marine Corps (Code RD-1) Washington, D.C. 20380
1	Director, Naval Research Laboratory Washington, D.C. 20390 Attn: Code 6100	1	Office of Naval Research Arlington, Virginia 22217 Attn: Dr. Richard S. Miller
1	The Asst. Secretary of the Navy (R&D) Department of the Navy Room 4E736, Pentagon Washington, D.C. 20350		
1	Commander, Naval Air Systems Command Department of the Navy Washington, D.C. 20360 Attn: Code 310C (H. Rosenwasser)		

TECHNICAL REPORT DISTRIBUTION LIST

<u>No. Copies</u>		<u>No. Copies</u>
	Dr. R. M. Grimes University of Virginia Department of Chemistry Charlottesville, Virginia 22901	1
	Dr. M. Tsutsui Texas A&M University Department of Chemistry College Station, Texas 77843	1
	Dr. C. Quicksall Georgetown University Department of Chemistry 37th & O Streets Washington, D.C. 20007	1
	Dr. M. F. Hawthorne University of California Department of Chemistry Los Angeles, California 90024	1
	Dr. D. B. Brown University of Vermont Department of Chemistry Burlington, Vermont 05401	1
	Dr. W. B. Fox Naval Research Laboratory Chemistry Division Code 6130 Washington, D.C. 20375	1
	Dr. J. Adcock University of Tennessee Department of Chemistry Knoxville, Tennessee 37916	1
	Dr. A. Cowley University of Texas Department of Chemistry Austin, Texas 78712	1
	Dr. W. Hatfield University of North Carolina Department of Chemistry Chapel Hill, North Carolina 27514	1
	Dr. D. Seyferth Massachusetts Institute of Technology Department of Chemistry Cambridge, Massachusetts 02139	1
	Dr. M. H. Chisholm Princeton University Department of Chemistry Princeton, New Jersey 08540	1
	Dr. B. Foxman Brandeis University Department of Chemistry Waltham, Massachusetts 02154	1
	Dr. T. Marks Northwestern University Department of Chemistry Evanston, Illinois 60201	1
	Dr. G. Geoffrey Pennsylvania State University Department of Chemistry University Park, Pennsylvania 16802	1
	Dr. J. Zuckerman University of Oklahoma Department of Chemistry Norman, Oklahoma 73019	1

