Lawrence Berkeley National Laboratory

Lawrence Berkeley National Laboratory

Title

SEASONAL OXYGEN DEPLETION IN CHESAPEAKE BAY

Permalink

https://escholarship.org/uc/item/1f65065c

Author

Taft, Jay L.

Publication Date





Lawrence Berkeley Laboratory UNIVERSITY OF CALIFORNIA, BERKELEY

EARTH SCIENCES DIVISION

Published in ESTUARIES, Volume 3, No. 4, December 1980,pp. 242-247

SEASONAL OXYGEN DEPLETION IN CHESAPEAKE BAY

Jay L. Taft, W. Rowland Taylor, Eric O. Hartwig, and Randy Loftus LAWRENCE

APR 7 1981

LIBRARY AND

January 1981

TWO-WEEK LOAN COPY

This is a Library Circulating Copy which may be borrowed for two weeks. For a personal retention copy, call Tech. Info. Division, Ext. 6782.

Prepared for the U.S. Department of Energy under Contract W-7405-ENG-48

DISCLAIMER

This document was prepared as an account of work sponsored by the United States Government. While this document is believed to contain correct information, neither the United States Government nor any agency thereof, nor the Regents of the University of California, nor any of their employees, makes any warranty, express or implied, or assumes any legal responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by its trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof, or the Regents of the University of California. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof or the Regents of the University of California.

Seasonal Oxygen Depletion in Chesapeake Bay

JAY L. TAFT

W. ROWLAND TAYLOR Chesapeake Bay Institute The Johns Hopkins University Baltimore, Maryland 21218

Eric O. Hartwig

Marine Sciences Group Lawrence Berkeley Laboratory Berkeley, California 94720

RANDY LOFTUS Chesapeake Bay Institute 4800 Atwell Road Shady Side, Maryland 20867

ABSTRACT: The spring freshet increases density stratification in Chesapeake Bay and minimizes oxygen transfer from the surface to the deep layer so that waters below 10 m depth experience oxygen depletion which may lead to anoxia during June to September. Respiration in the water of the deep layer is the major factor contributing to oxygen depletion. Benthic respiration seems secondary. Organic matter from the previous year which has settled into the deep layer during winter provides most of the oxygen demand but some new production in the surface layer may sink and thus supplement the organic matter accumulated in the deep layer.

Introduction

Oxygen depletion zones which approach anoxia deny use of that portion of the ecosystem to most plankton, nekton and benthos. Thus, oxygen depletion in coastal waters and estuaries impacts upon benthic and demersal populations, including commercially important species, and may adversely influence use of these waters by man for power plant cooling, waste disposal and recreation.

Summer deep-water oxygen depletion is common in fjords, in many lakes (Richards 1965), in some estuaries, including Chesapeake Bay (Newcombe and Horne 1938; Carpenter and Cargo 1957; Whaley, et al. 1966; Hobbie, et al. 1972; Taft and Taylor 1976a, 1976b), among the San Juan Islands (Johnson and Thompson 1929) and in parts of the Puget Sound system (Barnes and Collias 1958; Christensen and Packard 1976). However, large scale coastal ocean deoxygenation, such as observed on the Middle

© 1980 Estuarine Research Federation

Atlantic Shelf during the summer of 1976 (IDOE 1976), is not common.

Associated with summer oxygen depletion in Chesapeake Bay are accumulations of soluble reactive phosphorus (SRP) and ammonium. These nutrients are transported vertically to the euphotic zone where they support the annual open-bay primary productivity maximum and phytoplankton biomass maximum. Thus, summer deep-water oxygen depletion is an important facet of the annual nutrient cycle.

There are three processes which could influence deep water oxygen concentration: increased water column stratification during the spring freshet, which minimizes oxygen replenishment from surface to deep waters; increased plankton and benthic respiration rates due to warming in spring; increased organic loading to the deep water due to increased primary productivity in overlying waters during spring. The purpose of this study was to establish the relative impor-

0160-8347/80/040242-06\$01.50/0

Reprinted with permission from L. Harold Stevenson, Estuarine Research Federation, University of South Carolina, Columbia, SC 29208 from ESTUARIES, Vol.3,No.4,December 1980, pp. 242-247.

242

tance of these three processes to seasonal oxygen depletion in Chesapeake Bay.

This study was supported by Department of Energy contract number EY-76-S-02-3279, document number C00-3279-34 and by National Science Foundation grants GA-33445, OCE 76-11399 and OCE 76-01853. Chesapeake Bay Institute contribution number 274. We gratefully acknowledge the assistance of the crews of the R/V MAURY and R/V WARFIELD. We thank Dr. J. Heinbokel for critically reading the manuscript.

Methods

Data from two surveys of Chesapeake Bay (Whaley, et al. 1966; Taylor and Cronin 1974) were augmented with additional data collected during 1977. R/V MAURY cruises in 1964-66 and R/V WARFIELD cruises in 1969-71 and 1977 were made to established Chesapeake Bay Institute stations (Fig. 1). On all cruises a submersible pump was lowered from surface to near bottom to collect samples for profiling temperature and conductivity (Schiemer and Pritchard 1961) and dissolved oxygen (Carpenter 1965). Respiration measurements were made by Winkler titration (Hartwig and Michael 1978) on water collected with a Niskin sterile bag sampler. Precision estimates for replicate standardizations on board ship were $\pm 3.2\%$ and $\pm 0.5\%$ on two cruises. Particulate carbon (PC) was measured by the method of Gordon and Sutcliffe (1973), chlorophyll a by Strickland and Parsons (1972) and adenosine triphosphate (ATP) by luciferase assay after Strickland and Parsons (1972). Primary productivity was measured by ¹⁴C uptake during simulated in situ incubations (McCarthy, et al. 1974).

Results

Volume of Low O_2 Water. Data in Table 1 for August 1970 indicate that the southnorth extent of low oxygen deep-water, defined here as <20% saturated, was about $37^{\circ}44'N$ to $39^{\circ}04'N$ (stations 744 to 904N). The upper boundary of water less than 20% saturated occurred at about 9–10 m depth between $38^{\circ}04'N$ to $39^{\circ}04'N$. Lateral transects along $38^{\circ}04'N$, $38^{\circ}45'N$ and $39^{\circ}04'N$ in July 1977 (Fig. 2) also show the upper boundary for low oxygen water at about 10



Fig. 1. Station locations.

m or less. Therefore, the minimum volume of low oxygen water in mid-summer is estimated at 6×10^9 m³ (Cronin and Pritchard 1975) which is about 8% of the entire Bay volume and 26% of the volume north of 38°04'N.

Rate of Oxygen Depletion. During the three study periods oxygen concentrations were highest in December-February and generally began to decrease in March or April. Fig. 3 shows data for 1977. Minimum values were observed by June in each year except 1965 when the minimum was observed in early July. The rates of change in deep-water oxygen concentrations from the

J. L. Taft, et al.

Station	% O ₂ Saturation in Bottom Water	Approx. Depth of 20% Saturation (m)			
922Y	60.8	musikanika musikanika			
914S	33.1	consector-			
904N	1.1	` 9			
858C	U	9			
845 G	U	10			
834G	· U	9			
818P	4.5	10			
804C	12.8	10			
744	15.5	14			
724R	58.2	evenance			
707Ø	72.9	n an			

TABLE 1. Bottom water oxygen saturation in August 1970. U = Undet ctable. - = 20% saturation not observed.

winter-spring maximum to the summer minimum ranged from $-0.03 \text{ ml} \cdot l^{-1} \cdot d^{-1}$ in 1971 to -0.11 in 1969 (Table 2). Six of the seven values were between -0.06 and -0.11ml $\cdot l^{-1} \cdot d^{-1}$. Thus, when averaged over the same seasons for seven years, oxygen consumption in the deep water exceeded the replenishment rate by 0.08 ml $\cdot l^{-1} \cdot d^{-1}$.

Deep water oxygen depletion rates for Chesapeake Bay are considerably more rapid than for the Puget Sound basins examined by Barnes and Collias (1958) in which average oxygen utilization from winter to summer ranged $0.005-0.020 \text{ ml} \cdot \text{l}^{-1} \cdot \text{d}^{-1}$. Oxygen utilization in Dabob Bay, Washington, was $1.1 \ \mu\text{g-atom} \cdot \text{l}^{-1} \cdot \text{d}^{-1}$ (Barnes and Collias 1958) and $1.25 \ \mu\text{g-atom} \cdot \text{l}^{-1} \cdot \text{d}^{-1}$ (Christen-



Fig. 2. Summer oxygen saturation on three cross bay transects. White = greater than 20% saturation, cross hatch = 0-20% saturation, black = H_2S present.





sen and Packard 1976) in two studies as compared to an average of 7.3 μ g-atom·l⁻¹·d⁻¹ for Chesapeake Bay. Differences in the availability of readily oxidizable organic matter and in the rates of vertical exchange in Dabob and Chesapeake bays were probably the major factors contributing to these observations.

Spring Freshet. Water column density structure in partially mixed estuaries is largely controlled by salinity. Maximum salinity at all depths occurred in January-February in the years studied. During the spring the upper layer freshened from 12-15% to 5-10% at latitude 38°45'N for example, due to runoff from the Susquehanna River. Data for the seven years considered here show decreases in oxygen concentration immediately following the dilution of surface layer salinity by spring river runoff. The spring freshet was characterized by sustained surface salinity decrease averaging 0.07% · d⁻¹ at station 845G during spring of 1965 and 1966, and 0.06% · d⁻¹ in spring of 1970 and 1971. In some of the data, salt was diluted in two steps due to downstream displacement of water at all depths by the fresh water pulse entering the head of the Bay

TABLE 2. Deep water oxygen change at Station 845G measured from winter-spring concentration maximum to summer minimum.

	$\frac{\Delta O_2}{ml \cdot l^{-1} \cdot d^{-1}}$
1964	-0.10
1965	-0.08
1966	-0.06
1969	-0.11
1970	-0.06
1971	-0.03
1977	-0.09

Oxygen Depletion in Chesapeake Bay

	February 1976		April 1976		August 1976			
Station	mg-at C ⁻¹	mg Chl a ⁻¹	mg-at C ⁻¹	mg Chl 4 ^{−1}	mg ATP-1	mg-at C ⁻¹	mg Chl a ⁻¹	mg ATP-1
904N	0.45	4.4	0.54	12	54	0.52	6.4	38
854G						0.75	8.9	33
834G	0.77	9.1	0.43	16	1	0.49	8.4	61
818P			0.36	7	24	0.49	9.8	28
804C	0.71	15.7				1.0	32	64

TABLE 3. Respiration of whole plankton samples measured as oxygen loss expressed in mg-at $O_2 \cdot d^{-1} \cdot column$ heading. Incubation temperatures were 5 °C in February, 15 °C in April and 25 °C in August.

(Schubel 1972). Following initial displacement, seaward flowing fresh water diluted the surface salinity thereby increasing density differences between upper and lower layers of the estuary. This was evident in April 1964, May 1965, March 1966, May 1969 and 1977.

Respiration. Winter respiration rates (February 1976) ranged 0.23-0.89 ml · l⁻¹ · d⁻¹. spring rates (April 1976 and 1977) were 0.26-1.35 ml·l⁻¹·d⁻¹ and summer rates (August 1975 and 1976) were 0.54–1.91 ml·l⁻¹·d⁻¹. These broad ranges reflect differences in temperature, plankton biomass and composition, substrate availability and ambient oxygen concentration. The effects of containment on the natural populations could also contribute to the wide ranges (Venrick, et al. 1977). But, since the worst analytical precision expected is $\pm 3.2\%$ (Hartwig and Michael 1978) the variability in the respiration rate measurements was not due primarily to analytical technique.

Respiration rates normalized to biomass expressed as particulate carbon (PC), chlorophyll a and ATP are shown in Table 3. When normalized to PC, respiration rates were comparable in February, April, and August 1976 at incubation temperatures of 5, 15 and 25 °C respectively. Respiration normalized chlorophyll a ranged 4.4 to 15.7 mg-atom $O_2 \cdot mg$ Chl $a^{-1} \cdot d^{-1}$ in February, 7.0 to 12 mg-atom $O_2 \cdot mg$ Chl $a^{-1} \cdot d^{-1}$ in April and 6.4 to 32 mg-atom O2. mg Chl $a^{-1} \cdot d^{-1}$ in August. Respiration normalized to ATP was 24 and 54 mg-atom $O_2 \cdot mg$ ATP⁻¹·d⁻¹ in April and 28 to 64 mg-atom O₂·mg ATP⁻¹·d⁻¹ in August. No clear evidence for elevation in biomass specific oxygen consumption with temperature increase from February to August was observed.

Comparing water column respiration rates with average deep water oxygen change of $-0.08 \text{ ml} \cdot l^{-1} \cdot d^{-1}$ reveals that the seasonal net loss of oxygen from the deep water is about 10% of the maximum winter respiration rate and 4% of the maximum summer rate. Therefore, in Chesapeake Bay, the imbalance between oxygen flux to the deep water and water column respiration need be sustained at only 10% or less of the respiration rate in order to produce the observed seasonal depletion.

Primary Productivity. Monthly primary productivity data for April 1969 to April 1971 for station 845G and 834G are shown in Fig. 4. Productivity during February to May, the period of most rapid oxygen depletion, ranged 15 to 427 mg $C \cdot m^{-2} \cdot h^{-1}$ at 845G and 53 to 142 mg $C \cdot m^{-2} \cdot h^{-1}$ at 834G. The April 1971 value of 427 mg $C \cdot m^{-2} \cdot h^{-1}$ for 845G was unusually high in comparison to the more commonly observed rates of about 50 to 200 mg $C \cdot m^{-2} \cdot h^{-1}$ in February to May. Productivity increased about threefold from February to May. However, the annual maximum occurred in summer, after oxygen depletion was well underway, or in fall when mixing and reoxygenation were occurring. Therefore, oxygen depletion is not specifically a response to organic loading resulting from phytoplankton productiv-



Fig. 4. Primary productivity at two stations in upper Chesapeake Bay.

ity maxima but may be primarily supported by organic material accumulated in the deep water during the previous fall and winter. Hargrave (1972) found that benthic oxygen demand in Lake Esrom was related to epilimnetic productivity and Smith (1974) suggested that productive surface waters increase metabolism of underlying deep-sea sediments. Deep anoxia layer formation in Tokyo Bay has been related to the sinking of fecal material resulting from copepod grazing on phytoplankton in the euphotic zone (Seki, et al. 1974). Also, oxygen consumption may be closely coupled to primary productivity on the Atlantic Continental Shelf (Rowe, et al. 1975).

Discussion

Chesapeake Bay is a partially mixed estuary with thermohaline stratification through much of the year (Pritchard 1968). The water column density structure regulates advective exchange from the upper to lower layers. The primary factor controlling density structure is salinity which is graded vertically and longitudinally in the Bay. Rates of salinity change with depth and distance are influenced primarily by fresh water inflow. With low river flow during late summer and winter, sea water penetrates along the bottom of the estuary past Baltimore. River flow usually increases sharply between February and May, freshening the surface waters and increasing the vertical salinity gradient. Increased water column stability reduces advective transport of oxygen to deep water. As oxygen renewal declines, upsetting the balance between oxygen consumption and its input rate, biological and chemical utilization continues to reduce dissolved oxygen to very low concentrations. The water temperature increase in spring compounds the imbalance by reducing oxygen solubility, and further stabilizing the water column.

The data summarized here support this scenario but, as yet, a quantitative cause and effect relation between river flow and oxygen depletion rate cannot be established because the relation between river flow and salinity is itself complex. Boicourt (1969) modelled in one dimension the relation between upper Chesapeake Bay salinity distribution and Susquehanna River flow. Even

this relatively simple approach required the use of a separate predictor model to describe changes in the seaward boundary condition with time. Chesapeake Bay's response to a freshet is a function of the Bay's recent history and cannot be linearized or easily predicted. Two dimensional numerical models describing salinity response to river flow have been proposed (Elliott 1976; Wang and Kravits 1980) and may lead to acceptable quantitation of the salinity-river flow relation for Chesapeake Bay.

The wide ranges of biomass specific plankton respiration observed in winter, spring, and summer overlapped and the oxygen concentration declined at low temperatures during February to March 1966, suggesting that temperature alone has little effect on respiration. There needs to be only a 10% imbalance between respiration and replenishment for the daily oxygen loss from deep water to account for the observed oxygen depletion.

The existing data do not permit us to ascertain whether oxygen utilization in the water column or the sediments dominates the loss process, but an estimate can be made. Hargrave (1969) reported a regression of fresh water and marine sediment respiration data from the literature which estimated oxygen consumption at 360 ml $O_{3} \cdot m^{-2} \cdot d^{-1}$ at 10 °C which is within the range of observed values for Puget Sound sediments (Pamatmat 1971) and Atlantic Shelf sediments (Rowe, et al. 1975). Distributing this respiration through 1 m³ of overlying water gives 0.36 ml O₃·1⁻¹·d⁻¹ which is comparable to the minimum April water column consumption we observed. However, the minimum April water column consumption we measured (0.26 ml $O_2 \cdot l^{-1} \cdot d^{-1}$) integrated over a 10 m deep lower layer is 2,600 ml $O_2 m^{-2} \cdot d^{-1}$ or about 7 times the calculated sediment utilization. Thus it appears water column respiration dominates seasonal oxygen depletion in Chesapeake Bay deep water, at least until the oxygen concentration falls below the minimum required for healthy aerobic plankters. Christensen and Packard (1976) similarly found that plankton metabolism accounted for 70-85% of the oxygen depletion in Dabob Bay. Increased primary production in spring contributes oxidizable organic matter to the system, but is not the major factor directly driving oxygen depletion in Chesapeake Bay. Organic material from the previous summer and fall which collects in the deep water during winter is more likely providing the oxygen demand.

LITERATURE CITED

- BARNES, C. A., AND E. E. COLLIAS. 1958. Some considerations of oxygen utilization rates in Puget Sound, J. Mar. Res. 17:68-80.
- BOICOURT, W. C. 1969. A numerical model of the salinity distribution in Upper Chesapeake Bay. Master's Essay. Johns Hopkins University. 59 p.
- CARPENTER, J. H. 1965. The Chesapeake Bay Institute technique for the Winkler dissolved oxygen method. *Limnol. Oceanogr.* 10:141-143.
- , AND D. G. CARGO. 1957. Oxygen requirement and mortality of the blue crab in the Chesapeake Bay. Ches. Bay Inst. Tech. Rept. No. 13.
- CHRISTENSEN, J. P., AND T. T. PACKARD. 1976. Oxygen utilization and plankton metabolism in a Washington fjord. *Estuarine Coastal Mar. Sci.* 4:339–347.
- CRONIN, W. B., AND D. W. PRITCHARD. 1975. Additional statistics on the dimensions of the Chesapeake Bay and its tributaries; Cross section widths and segment volumes per meter depth. Ches. Bay Inst. Spec. Rept. No. 42.
- ELLIOTT, A. J. 1976. A numerical model of the internal circulation in a branching tidal estuary. Ches. Bay Inst. Spec. Rept. No. 54.
- GORDON, D. C., AND W. H. SUTCLIFFE. 1973. A new dry combustion method for the simultaneous determination of total organic carbon and nitrogen in seawater. *Mar. Chem.* 1:231–244.
- HARGRAVE, B. T. 1969. Similarity of oxygen uptake by benthic communities. *Limnol. Oceanogr.* 14:801– 805.
- . 1972. A comparison of sediment oxygen uptake, hypolimnetic oxygen deficit and primary production in Lake Esrom, Denmark. Verh. Internat. Verein. Limnol. 18:134-139.
- HARTWIG, E. O., AND J. A. MICHAEL. 1978. A sensitive Winkler Titrator for respiration measurements. *Envir. Sci. Tech.* 12:712–715.
- HOBBIE, J. E., B. J. COPELAND, AND W. G. HARRISON. 1972. Nutrients in the Pamlico River Estuary, N.C. 1969-1971. Water Resources Research Institute, Univ. N.C. Rept. No. 76.
- IDOE. 1976. Anoxia on the middle Atlantic Shelf during the summer of 1976. Report of NSF Workshop on October 15-16, 1976, in Washington, D.C.
- JOHNSON, M. W., AND T. G. THOMPSON. 1929. The sea water at the Puget Sound Biological Station from September 1927 to September 1928. Pub. Puget Sound Biol. Sta. 7:119-128.
- McCARTHY, J. J., W. R. TAYLOR, AND M. E. LOFTUS. 1974. Significance of nannoplankton in the Chesapeake Bay estuary and problems associated with the

measurement of nannoplankton productivity. Mar. Biol. 24:7-16.

- NEWCOMBE, C. L., AND W. A. HORNE. 1938. Oxygenpoor waters of the Chesapeake Bay. *Science* 88:80– 81.
- PAMATMAT, M. M. 1971. Oxygen consumption by the seabed. VI. Seasonal cycle of chemical oxidation and respiration in Puget Sound. Int. Revue Ges. Hydro. Biol. 56:769-793.
- PRITCHARD, D. W. 1968. Chemical and physical oceanography of the Bay. In: Proceedings of the Governors Conference on Chesapeake Bay. 11-13 Sept., 1968. Session IV. p. 49-74. Wye Institute, Wye, Maryland.
- RICHARDS, F. A. 1965. Anoxic basins and fjords, p. 611-645. In: J. P. Riley and G. Skirrow (eds.), Chemical Oceanography, Academic Press, N.Y.
- Rowe, G. T., C. H. CLIFFORD, K. L. SMITH, AND P. L. HAMILTON. 1975. Benthic nutrient regeneration and its coupling to primary productivity in coastal waters. *Nature* 255:215–217.
- SCHIEMER, E. W., AND D. W. PRITCHARD. 1961. An induction conductivity indicator. Ches. Bay Inst. Tech. Rept. No. 25.
- SCHUBEL, J. R. 1972. The physical and chemical conditions of Chesapeake Bay. J. Wash. Acad. Sci. 62:56-87.
- SEKI, H., T. TSUJI, AND A. HATTORI. 1974. Effect of zooplankton grazing on the formation of the anoxic layer in Tokyo Bay. *Estuarine Coastal Mar. Sci.* 2:145-151.
- SMITH, K. L. 1974. Oxygen demands of San Diego trough sediments: An in situ study. Limnol. Oceanogr. 19:939-944.
- STRICKLAND, J. D. H., AND T. R. PARSONS. 1972. A practical handbook of sea water analysis. 2nd ed. Bull. Fish. Res. Board Can. 167.
- TAFT, J. L., AND W. R. TAYLOR. 1976a. Phosphorus distribution in the Chesapeake Bay. *Chesapeake Sci.* 17:67–73.
- ———, AND ———. 1976b. Phosphorus dynamics in some coastal plain estuaries. *In*: M. Wiley (ed.), Estuarine Processes. Vol. 1. Academic Press, New York.
- TAYLOR, W. R., AND W. B. CRONIN. 1974. Station data, AESOP cruises. April 1969 to April 1971. Ches. Bay Inst. Special Report 38. 228 p.
- VENRICK, E. L., J. R. BEERS, AND J. F. HEINBOKEL. 1977. Possible consequences of containing microplankton for physiological rate measurements. J. Exp. Mar. Biol. Ecol. 26:57-76.
- WANG, D. P., AND D. W. KRAVITS. 1980. A semiimplicit two-dimensional model of estuarine circulation. J. Phys. Oceanogr. 10:441-454.
- WHALEY, R. C., J. H. CARPENTER, AND R. L. BAKER. 1966. Nutrient data summary 1964, 1965, 1966: Upper Chesapeake Bay (Smith Point to Turkey Point), Potomac, South, Severn, Magothy, Back, Chester, and Miles rivers; and Eastern Bay. Ches. Bay Inst. Spec. Rept. 12.

This work was partially supported by the U.S. Department of Energy under Contract W-7405-ENG048.