



United States
Department of the Interior
U. S. GEOLOGICAL SURVEY
345 Middlefield Road, MS #496
Menlo Park, California, USA 94025
Phone: (650) 329-4588 Fax: (650) 329-4327
<http://sfbay.wr.usgs.gov/access/wqdata/>

August 10, 2012

To: 2012 Brown-Nichols Science Award Committee

From: Lisa Lucas and Tara Schraga

Please see attached nomination package for Dr. James E. Cloern, containing:

- 1) Nomination Letter (Lucas & Schraga)
- 2) Cloern CV and Bibliography
- 3) Cloern & Jassby, "Drivers of Change in Estuarine-Coastal Ecosystems: Discoveries from Four Decades of Study in San Francisco Bay" (In press, Reviews of Geophysics)
- 4) Luoma Letter of Support
- 5) Jassby Letter of Support
- 6) Wolfe Letter of Support
- 7) Brown Letter of Support
- 8) Connor Letter of Support
- 9) Dettinger & Cayan Letter of Support

(Letters of Support submitted via separate upload function.)

Please do not hesitate to contact us with any questions. Many thanks for your consideration.



United States
Department of the Interior
U. S. GEOLOGICAL SURVEY
345 Middlefield Road, MS #496
Menlo Park, California, USA 94025
Phone: (650) 329-4588 Fax: (650) 329-4327
<http://sfbay.wr.usgs.gov/access/wqdata/>

August 10, 2012

Dear 2012 Brown-Nichols Science Award Committee Members,

It is with great pleasure that we nominate James E. Cloern for the Delta Science Program's Brown-Nichols Science Award. The nominators are fortunate to have been members of Dr. Cloern's research group at the U.S. Geological Survey for the past 15 years. Dr. Cloern's 36 year career has been spent 1) conducting significant scientific research in the San Francisco Bay-Delta and Watershed, 2) making significant scientific contributions to the development of management strategies and policies, and 3) facilitating good science practices by others through management, mentoring, and collaboration. In the following, we aim to demonstrate the breadth, depth, and impact of Dr. Cloern's past and continued contributions, and why he is most deserving of the Brown-Nichols Science Award.

High-Impact Research on the San Francisco Bay-Delta-Watershed

Dr. Cloern's research and publication record is of a quality, quantity, diversity, and impact that few working in our San Francisco Estuary system have achieved. In his more than 130 publications (94 peer-reviewed), Dr. Cloern has addressed an impressive breadth of estuarine science topics including: primary production, phytoplankton community ecology, ecosystem metabolism, carbon biogeochemistry, resource limitation of algal growth, benthic grazing, zooplankton dynamics, disturbance by introduced species, environmental toxicology, impacts of climatic and hydrologic variability, nutrient regeneration, coastal eutrophication, hydrodynamics and mixing processes, and ocean-estuary-watershed coupling. As one measure of the broad impact of Dr. Cloern's research, his peer-reviewed publications alone have been collectively cited more than 6200 times¹.

Dr. Cloern employs a broad toolkit of scientific approaches, including implementation of intense short-term field campaigns, sustained long-term monitoring, numerical modeling, laboratory experimentation, and remote sensing to learn about our coastal ecosystem. Of his 103 publications pertaining to the San Francisco Bay-Delta, we highlight a few that have transformed the way scientists conceptualize both our ecosystem and global estuarine ecological processes. These publications have provided motivation and a framework for numerous studies of San Francisco Bay and other estuaries around the world.

Dr. Cloern's 1982 paper "Does the benthos control phytoplankton biomass in San Francisco Bay (USA)?" (465 citations) was one of the first to systematically demonstrate that grazing by benthic bivalves can control phytoplankton biomass in aquatic systems. This publication transformed the way Bay Area scientists viewed the ecological function of this estuary, and similarly revolutionized the conceptual models of estuarine functionality for scientists around the world. Alpine and Cloern (1992, 237 citations) built on this work to show the impact of invasive benthic filter feeders on seasonal

variability of phytoplankton biomass and productivity. These papers formed a basis for understanding “top-down” controls that 1) help explain how the Bay works and how it continues to change, and 2) bear directly on current concerns in the Delta.

Dr. Cloern and his collaborators have revealed critical physical processes and their interactions with biological processes in San Francisco Bay. He built a first-generation numerical model of phytoplankton dynamics and turbulent mixing in an estuarine water column and demonstrated how, through interactions with turbidity and benthic grazing, the rate of vertical mixing can control phytoplankton dynamics (Cloern 1991, 127 citations). This and his early recognition of the importance of density stratification for phytoplankton bloom development in South San Francisco Bay (Cloern 1984), formed the basis for several more modeling and field studies of physical-biological linkages operating in the vertical dimension (e.g. Koseff et al. 1993, Lucas et al. 1998, May et al. 2003). His early work on shoal-channel exchange (Cloern and Cheng 1981) provided motivation for later research examining horizontal transport and habitat connectivity in the Bay (e.g. Huzzey et al. 1990, Lucas 1997, Lucas et al. 1999a/b, 2009, Thompson et al. 2008) and Delta (Lucas et al. 2002, Lopez et al. 2006, Lucas et al. 2006). Inspired by questions regarding the influence of habitat connectivity on productivity in the Delta, Dr. Cloern developed a general model of a productive (autotrophic) habitat connected to an unproductive (heterotrophic) habitat; he showed that dispersive connectivity between such disparate habitats can amplify overall system production in metazoan food webs (Cloern 2007).

Other recent work has highlighted the importance of searching outside the San Francisco Estuary to understand the dynamics within. In their 2005 paper (*Geophys. Res. Letters*), Dr. Cloern and his co-authors described an unprecedented event within our estuary—a summertime red tide in South and Central San Francisco Bay. They showed that this event was seeded by a coastal dinoflagellate bloom generated by upwelling anomalies in the neighboring coastal ocean. In a 2007 publication (*Proc. Nat. Acad. Sci.*), he and co-authors revealed their discovery of a trophic cascade that began with record-high abundances of marine shrimp and juvenile flatfish and crab in the Pacific and resulted in increased phytoplankton biomass and the appearance of autumn blooms within San Francisco Bay. In the 2010 follow-up publication (*Geophysical Research Letters*), he and his co-authors detail the state change in climate and oceanic forcing across the Pacific Ocean that underlay the trophic cascade. These papers have been exceptionally important not only in demonstrating the criticality of the estuary-ocean connection, but also in alerting the community to the trend of increasing phytoplankton biomass within San Francisco Bay.

The San Francisco Bay-Delta system has long functioned as Dr. Cloern’s laboratory, and he has used his locally learned lessons to illuminate the functioning of estuaries around the world. His review of coastal phytoplankton bloom dynamics (Cloern 1996, 296 citations) synthesizes what he and colleagues discovered through decades of local research. Knowing that one of the best ways to learn about our “backyard” ecosystem is through comparing it to others, Dr. Cloern has performed cross-cutting comparisons of San Francisco Bay and numerous other estuaries (e.g., Cloern 1987b, Cloern 2001). For example, he conceived and organized the first international conference on multi-decadal records of phytoplankton variability across the world’s coastal ecosystems (AGU Chapman Conference, Rovinj, Croatia, 2007). This conference led to a 2010 special issue in the journal *Estuaries & Coasts* and three lead- or co-authored publications synthesizing patterns of phytoplankton variability across the world’s coastal ecosystems (Cloern and Jassby 2008, 2009) and across lake, coastal, and ocean environments (Winder and Cloern, 2010). The groundbreaking 2008 paper documented the surprising absence of any canonical phytoplankton seasonal patterns and, instead, the presence of a broad continuum of seasonal patterns across the world’s coastal ecosystems: annual algal biomass peaks can

occur during any time of year, in both hemispheres. Dr. Cloern's most highly cited publication "Our evolving conceptual model of the coastal eutrophication problem" (Mar. Ecol. Prog. Ser. 2001, 835 citations) has become a definitive resource on the subject for researchers, managers, educators, and students. Following this important contribution to the coastal ecology literature, Dr. Cloern was asked to co-author the entry on "Eutrophication" for the online Encyclopedia of Earth (<http://www.eoearth.org/article/Eutrophication>).

The frequency and importance of Dr. Cloern's publications, if anything, increase over time. As he parlays his decades of study toward continued and increasingly more significant scientific contributions, he is helping the San Francisco Estuary community better understand and manage our ecosystem. He lead-authored the paper synthesizing the first phase of the CASCaDE (Computational Assessments of Scenarios of Change for the Delta Ecosystem, <http://sfbay.wr.usgs.gov/cascade/>) project (PLoS ONE 2011; over 5000 page views and almost 1400 downloads since publication less than one year ago). This paper is the result of Dr. Cloern's leadership and persistence in wrangling multiple disparate model-generated datasets into digestible, rigorously derived environmental indicators of our Estuary-Watershed's possible responses to climate change in the coming century. Among the many significant findings presented, this study highlighted 1) how hydroclimatic events considered extreme by today's standards will become much more common in the future; and 2) the inevitability of changes in our system's biological communities due to shifts in key environmental variables. As one measure of the broad interest in and impact of this paper, Dr. Cloern participated in numerous radio, television and press interviews when it was first published. This hyper-disciplinary effort is a testament to Dr. Cloern's rare gift for extreme synthetic thinking.

Since 1976, Dr. Cloern has sustained one of the longest running programs of water quality research and monitoring in the United States. With the advent of the internet, he was determined to make these data easily accessible to all, creating the U.S.G.S. Water Quality of San Francisco Bay website (<http://sfbay.wr.usgs.gov/access/wqdata>). The website's user-configurable database now contains 43 years of chemical, biological, and ecological data collected at 45 current and historical stations on cruises occurring every month. These data are of the highest quality due to Dr. Cloern's rigorous standards for thorough data collection, precise analysis, and meticulous quality control. In 2011, the website had 334-thousand hits and almost 10-thousand database queries from national, state and local government agencies, research institutions, schools, non-profit groups, and private industry, in addition to institutions in over 40 other countries. For the past 18 years, this monitoring has been conducted in cooperation with the San Francisco Estuary Institute and represents one component of the San Francisco Bay Regional Monitoring Program for Trace Substances mandated by the Regional Water Quality Control Board.

Results Dr. Cloern has derived from this extensive data collection have contributed considerably to our understanding of the San Francisco Estuary and led to discoveries that could only be revealed through decadal scale observation (e.g. Cloern 1991, Alpine and Cloern 1992, Cloern and Dufford 2005, Cloern et al. 2005, 2007, 2010, to name a few). In addition to the numerous papers, reports, and chapters authored by Dr. Cloern, data from this website have been included in a minimum of 25 other peer reviewed publications and 20 Ph.D. and M.S. dissertations (that we know of) in the past decade alone. It is easy to comprehend the significance of Dr. Cloern's program as we continue to require these data for separating the effects of global climate change from the responses to regional natural and anthropogenic drivers in the San Francisco Bay area.

This remarkable dataset represents Dr. Cloern's commitment to long-term data collection in the San Francisco Estuary, even when funding waned and other research projects competed for resources. Dr.

Cloern has persistently and patiently educated agency leaders, academicians, funders, and politicians on the value of and need for continued long-term monitoring in San Francisco Estuary and beyond. In 1985, Drs. Cloern and Nichols wrote in their preface to Temporal Dynamics of an Estuary – San Francisco Bay:

“Ideally, we would like to understand how all components [of the Estuary] change over all time scales. In reality, our understanding in some areas is limited by the lack of comprehensive, long-term studies and/or the relative difficulty in achieving understanding of all the intricate interrelations among components of the estuarine system.”

Because Dr. Cloern is so dedicated to the cause of continuous observations in our system, he is already working to insure their future beyond his tenure at the U.S.G.S. For three years now, he has been working with public and private researchers, managers, stakeholders, and policy makers, discussing strategies for regional and local funding for future water quality monitoring of the Bay. It is understood by these entities that Dr. Cloern’s program has historically relieved the state and local authorities from needing to monitor fundamental water quality in San Francisco Bay.

A very recent paper (Cloern and Jassby, Rev. Geophys., In press) represents perhaps Dr. Cloern’s crowning achievement. This manuscript is the culmination of his nearly four decades of research in this ecosystem. In it, Drs. Cloern and Jassby analyze 40+ years of collected data and discuss them in the context of six primary drivers of change. The authors describe how the Estuary has responded to these agents of change and demonstrate that continuous monitoring enabled detection and projection of ecosystem changes, which then catalyzed key policy responses. One can summarize this manuscript as the major “lessons learned” from 4 decades of observations and study in the San Francisco Bay-Delta. We expect this paper will become essential reading for anyone working in, studying, or managing the San Francisco Estuary-Watershed. We have included a preprint of this manuscript in our nomination package, as it is not yet publicly available.

Significant scientific contributions to the development of management strategies and policies

Dr. Cloern is dedicated to performing objective, relevant research that informs management of and decision making for our Estuary and Watershed. The long-term water quality monitoring program described above has provided a “scientific basis for prioritizing management actions so that regulatory efforts can identify and focus on pollutants posing the greatest threats...” (Cloern et al. 2003). By documenting the disappearance of summer phytoplankton blooms in Suisun Bay and the five-fold reduction in primary production following invasion by the clam *Potamocorbula amurensis* (Alpine and Cloern 1992), the monitoring program presented a direct measure of the ecological impact caused by this alien species. The California Regional Water Quality Control Board cited this as evidence in its TMDL decision to put exotic species on the 303(d) list: “The ecological impacts of the *Potamocorbula* invasion are as dramatic as any documented in the literature, and provided the impetus for the Board’s 303(d) listing decision.”²

The long-term monitoring program documented trends in dissolved oxygen in the Bay before and after regulation of wastewater inputs by the 1972 Federal Clean Water Act, revealing steadily increasing dissolved oxygen and near elimination of low oxygen conditions in response to investments in advanced wastewater processes (Cloern et al. 2003). These data provided strong support for a regulatory decision to remove San Francisco Bay from the list of California water bodies impaired by low oxygen. Dr. Cloern’s analysis of long-term data has further shown that, unlike in the Chesapeake Bay, San Francisco Bay’s comparably high nutrient loads have not yet created excessive blooms because

“resistive mechanisms” such as consumption by clams, turbidity, and turbulent mixing have limited the conversion of nutrients into phytoplankton standing stocks (Cloern 2001). This taught the community why we cannot expect a “one-size-fits-all” cross-system rule to govern coastal ecosystem response to nutrient loading. More recently, Dr. Cloern’s ongoing analysis of the long-term dataset and consideration of other colleagues’ recent findings (see Schoellhamer 2009³) show that turbidity and grazing pressure are changing in ways that could make the system less resilient to nutrient enrichment. Dr. Cloern was the first to raise the alarm that the Bay could become more sensitive to nutrient enrichment in the future. He has organized meetings and colloquia with entities concerned with Bay water quality and wastewater inputs, and he is working with the State Water Board, the Southern California Coastal Water Research Program (SCCWRP) and the San Francisco Estuary Institute (SFEI) to develop nutrient criteria for the San Francisco Bay as part of the California initiative to address concerns of over-enrichment.

Under Dr. Cloern’s leadership, the long-term monitoring program additionally provides management-relevant information through: 1) documenting changes in phytoplankton dynamics, which affect the rates of transformation and bioavailability of contaminants to upper trophic levels; 2) measuring changes in Bay salinity as an indicator of shifts in river flow, contaminant inputs and metal availability for uptake by aquatic organisms; 3) establishing baseline information to compare against future changes in our Bay such as salt pond restoration, climate change, and sea level rise.

Dr. Cloern has generously contributed his expertise to the South San Francisco Bay Salt Pond Restoration Program. He has served as an advisor to managers and administrators on issues such as designing relevant monitoring programs, and has materially supported other groups in collecting high quality data, even in the absence of available funding. He funded the creation of an inventory of water quality data collected in and around South Bay salt ponds and sloughs to enhance data access for researchers. Dr. Cloern recruited a post-doc to estimate and document primary production for the first time in the decommissioned salt ponds. This study’s findings are a significant contribution to the adaptive management of the ponds, demonstrating the ponds’ food supply benefit along with risks of hypoxia and harmful algal blooms (HABs) (Thebault et al., 2008). This study was the first to discover the prevalence of HAB species in the ponds, and motivated Dr. Cloern’s preliminary study aimed at understanding the potential threat HABs pose to the greater estuarine system. Historically, San Francisco Bay has not sustained HAB episodes, but Dr. Cloern’s research has revealed that in the past decade a number of species previously undetected in the system have maintained significant and potentially deleterious populations. In response to this threat, he has initiated a year long project in collaboration with Prof. Raphael Kudela (UC Santa Cruz) measuring algal toxins on all of the regular monitoring cruises in San Francisco Bay. This comprehensive phytoplankton toxin sampling is the first of its kind in the estuary. The results will answer key questions regarding the threat of various HAB species to organisms that live in and around the Bay.

Dr. Cloern has made numerous scientific contributions to the management of the Delta. Organic carbon work performed under his project leadership and CALFED supported collaborations informed managers of the revelation that phytoplankton are the key, and often limiting, food source for the Delta’s planktonic food web (Sobczak et al. 2002, 2006; Mueller-Solger et al. 2002; Jassby and Cloern 2000; Jassby et al. 2002). As part of those endeavors, the first cross-disciplinary studies of *Corbicula fluminea* in the Delta were conducted and established this freshwater clam as a significant limit on the availability of quality organic carbon to pelagic consumers in parts of the Delta (Lucas et al. 2002; Lopez et al. 2006); also, the effect on Delta primary productivity of the estuarine clam *P. amurensis* in the western Delta was assessed (Jassby, Cloern and Cole 2002). These studies called attention to and

quantified the “phytoplankton” and “clam” issues that are now widely understood as matters of consequence for the Delta’s future and play prominently in ecosystem management planning. Other work conducted under Dr. Cloern’s mentorship describes model-based case studies of Delta water operations, demonstrating how local diversions can change Delta-scale circulation patterns, flushing times, water source mixtures, and water quality (Monsen et al. 2007). Dr. Cloern’s modeling study of the influence of habitat connectivity on ecosystem productivity (Cloern 2007) is pertinent to current discussions within the Delta community regarding habitat restoration, levee stability, and ecological implications of transport parameters such as residence time and flushing time. This paper is frequently cited in documents and manuscripts concerned with Bay-Delta science and management.

More than nine years ago, Dr. Cloern conceived of the visionary CASCaDE project to provide science based information to resource managers who face increasingly tough choices as they search for strategies to stabilize water supplies, provide safe drinking water, and sustain native species while contending with key forces of change such as climate, sea level rise, water diversion, and land use. Dr. Cloern organized a 2003 public workshop to solicit comments and suggestions from citizens, resource managers, and stake holders on the proposed project. He assembled a diverse team of researchers specializing in climate, hydrology, hydrodynamics, phytoplankton ecology, benthic ecology, sediment dynamics, geomorphology, toxic contaminants, and fish population dynamics to develop a series of linked models and skillfully guided the group to maximize cohesion and relevance to decision makers. CASCaDE is an ambitious and complicated project assessing how the Delta system might respond under a range of future scenarios, and has already contributed results and several publications valuable to California’s decision makers as they contend with the next century of anticipated change (see CASCaDE publications at <http://cascade.wr.usgs.gov/publications.shtm>). The primary CASCaDE synthesis paper (Cloern et al., PLoS ONE 2011, described above) was geared from the start to provide resource managers and planners integrated, useable scenarios of how the changing climate can affect water quality and availability, aquatic communities, and the environment for humans. This paper is being cited in reports and manuscripts concerning management of the Bay-Delta and other coastal ecosystems. It was highlighted in a report by the Delta Science Program’s lead scientist, who stated: “The resulting synthesis paper serves as an example of integration among multiple disciplines needed to inform management decisions. The analyses presented in the paper will be useful for guiding the development of performance measures for the Bay-Delta-River system.”

In 2010, Dr. Cloern gave an invited presentation to the National Academies’ *Committee on Sustainable Water and Environmental Management in the California Bay-Delta* in Sacramento. In his talk entitled “*Historical perspective on human disturbance in the Sacramento-San Joaquin Delta Ecosystem*”, he brought to the prestigious panel his decades of Bay-Delta experience and broad understanding of interlinked physical, biological, and chemical changes in the system to provide a multi-dimensional perspective of the many stressors that have been acting on, and transforming, the Bay-Delta for decades. He has served as advisor or invited committee member on many Bay-Delta issues including, for example: CA Coastal Conservancy/South Bay Salt Pond Restoration, National Center for Ecological Analysis and Synthesis POD Working Group, Bay Area Ecosystems Climate Change Consortium, BCDC-NOAA-NMFS Subtidal Habitat Goals, CALFED Environmental Water Account, NOAA SFO Runway Expansion, CA Coastal Conservancy-NOAA SF Bay Modeling Needs & Strategies, CALFED-Delta Vision Blue Ribbon Task Force. Dr. Cloern recently provided a Capitol Hill (Washington D.C.) briefing on the value of coastal long-term monitoring for staff members to the U.S. Senate Committee on Environment and Public Works, Office of Management and Budget, House Science Committee, and Congressional Research Service. Dr. Cloern has enthusiastically agreed on two occasions to present

plenary talks at CALFED Science Conferences (2003, 2008).

Facilitation of good science practices by others through management, mentoring, and collaboration

Regarded as a world expert in estuarine ecology, Dr. Cloern is sought after as an advisor to scientists in the Bay Area and around the world. For emerging scientists, he is generous with his time and knowledge. He has mentored 12 post-doctorates conducting research in the San Francisco Bay and Delta, all of whom on completion of their post-docs went to faculty or scientific positions in government agencies, academic institutions, or private consulting. A few high impact outcomes of Dr. Cloern's mentorship are: Canuel et al. 1995, Lucas et al. 1998, 1999a/b, Sobczak et al. 2002, 2006, Mosen et al. 2002, May et al. 2003, Lopez et al. 2006; these nine publications have over 550 citations combined. In the past 20 years he has sat on 20 graduate thesis committees and has mentored 4 high school interns. Dr. Cloern has committed to serve as community mentor for several young applicants for CALFED/DSP fellowships. He provides a great deal of latitude to the younger scientists under his own direction, encouraging everyone including his lab technicians to take the lead in publishing and giving public presentations. Dr. Cloern's research group members consistently present at CALFED/DSP Science Conferences. He often reminds us of the investment in our research from funding agencies and taxpayers and our responsibility to provide ample scientific and educational returns on those investments. He regularly invites and welcomes graduate students, post-docs, and visiting scientists to conduct their own research aboard monthly U.S.G.S. Research Vessel Polaris water quality cruises.

Dr. Cloern further shares his professional experience by instructing the young scientific community at large; his tutorial seminar entitled "A Brief Guide to Scientific Writing and Publishing" originally given in 2006 is available as a video presentation via his website at <http://sfbay.wr.usgs.gov/access/wqdata/presentations/>. He also presented a short course on the same topic at the Universite de Bretagne Occidentale (2006), two (Coastal and) Estuarine Research Federation Conferences (2007, 2009), Woods Hole Oceanographic Institution (2010), and most recently at University of Maryland Horn Point Laboratory, where he spent an invited week teaching and interacting with students as the 2012 "Scholar in Residence." (He was chosen for this honor by Horn Point graduate students, in consultation with faculty.) Dr. Cloern regularly gives invited lectures at universities around the world, to the intense interest and appreciation of students and established scientists alike.

Realizing that the challenging environmental questions facing our Bay-Delta require interdisciplinary science, Dr. Cloern has forged many productive science collaborations and built and led several multi-disciplinary research teams. For example, he has provided invaluable leadership to three large CALFED/DSP supported multi-disciplinary projects resulting in findings and publications critical to understanding ecological function of the Delta: 1) 1999-2002 study on Sources and Quality of Organic Carbon; 2) 2001-2005 Carbon-Selenium-Transport Project; 3) 2006-2009 CASCaDE. His collaborations in the San Francisco Bay-Delta have transcended the federal government, reaching to state agencies and academic institutions (e.g. CA Departments of Fish and Game and Water Resources, Stanford University, UC Berkeley, UC Davis, UC Santa Cruz, UC Santa Barbara, San Francisco State University, State University of New York, Virginia Institute of Marine Science, Universite de Bretagne Occidentale (France), Stazione Zoologica Anton Dohrn (Italy)).

One of Dr. Cloern's most fruitful collaborations has been with Dr. Alan Jassby (UC Davis). Together, they have published seminal analyses of Delta organic carbon dynamics (e.g., 2000, 2002) that

are heavily cited by Delta scientists and planners. They also recently developed and documented a suite of tools for analyzing water quality time series data in “R” (Jassby and Cloern, 2010; <http://cran.r-project.org/web/packages/wq/index.html>). This suite is publically available at no cost, and Dr. Cloern actively mentors others in using this tool suite with his San Francisco Bay-Delta dataset, encouraging them to discover new trends.

Dr. Cloern spearheaded the creation of the joint USGS-CALFED Bay-Delta Authority video documentary “Delta Revival: Restoring a California Ecosystem” (<http://gallery.usgs.gov/videos/443#.UCQmjaMXKU9>), winner of a 2003 Golden Screen Award from the National Association of Government Communicators and a 2004 First Place Silver Telly Award. This video describes the collaborative science Dr. Cloern led in support of ecosystem restoration and was broadcast in August 2003 by the San Francisco CBS television affiliate. Dr. Cloern has provided hundreds of DVD copies to teachers, professors, journalists, researchers and government agencies in the U.S. and beyond. He has served as Associate Editor for several scientific journals (including *San Francisco Estuary and Watershed Science*) and as Co-Editor-in-Chief for the journal *Estuaries and Coasts* (2007-2012), providing additional examples of Dr. Cloern’s generosity in sharing his experience and knowledge in order to facilitate the sound scientific practices of others.

A Career of San Francisco Estuary and Sacramento-San Joaquin Delta Stewardship

Jim’s love of the San Francisco Estuary transcends academic and institutional walls — it permeates all aspects of his life. He routinely bikes to work across San Francisco Bay and takes daily walks along its fringes and sloughs. We know he draws scientific inspiration from these activities, the tangible results of which are his creative scientific contributions. Jim has devoted his nearly four decade career to increasing understanding of the San Francisco Bay-Delta-Watershed to support its scientifically informed management. Along the way, he has provided an example to many young scientists of how to work with focus and integrity. Through his research based in our Bay-Delta, he has become a locally and internationally recognized leader in estuarine research, teaching valuable lessons to scientists and managers around the world. Jim embodies both the definition and the spirit of this prestigious award honoring the achievements and stewardship of Randy Brown and Fred Nichols. We sincerely thank the committee for considering this nomination of Dr. James Cloern for the Delta Science Program’s Brown-Nichols Science Award.

Sincerely,



Lisa V. Lucas, Research Engineer
U.S. Geological Survey



Tara Schraga, Oceanographer
U.S. Geological Survey

¹ Citation tallies were obtained from Web of Knowledge whenever possible and from Google Scholar otherwise.

² California Regional Water Quality Control Board, San Francisco Region. Prevention of Exotic Species Introductions to the San Francisco Bay Estuary: A Total Maximum Daily Load Report to U.S. EPA. May 8, 2000.

³ Suspended sediment in the Bay: Past a tipping point, in: *The Pulse of the Estuary: Monitoring and managing water quality in the San Francisco Estuary*. 2009. San Francisco Estuary Institute, Oakland, CA.

BIOGRAPHICAL SKETCH - JAMES E. CLOERN

Updated 1 June 2012

EDUCATION

University of Wisconsin-Madison, B.S. 1970, Zoology
University of Wisconsin-Milwaukee, M.S. 1973, Zoology (fisheries biology, limnology)
Washington State University, Ph.D. 1976, Zoology (limnology, ecological modeling)

RESEARCH AND PROFESSIONAL EXPERIENCE

1999-present Senior Research Scientist, U.S. Geological Survey, Menlo Park, CA
1997-present Consulting Professor, Stanford University, Department of Civil & Env. Engineering
2005 Chercheur Associé, Centre National de la Recherche Scientifique, Brest, France
2005 Instructor, Ecole Doctorale, Université de Bretagne Occidentale, Brest, France
2000 Visiting Scientist, National Environmental Research Institute, Roskilde, Denmark
1976-1999 Research Scientist, U.S. Geological Survey, Menlo Park, CA
1997-1998 Lecturer, University of California-Santa Cruz, Department of Earth Sciences
1996 Visiting Lecturer, Second Institute of Oceanography, Hangzhou, China
1993 Distinguished Visiting Scientist, National Institute of Water and Atmospheric Research, Hamilton, New Zealand
1993-1994 Directeur de Recherche, Université d'Aix-Marseille II, France

RESEARCH INTERESTS

Comparative ecology and biogeochemistry of estuaries, directed to understand how they respond as ecosystems to climatic-hydrologic variability and human disturbance. Research is focused around a long-term (36 year) team investigation of San Francisco Bay that has included study of primary production, nutrient cycling, algal and zooplankton community dynamics, ecosystem metabolism and food web dynamics, disturbance by introduced species, Bay-Ocean connectivity, ecosystem restoration, and projected responses to climate change.

AWARDS AND HONORS

Ian Morris Scholar in Residence, Horn Point Laboratory, University of Maryland Center for Environmental Science, 2012 <http://www.umces.edu/hpl/ian-morris-scholar-residence>
B.H. Ketchum Award, Woods Hole Oceanographic Institution, 2010 <http://www.whoi.edu/page.do?pid=7545&tid=3622&cid=72786>
Laura Randall Schwappe Endowed Lecturer, University of Texas Marine Science Institute, 2009
Fulbright Senior Specialist, 2008-present
Golden Screen Award, National Association of Government Communicators, 2003
U.S. Department of Interior Distinguished Service Award, 2000
USGS Shoemaker Communications Award, 1998
Fulbright Research Scholar, 1993-94
NATO Collaborative Research Grant Award, 1993
U.S. Department of Interior Meritorious Service Award, 1991

POSTDOCTORATES:

Dr. Linda Huzzey, Professor of Marine Science, U.S. Coast Guard Academy
Dr. Richard L. Miller, Research Scientist, NASA Stennis Space Center
Dr. Jane Caffrey, Assistant Professor, Center for Bioremediation and Diagnostics, University of West Florida
Dr. Joseph Rudek, Senior Scientist, Land, Water and Wildlife Programs, Environmental Defense Fund
Dr. Elizabeth Canuel, Professor of Marine Science, Virginia Institute of Marine Science

Dr. Lisa Lucas, Research Engineer, U.S. Geological Survey
Dr. William Sobczak, Associate Professor of Environmental Studies, College of Holy Cross
Dr. Jean-Marc Guarini, Professor of Oceanography, Université Pierre et Marie Curie (Paris VI)
Dr. Laurent Chauvaud, Directeur de Recherche, Institut Universitaire Européen de la Mer, Brest, France
Dr. Nancy Monsen, Research Associate, Dept. Civil & Environmental Engineering, Stanford University
Dr. Julien Thébault, Lecturer, Université de Bretagne Occidentale, Brest, France
Dr. Rochelle Labiosa, Research Scientist, Pacific Northwest National Laboratory, Seattle

THESIS/DISSERTATION COMMITTEES:

Bertrand Millet, Université d'Aix-Marseille II, France, 1992 (Habilitation)
Anne Lefebvre, Université de Montpellier II, France, 1993 (MS)
Kedong Yin, University of British Columbia, Canada, 1993(PhD)
Frederic Jean, Université de Bretagne Occidentale, Brest, France, 1994 (PhD)
Olivier Ragueneau, Université de Bretagne Occidentale, Brest, France, 1994 (PhD)
Nabila Mazouni, Université d'Aix-Marseille II, France, 1995 (PhD)
Jody Edmunds, Moss Landing Marine Laboratories, 1997 (MS)
Lisa Lucas, Stanford University, 1997 (PhD)
Laurent Chauvaud, Université de Bretagne Occidentale, Brest, France, 1998 (PhD)
Anne Lorrain, Université de Bretagne Occidentale, Brest, France, 2002 (PhD)
Christopher Carter, University of Canterbury, New Zealand, 2004 (PhD)
Christine May, Stanford University, 2005 (PhD)
Stephanie Moore, University of New South Wales, Australia, 2005 (PhD)
Olivier Ragueneau, Université de Bretagne Occidentale, Brest, France, 2005 (Habilitation)
Christel Pinazo, Université d'Aix-Marseille II, France, 2005 (Habilitation)
Laurent Chauvaud, Université de Bretagne Occidentale, Brest, France, 2005 (Habilitation)
Sophie Martin, Université de Bretagne Occidentale, Brest, France, 2005 (PhD)
Julien Thebault, Université de Bretagne Occidentale, Brest, France, 2005 (PhD)
Diana Escobedo, Centro Interdisciplinario de Ciencias Marinas, La Paz, Mexico, 2007 (PhD)
Eric Goberville, Université de Bordeaux I, Bordeaux, France, 2010 (PhD)

ADVISORY SERVICE (examples):

Member, Florida Bay Science Oversight Panel, 1996-1998
Advisor, Swan River Science Program, Perth, Australia, 1998
Committee of International Experts, Programme Rade de Brest, France, 1998-2000
Panelist, NSF Cooperative Activities in Environmental Research between the National Science Foundation and the European Commission, 2003
Evaluator, Heinz Center's *State of the Nation's Ecosystems*, 2002-2003
Advisor, California Coastal Conservancy, South Bay Salt Pond Restoration Project, 2004
Advisor, USEPA STAR Grant Program, 2004
Panelist, NOAA Coastal Ocean Research, Ecological Forecasting, 2004
Advisor, USGS Tampa Bay Studies Program, 2004-2005
Advisor, USEPA National Nutrient Criteria Program, 2005-2006
Chesapeake Bay Program, Scientific Advisory Committee Workshop, 2007
Work Group Member, National Center for Ecological Analysis and Synthesis, 2008-2010
Scientific Committee for Ocean Research, Working Group 137, 2009-2012
External Review Panel, Ecology Program, Stazione Zoologica Anton Dohrn, Naples, Italy, 2009
Member, USGS Science Strategy Planning Team for Ecosystems, 2010-2012
Panelist, National Science Foundation, Long Term Ecosystem Research Program, 2012
International Scientific Advisory Committee, LabexMER (Marine Excellence Research), Brest, France
Science Advisor to Bay Area Ecosystems Climate Change Consortium, 2012-

SCIENTIFIC SOCIETIES (examples):

Editorial Board, *Limnology and Oceanography*, 1989-1992
Editorial Board, *Estuaries*, 1989-1994
Program Chair, Estuarine Research Federation, Biennial Meeting, 1991, San Francisco
Steering Committee, American Society of Limnology and Oceanography, 1997 Aquatic Sciences Meeting
Outstanding Reviewer, *Limnology and Oceanography*, 2004
Session Organizer, 2004 ASLO/TOS Ocean Research Conference, Honolulu
Associate Editor, *Limnology and Oceanography*, 2004-2005
Scientific Program Committee, 8th International Conference on Shellfish Restoration, Brest, France, 2005
Co-Editor in Chief, *Estuaries and Coasts*, 2007-2012
Convener and Organizer, AGU Chapman Conference on phytoplankton time series, Rovinj, Croatia, 2007
Session Organizer, 2009 Coastal and Estuarine Research Federation Meeting, Portland OR

KEYNOTE TALKS (examples):

University of Connecticut, Dedication of the Marine Sciences Building, 2001
Forum Québécois en Sciences de la Mer, Québec, Canada, 2002
CALFED Science Conference, Sacramento CA, 2003
National Water Quality Monitoring Council, San Jose CA, 2005
CALFED Science Conference, Sacramento CA, 2008
State of the San Francisco Estuary Conference, Oakland CA, 2009
Brazilian Congress of Oceanography, Rio Grande, Brazil 2010
Third International Symposium on Research and Management of Eutrophication in Coastal Ecosystems, Nyborg, Denmark 2010
International Council for Exploration of the Sea (ICES), Annual Science Conference, Gdansk, Poland 2011

INVITED TALKS (since 2006):

Marine Biological Laboratory, Woods Hole MA, Distinguished Scientist Seminar 2006
Estuarine Research Federation Meeting, Providence RI, 2007
AGU Meeting, San Francisco CA, 2008
CSIRO Marine and Atmospheric Research, Hobart, Australia 2008
ASLO Ocean Sciences Meeting, Nice, France, 2009
AAAS Pacific Division Symposium, San Francisco CA, 2009
National Environmental Research Institute, Roskilde, Denmark 2010
Centro Interdisciplinario de Ciencias Marinas, La Paz, Mexico 2010
Scientific Committee for Ocean Research, WG 137 workshop, Hangzhou, China, 2010
Ketchum Award acceptance lecture, Woods Hole Oceanographic Institution, Woods Hole MA, 2010
NRC Committee on Sustainable Water and Environmental Management in the California Bay-Delta, Sacramento CA, 2010
Scientific Committee for Ocean Research, WG 137 workshop, Sorrento, Italy, 2011
Stazione Zoologica Anton Dohrn, Naples, Italy, 2011
Coastal and Estuarine Research Federation Meeting, Daytona FL, 2011
Chesapeake Biological Laboratory, UMCEES, Solomons MD, 2012
Horn Point Laboratory, UMCEES, Cambridge MD, 2012 (3 lectures)
North American Congress for Conservation Biology, Oakland CA, 2012
University Lectures: Ecole Doctorale, Université de Bretagne Occidentale, Brest, France; Université de Rennes, France; Université de la Méditerranée, Marseille, France; College of Holy Cross, Biology Seminar; University of Washington, Department of Civil & Environmental Engineering; University of North Carolina, Institute of Marine Sciences; University of Wisconsin, Zoology Colloquium; University of

Texas at Austin, Marine Science Institute; University of California-Santa Cruz, Ocean Sciences Department; State University of New York – Stony Brook, School of Marine and Atmospheric Sciences; Stanford University, Dept. of Civil & Environmental Engineering; Scripps Institute of Oceanography; Griffith University-Gold Coast, Australia; Griffith University, Brisbane, Australia; Stanford University, Earth Systems Science; University of Wisconsin, Water Chemistry Program; University of California-Davis, Department of Environmental Science & Policy; McQuarie University, Sydney, Australia

OUTREACH AND COMMUNICATION

Public Lecture, USGS Menlo Park, CA 2004 [mms://video.wr.usgs.gov/science/may04.wmv](https://video.wr.usgs.gov/science/may04.wmv)

Tutorial on Scientific Writing and Publishing, Estuarine Research Federation Meeting, Providence RI, 2007

Public Lecture, University of Texas Marine Science Institute, Port Aransas TX, 2009

Tutorial on Scientific Writing and Publishing, Coastal & Estuarine Research Federation, Portland OR, 2009

Interview on NPR series *Living on Earth*, 2009, <http://www.livingonearth.org/shows/segments.htm?programID=09-P13-00004&segmentID=6>

Tutorial on Scientific Writing and Publishing, Woods Hole Oceanographic Institution, 2010

Organizer and Instructor, International Summer School: “Facing Climate Change”, Brest, France 2010

Published R package *wq0.2-2: Exploring water quality monitoring data* <http://cran.r-project.org/web/packages/wq/vignettes/wq-package.pdf>

Public Lecture, USGS Menlo Park, CA, 2011 <http://online.wr.usgs.gov/calendar/2011/jul11.html>

Interview published in *Estuary News* 20(5), 2011 <http://www.sfestuary.org/PDF/newsletters/October2011ESTUARYNEWS.pdf>

USGS Press Release about a 2011 scientific article http://www.usgs.gov/newsroom/article.asp?ID=3022#.T8jX_-1ORA4

Numerous radio, television and press interviews (e.g., KQED and KRCB radio, PBS.org, Sacramento Bee, San Jose Mercury News, The Tribune, The Telegraph, ClimateWire, The Sun Herald, The Miami Herald, Scientific American, Central Valley Business Times, Kansas City Star, McClatchy-Tribune News Service)

Briefings on the value of long term monitoring for staff members to the US Senate Committee on Environment and Public Works, Office of Management and Budget, House Science Committee, and Congressional Research Service, Washington DC, 2012 <http://www.compassonline.org/policy/May15/ScienceBriefing>

Tutorial on Scientific Writing and Publishing, Horn Point Laboratory, UMCEES, Cambridge MD, 2012

Public Lecture, Horn Point Laboratory, UMCEES, Cambridge MD, 2012

Public Lecture, DC Science Café, Washington DC, 2012 <http://www.dcswa.org/dc-science-cafe-11---two-bays>

BIBLIOGRAPHY – JAMES E. CLOERN

- 1 Cloern, J.E., 1976. The survival of coho salmon (*Oncorhynchus kisutch*) eggs in two Wisconsin tributaries of Lake Michigan: *American Midland Naturalist*, 96:451-461.
- 2 Cloern, J.E., 1976. Recent limnological changes in southern Kootenay Lake, British Columbia: *Canadian Journal of Zoology*, 54:1571-1578.
- 3 Cloern, J.E., 1977. Effects of light and temperature on the growth and nutrient-uptake kinetics of *Cryptomonas ovata*: *Journal of Phycology*, 13:389-395.
- 4 Cloern, J.E., 1978. Simulation model of *Cryptomonas ovata* population dynamics in southern Kootenay Lake, British Columbia: *Ecological Modeling*, 4:133-149.
- 5 Cloern, J.E., 1978. Empirical model of *Skeletonema costatum* photosynthetic rate, with applications in the San Francisco Bay estuary: *Advances in Water Resources*, 1:267-274.
- 6 Cloern, J.E., and Nichols, F.H., 1979. A von Bertalanffy growth model with a seasonally varying coefficient: *Journal of the Fisheries Research Board of Canada*, 35:1479-1482.
- 7 Cloern, J.E., 1979. Phytoplankton ecology of San Francisco Bay: The status of our current understanding, in Conomos, T.J., ed., *San Francisco Bay: The Urbanized Estuary*: Pacific Division, AAAS, San Francisco, page 247-264.
- 8 Cloern, J.E., and Cheng, R.T., 1981. Simulation model of *Skeletonema costatum* population dynamics in northern San Francisco Bay: *Estuarine and Coastal Shelf Science*, 12:83-100.
- 9 Alpine, A.E., Cloern, J.E., and Cole, B.E., 1981. Plankton studies in San Francisco Bay. I. Chlorophyll distributions and hydrographic properties of the San Francisco Bay estuary, July 1977-December 1979: U.S. Geological Survey Open-File Report 81-213.150 p.
- 10 Wong, R.L.J., and Cloern, J.E., 1981. Plankton Studies in San Francisco Bay. II. Phytoplankton abundance and species composition, July 1977 - December 1979: U.S. Geological Survey Open-File Report 81-214, 103 p.
- 11 Cloern, J.E., 1982. Does the benthos control phytoplankton biomass in South San Francisco Bay (USA)?: *Marine Ecology-Progress Series*, 9:191-202.
- 12 Luoma, S.N., and Cloern, J.E., 1982. The impacts of waste water discharge on biological communities in San Francisco Bay, in Kockelman, W.J., Conomos, T.J., and Leviton, A.E., eds., *San Francisco Bay: Use and Protection*: Pacific Division, AAAS, San Francisco, page 137-161.
- 13 Wong, R.L.J., and Cloern, J.E., 1982. Plankton studies in San Francisco Bay. IV. Phytoplankton abundance and species composition, January 1980 -February 1981: U.S. Geological Survey Open-File Report 82-443, 152 p.
- 14 Cloern, J.E., Cole, B.E., and Oremland, R.S., 1983. Chemistry and microbiology of a sewage spill in South San Francisco Bay: *Estuaries*, 6:399-406.
- 15 Cloern, J.E., Alpine, A.E., Cole, B.E., Wong, R.L.J., Arthur, J.F., and Ball, M.D., 1983. River discharge controls phytoplankton dynamics in the northern San Francisco Bay estuary: *Estuarine, Coastal and Shelf Science*, 16:415- 429.
- 16 Cloern, J.E., Cole, B.E., and Oremland, R.S., 1983. Autotrophic processes in meromictic Big Soda Lake, Nevada: *Limnology and Oceanography*, 28:1049- 1061.
- 17 Cloern, J.E., Cole, B.E., and Oremland, R.S., 1983. Seasonal changes in the chemical and biological nature of a meromictic lake (Big Soda Lake, Nevada, USA): *Hydrobiologia*, 105:195-206.
- 18 Cloern, J.E., 1984. Temporal dynamics and ecological significance of salinity stratification in an estuary: *Oceanologica Acta*, 7:137-141.
- 19 Cole, B.E., and Cloern, J.E., 1984. Significance of biomass and light availability to phytoplankton productivity in San Francisco Bay: *Marine Ecology- Progress Series*, 15:15-24.
- 20 Catts, G., Khorram, S., Cloern, J.E., Knight, A.W., and DeGloria, S.D., 1985. Remote sensing of tidal chlorophyll a variations in estuaries: *International Journal of Remote Sensing*, 6:1685-1706.
- 21 Alpine, A.E., and Cloern, J.E., 1985. Differences in in-vivo fluorescence yield between three phytoplankton size classes: *Journal of Plankton Research*, 7:381-390.
- 22 Ambler, J.W., Cloern, J.E., and Hutchinson, A., 1985. Seasonal cycles of zooplankton from San Francisco Bay: *Hydrobiologia*, 129:177-197.
- 23 Cloern, J.E., Cole, B.E., Wong, R.L.J., and Alpine, A.E., 1985. Temporal dynamics of estuarine phytoplankton: A case study of San Francisco Bay: *Hydrobiologia*, 129:153-176.
- 24 Cloern, J.E., and Nichols, F.H., 1985. Time scales and mechanisms of estuarine variability - a synthesis from studies of San Francisco Bay: *Hydrobiologia*, 129:229-237.

- 22 Cloern, J.E., and Nichols, F.H., eds., 1985. *Temporal dynamics of an estuary - San Francisco Bay*: Developments in Hydrobiology No. 30, Kluwer Academic Publishers, Dordrecht, 237 pages.
Alpine, A.E., Wienke, S.M., Cloern, J.E., and Cole, B.E., 1985. Plankton studies in San Francisco Bay. VIII. Chlorophyll distributions and hydrographic properties in South San Francisco Bay, 1983: U.S. Geological Survey Open-File Report 82-443, 152 p.
- 23 Nichols, F.H., Cloern, J.E., Luoma, S.N., and Peterson, D.H., 1986. Modification of an estuary: *Science*, 231:567-573.
- 24 Cole, B.E., Cloern, J.E., and Alpine, A.E., 1986. Biomass and productivity of three phytoplankton size classes in San Francisco Bay: *Estuaries*, 9:117-126.
- 25 Powell, T.M., Cloern, J.E., and Walters, R.A., 1986. Phytoplankton spatial distribution in South San Francisco Bay -- mesoscale and small-scale variability, in D.A. Wolfe, ed., *Estuarine Variability*: Academic Press, N.Y., page 369-383.
- 26 Cole, B.E., and Cloern, J.E., 1987. An empirical model of phytoplankton productivity in estuaries: *Marine Ecology Progress Series*, 36:299-305.
- 27 Wienke, S.M., and Cloern, J.E., 1987. The phytoplankton component of seston in San Francisco Bay: *Netherlands Journal of Sea Research*, 21:25-33.
- 28 Khorram, S., Catts, G.P., Cloern, J.E., and Knight, A.W., 1987. Modeling of chlorophyll *a* from an airborne scanner: *IEEE Transactions on GeoScience and Remote Sensing*, v. GE-25, page 662-669.
Cloern, J.E., 1987. Review of the book "Synthesis and Modeling of Intermittent Estuaries", by W.R. Cuff and M. Tomczak, Jr.: *Marine Geology*, 76:332-333.
- 29 Zehr, J.P., Harvey, R.W., Oremland, R.S., Cloern, J.E., George, L., and Lane, J.L., 1987. Big Soda Lake (Nevada). 1. Pelagic bacterial heterotrophy and biomass: *Limnology and Oceanography*, 32:781-793.
- 30 Cloern, J.E., Cole, B.E., and Wienke, S.M., 1987. Big Soda Lake (Nevada). 4. Vertical fluxes of particulate matter - seasonality and variations across the chemocline: *Limnology and Oceanography*, 32:815-824.
- 31 Cloern, J.E., 1987. Turbidity as a control on phytoplankton biomass and productivity in estuaries: *Continental Shelf Research*, 7:1367-1381.
- 32 Powell, T.M., Willmot, S., Murray, J.D., Manoranjan, V.S., and Cloern, J.E., 1988. Transient spatial patterns in plankton communities: blooms and traveling waves of phytoplankton in estuaries, in Hallam, T.G., Gross, L.J. and Levin, S.A., eds., 1986 Proceedings of Trieste Research Conference on Mathematical Ecology: World Scientific Publishing, page 236-267.
- 33 Alpine, A.E., and Cloern, J.E., 1988. Phytoplankton growth rates in a light- limited environment, San Francisco Bay. *Marine Ecology Progress Series*, 44:167-173.
- 34 Oremland, R.S., Cloern, J.E., Sofer, Z., Smith, R.L., Culbertson, C.W., Zehr, J., Miller, L., Cole, B.E., Harvey, R., Iversen, N., Klug, M., Des Marais, D.J., and Rau, G., 1988. Microbial and biogeochemical processes in Big Soda Lake, Nevada, in A.J. Fleet, K. Kelts, and M.R. Talbot, eds., *Lacustrine Petroleum Source Rocks*: Geological Society Special Publication no. 40, page 59-75.
Alpine, A.E., Wienke, S.M., Cloern, J.E., and Cole, B.E., 1988. Plankton studies in San Francisco Bay. IX. Chlorophyll distributions and hydrographic properties in South San Francisco Bay, 1984-1986: U.S. Geological Survey Open-File Report 88-319, 86 p.
- 35 Kuwabara, J.S., Chang, C.C.Y., Cloern, J.E., Fries, T.L., Davis, J.A., and Luoma, S.N., 1989. Trace metal associations in the water column of South San Francisco Bay, California: *Estuarine, Coastal and Shelf Science*, 28:307- 325.
Cloern, J.E., 1989. Phytoplankton, in Britton, L.J. and Greeson, P.E., eds., *Methods for collection and analysis of aquatic biological and microbiological samples: Techniques of Water Resources Investigations of the U.S. Geological Survey*, Chapter A-4, p. 99-115.
- 36 Powell, T.M., Cloern, J.E., and Huzzey, L.M., 1989. Spatial and temporal variability in South San Francisco Bay. I. Horizontal distributions of salinity, suspended sediments, and phytoplankton biomass and productivity: *Estuarine, Coastal and Shelf Science*, 28:583-597.
- 37 Cloern, J.E., Powell, T.M., and Huzzey, L.M., 1989. Spatial and temporal variability in South San Francisco Bay. II. Temporal changes in salinity, suspended sediments, and phytoplankton biomass and productivity over tidal time scales: *Estuarine, Coastal and Shelf Science*, 28:599-613.
- 38 Huzzey, L.M., Cloern, J.E., and Powell, T.M., 1990. Episodic changes in lateral transport and phytoplankton distribution in South San Francisco Bay: *Limnology and Oceanography*, 35:472-478.
Wienke, S.M., Alpine, A.E., Cloern, J.E., and Cole, B.E., 1990. Plankton studies in San Francisco Bay. X. Chlorophyll distributions and hydrographic properties in San Francisco Bay, 1987: U.S. Geological Survey Open-File Report 90- 145, 50 p.
Wienke, S.M., Cloern, J.E., and Cole, B.E., 1990. Plankton studies in San Francisco Bay. XI. Chlorophyll distributions and hydrographic properties in San Francisco Bay, 1988-1989: U.S. Geological Survey Open-File Report 90-562, 212 p.

- 39 Cloern, J.E., 1991, Annual variations in river flow and primary production in the South San Francisco Bay Estuary, in Elliott, M. and Ducrottoy, D., eds., *Estuaries and Coasts: Spatial and Temporal Intercomparisons*: Olsen and Olsen Publishers, Denmark, p. 91-96.
- 40 Cloern, J.E., 1991, Tidal stirring and phytoplankton bloom dynamics in an estuary: *Journal of Marine Research*, 49:203-221.
- Wienke, S.M., Cole, B.E., Cloern, J.E., and Alpine, A.E., 1991, Plankton studies in San Francisco Bay. XII. Chlorophyll distributions and hydrographic properties in San Francisco Bay, 1990: U.S. Geological Survey Open-File Report 91-476, 85 p.
- 41 Alpine, A.E., and Cloern, J.E., 1992, Trophic interactions and direct physical effects control phytoplankton biomass and production in an estuary: *Limnology and Oceanography*, 37:946-955.
- 42 Cole, B.E., Thompson, J.K., and Cloern, J.E., 1992, Measurement of filtration rates by infaunal bivalves in a recirculating flume: *Marine Biology*, 113:219-225.
- 43 Cloern, J.E., Alpine, A.E., Cole, B.E., and Heller, T., 1992, Seasonal changes in the spatial distribution of phytoplankton in small, temperate-zone lakes: *Journal of Plankton Research*, 14:1017-1024.
- Wienke, S.M., Cole, B.E., Cloern, J.E., and Alpine, A.E., 1992, Plankton studies in San Francisco Bay. XIII. Chlorophyll distributions and hydrographic properties in San Francisco Bay, 1991: U.S. Geological Survey Open-File Report 92-158, 116 p.
- 44 Jassby, A.D., Cloern, J.E., and Powell, T.M., 1993, Organic carbon sources and sinks in San Francisco Bay: variability induced by river flow. *Marine Ecology Progress Series*, 95:39-54.
- McBride, G.B., Vant, W.N., Cloern, J.E., and Liley, J.B., 1993, Development of a model of phytoplankton blooms in Manukau Harbour: NIWA Ecosystems Publication No. 3, Hamilton, New Zealand, 52 pp..
- 45 Koseff, J.R., Holen, J.K., Monismith, S.G., and Cloern, J.E., 1993, Effects of vertical mixing and benthic grazing on phytoplankton populations in shallow, turbid estuaries: *Journal of Marine Research*, 51:843-868.
- Cloern, J.E., Canuel, E.A., and Wienke, S.M., 1993, Particulate organic matter in the San Francisco Bay estuary, California: chemical indicators of its origin and assimilation into the benthic food web: U.S. Geological Survey Open-File Report 93-146, 42 p.
- Wienke, S.M., Cole, B.E., and Cloern, J.E., 1993, Plankton studies in San Francisco Bay. XIV. Chlorophyll distributions and hydrographic properties in San Francisco Bay, 1992: U.S. Geological Survey Open-File Report 93-423, 175 p.
- 46 Cloern, J.E., Cole, B.E. and Hager, S.W. 1994. Notes on a *Mesodinium rubrum* Red Tide in South San Francisco Bay (USA): *Journal of Plankton Research*, 16:1269-1276.
- Jassby, A., Cloern, J.E., Caffrey, J., Cole, B., and Rudek, J., 1994, San Francisco Bay/Delta Regional Monitoring Program plankton and water quality pilot study, 1993, in 1993 Annual Report. San Francisco Estuary Regional Monitoring Program for Trace Substances. San Francisco Estuary Institute, p. 117-128
- Caffrey, J.M., Cole, B.E., Cloern, J.E., Rudek, J.R., Tyler, A.C., and Jassby, A.D., 1994, Studies of the plankton and its environment in the San Francisco Bay Estuary, California. Regional Monitoring Results, 1993: U.S. Geological Survey Open-File Report 94-82, 411 p.
- 47 Cloern, J.E. and Jassby, A.D., 1995, Year-to-year fluctuation of the spring phytoplankton bloom in South San Francisco Bay: An example of ecological variability at the land-sea interface, in Steele, J.H, Powell, T.M., and Levin, S., eds., *Ecological Time Series*, Chapman Hall, p. 139-149.
- 48 Canuel, E.A., Cloern, J.E., Ringelberg, D., Guckert, J., and Rau, G., 1995, Molecular and isotopic tracers used to understand sources of organic matter and trophic relationships in the San Francisco Bay estuary: *Limnology and Oceanography*, 40:67-81.
- Cloern, J.E., 1995, Ecosystem responses to phytoplankton blooms -- examples from South San Francisco Bay: Newsletter, Interagency Ecological Program for the Sacramento-San Joaquin Estuary, Spring 1995, p. 10-14.
- Cloern, J.E., Luoma, S.N., and Nichols, F.H., 1995, The United States Geological Survey San Francisco Bay Program: Lessons Learned for Managing Coastal Water Resources: U.S. Geological Survey, Fact Sheet FS-053-95
- 49 Jassby, A.D., Kimmerer, W.J., Monismith, S., Armor, C., Cloern, J.E., Powell, T.M., Schubel, J.R., and Vendlinski, T. 1995, Isohaline position as a habitat indicator for estuarine resources: San Francisco Bay-Delta, California, U.S.A.: *Ecological Applications*, 5:272-289..
- Jassby, A.D., Cloern, J.E., Cole, B.E., and Caffrey, J.M., 1995, Choosing optimum station configurations for summarizing water quality characteristics, in 1994 Annual Report, San Francisco Estuary Regional Monitoring Program for Trace Substances: San Francisco Estuary Institute, p. 157-169.
- 50 Cloern, J.E., Grenz, C., and Vidregar-Lucas, L., 1995. An empirical model of the phytoplankton chlorophyll/carbon ratio -- the conversion factor between productivity and growth rate: *Limnology and Oceanography*, 40:1313-1321.

- Edmunds, J.L., Cole, B.E., Cloern, J.E., Caffrey, J.M., and Jassby, A.D., 1995, Studies of the San Francisco Bay, California, Estuarine Ecosystem. Pilot Regional Monitoring Program Results, 1994: U.S. Geological Survey Open-File Report 95-378, 436 p.
- 51 Canuel, E.A., and Cloern, J.E., 1996, Regional differences in the origins of organic matter in the San Francisco Bay ecosystem. Evidence from lipid biomarkers, in J.T. Hollibaugh, editor, *San Francisco Bay: The Ecosystem*, Pacific Division, AAAS, San Francisco, p. 305-324.
- 52 Rudek, J., and Cloern, J.E., 1996, Planktonic respiration rates in San Francisco Bay, in J.T. Hollibaugh, editor, *San Francisco Bay: The Ecosystem*, Pacific Division, AAAS, San Francisco, p. 289-304.
- 53 Cloern, J.E., 1996. Phytoplankton bloom dynamics in coastal ecosystems: A review with some general lessons from sustained investigation of San Francisco Bay, California: *Reviews of Geophysics*, 34:127-168.
- Cole, B.E., Cloern, J.E., and Alpine, A.E., 1997, The photosynthetic response of phytoplankton in Shingobee Lake and Williams Lake: USGS Water-Resources Investigations Report 96-4215, pages 105-110.
- Cloern, J.E., Alpine, A.E., and Cole, B.E., 1997, Seasonal comparisons of seston abundance and sedimentation rate in a closed-basin lake (Williams) and an open-basin lake (Shingobee): USGS Water-Resources Investigations Report 96-4215, pages 111-117.
- 54 Jassby, A.D., Cole, B.E., and Cloern, J.E., 1997. The design of sampling transects for characterizing water quality in estuaries: *Estuarine, Coastal and Shelf Science*, 45: 285-302.
- Cloern, J.E., Cole, B.E., and Edmunds, J.L., 1997, Seasonal fluctuations of water quality in San Francisco Bay during the first three years of the Regional Monitoring Program 1993-1995, in 1995 Annual Report, San Francisco Estuary Regional Monitoring Program for Trace Substances: San Francisco Estuary Institute, pages 5.1.2 to 5.1.15.
- Rodgers, K.M., Garrison, D.L., and Cloern, J.E., 1997, Toxic phytoplankton in San Francisco Bay, in 1995 Annual Report, San Francisco Estuary Regional Monitoring Program for Trace Substances: San Francisco Estuary Institute, pages 7.4.1 to 7.4.15.
- Edmunds, J.L., Cole, B.E., Cloern, J.E., and Dufford, R.G., 1997, Studies of the San Francisco Bay, California, Estuarine Ecosystem. Pilot Regional Monitoring Program Results, 1995: U.S. Geological Survey Open-File Report 97-15, 380 p.
- Baylous, J.I., Edmunds, J.L., Cole, B.E., and Cloern, J.E., 1997, Studies of the San Francisco Bay, California, Estuarine Ecosystem. Pilot Regional Monitoring Program Results, 1996: U.S. Geological Survey Open-File Report 97-598, 218 p.
- Cloern, J.E., B.E. Cole, J.L. Edmunds, and J.I. Baylous. 1997. Water-quality variability in San Francisco Bay, some general lessons from 1996 sampling, in 1996 Annual Report, San Francisco Estuary Regional Monitoring Program for Trace Substances: San Francisco Estuary Institute, 46-60.
- 55 Luoma, S.N., van Geen, A., Lee, B.-G., and Cloern, J.E., 1998, Metal uptake by phytoplankton during a bloom in South San Francisco Bay: Implications for metal cycling in estuaries: *Limnology and Oceanography*, 43:1007-1016.
- 56 Lucas, L, J.E. Cloern, J.R. Koseff, S.G. Monismith, and J.K. Thompson, 1998, Does the Sverdrup Critical Depth Model explain bloom dynamics in estuaries? *Journal of Marine Research*, 56:1-41.
- Cloern, J.E. 1998. Book Review: "Eutrophication in Coastal Marine Ecosystems", Jørgensen, B.B. and K. Richardson [Eds.]. *Limnology and Oceanography*, 43:1018-1019.
- Baylous, J.I., Cole, B.E., and Cloern, J.E., 1998, Studies of the San Francisco Bay, California, Estuarine Ecosystem. Pilot Regional Monitoring Program Results, 1997: U.S. Geological Survey Open-File Report 98-168,
- 57 Caffrey, J.M., C. Grenz, and J.E. Cloern, 1998. Changes in production and respiration during a spring phytoplankton bloom in San Francisco Bay, California: implications for net ecosystem metabolism. *Marine Ecology Progress Series* 172:1-12.
- 58 Cloern, J.E. 1999. The Relative Importance of Light and Nutrient Limitation of Phytoplankton Growth: A Simple Index of Coastal Ecosystem Sensitivity to Nutrient Enrichment: *Aquatic Ecology* 33: 3-15.
- 59 Lucas, L.V., J.R. Koseff, J.E. Cloern, S.G. Monismith and J.K. Thompson. 1999. Processes governing phytoplankton blooms in estuaries. Part I. The local production-loss balance. *Marine Ecology Progress Series* 187:1-15.
- 60 Lucas, L.V., J.R. Koseff, J.E. Cloern, S.G. Monismith and J.K. Thompson. 1999. Processes governing phytoplankton blooms in estuaries. Part II. The role of transport in global dynamics. *Marine Ecology Progress Series* 187: 17-30.
- Cloern, J.E., Cole, B.E., Edmunds, J.L., and J.I. Baylous. 1999. Water-quality variability in San Francisco Bay: general patterns of change during 1997. in 1997 Annual Report, San Francisco Estuary Regional Monitoring Program for Trace Substances: San Francisco Estuary Institute, pp. 67-81.
- Edmunds, J.L., Kuivila, K.M., Cole, B.E., and J.E. Cloern. 1999. Do Herbicides Impair Phytoplankton Primary Production in the Sacramento-San Joaquin River Delta? in Morganwalp, D.W., and Buxton, H.T., eds., U.S. Geological Survey Toxic Substances Hydrology Program--Proceedings of the Technical Meeting, Charleston, South Carolina, March

- 8-12, 1999--Volume 2 of 3--Contamination of Hydrologic Systems and Related Ecosystems: U.S. Geological Survey Water-Resources Investigations Report 99-4018B, pp. 81-88.
- 61 Grenz C., Cloern J.E., Hager S.W., Cole B.E. 2000. Dynamics of nutrient cycling and related benthic nutrient and oxygen fluxes during a spring phytoplankton bloom in South San Francisco Bay (USA). *Marine Ecology Progress Series* 197: 67-80
- 62 Ning, X., Cloern, J.E., and Cole, B.E. 2000. Spatial and temporal variability of picocyanobacteria *Synechococcus* sp. in San Francisco Bay. *Limnology and Oceanography* 45: 695-702.
- 63 Jassby, A.D. and J.E. Cloern. 2000. Organic matter sources and rehabilitation of the Sacramento-San Joaquin Delta (California, USA). *Aquatic Conservation: Marine and Freshwater Ecosystems* 10: 323-352.
- 64 Howarth, R., D. Anderson, J. Cloern, C. Hopkinson, B. LaPointe, T. Malone, N. Marcus, K. McGlathery, and A. Sharpley. 2000. Nutrient Pollution of Coastal Rivers, Bays, and Seas. *Issues in Ecology* 7:1-15.
- 65 Cloern, J.E. 2001. Our evolving conceptual model of the coastal eutrophication problem. *Marine Ecology Progress Series* 211: 223-253.
- 66 Cloern, J.E., E.A. Canuel, D. Harris. 2002. Stable carbon and nitrogen isotope composition of aquatic and terrestrial plants of the San Francisco Bay estuarine system. *Limnology and Oceanography* 47: 713-729.
- 67 Jassby, A.D., J.E. Cloern, B.E. Cole. 2002. Annual primary production: patterns and mechanisms of change in a nutrient-rich tidal ecosystem. *Limnology and Oceanography* 47: 698-712.
- 68 Guarini, J.-M., J.E. Cloern, J. Edmunds, P. Gros. 2002. Microphytobenthic potential productivity estimated in three tidal embayments of the San Francisco Bay: A comparative study. *Estuaries* 25: 409-417.
- 69 Lucas, L.V., J.E. Cloern, J.K. Thompson, N.E. Monsen. 2002. Functional variability of habitats within the Sacramento-San Joaquin Delta: restoration implications. *Ecological Applications* 12: 1528-1547.
- 70 Lucas, L.V. and J.E. Cloern. 2002. Effects of tidal shallowing and deepening on phytoplankton production dynamics: a modeling study. *Estuaries* 25: 497-507.
- 71 Sobczak, W.S., Cloern, J.E., Jassby, A.D and Mueller-Solger, A. 2002. Bioavailability of organic matter in a highly disturbed estuary. The role of detrital and algal resources. *Proceedings of the National Academy of Sciences* 99: 8101-8105.
- 72 Monsen, N.E., Cloern, J.E., Lucas, L.V., and Monismith, S.G. 2002. A comment on the use of flushing rate, residence time and age as transport time scales. *Limnology and Oceanography* 47: 1545-1553.
- 73 Chauvaud, L., Thompson, J.K., Cloern, J.E. and Thouzeau, G. 2003. Clams as CO₂ generators: The *Potamocorbula amurensis* example in San Francisco Bay. *Limnology and Oceanography* 48: 2086-2092.
- 74 Jassby, A.D., J.E. Cloern, A. Mueller-Solger. 2003. Phytoplankton fuels the food web in Delta waterways. *California Agriculture* 57:104-109.
- 75 May, C.L., J.R. Koseff, L.V. Lucas, J.E. Cloern, and D.H. Schoellhamer. 2003. Effects of spatial and temporal variability of turbidity on phytoplankton blooms. *Marine Ecology Progress Series* 254: 111-128.
- Cloern, J.E., T.S. Schraga, C.B. Lopez, R. Labiosa. 2003. Lessons from monitoring water quality in San Francisco Bay. In 2003 Pulse of the Estuary, San Francisco Estuary Institute, pp. 15-20.
- 76 Cloern, J.E., R. Dufford. 2005. Phytoplankton community ecology: principles applied in San Francisco Bay. *Marine Ecology Progress Series*, 285:11-28.
- 77 Sobczak, W.V., J.E. Cloern, A.D. Jassby, B.E. Cole, T.S. Schraga, and A. Arnsberg. 2005. Detritus fuels ecosystem metabolism but not metazoan food webs in San Francisco Estuary's freshwater Delta. *Estuaries* 28(1):122-135.
- 78 Cloern, J.E., T.S. Schraga and C. Burns Lopez. 2005. Heat wave brings a red tide to San Francisco Bay. *Eos Transactions of the American Geophysical Union* 86(7):66
- 79 Cloern, J.E., T.S. Schraga, C.B. Lopez, N. Knowles, R.G. Labiosa, and R. Dugdale. 2005. Climate anomalies generate an exceptional dinoflagellate bloom in San Francisco Bay. *Geophysical Research Letters* 32, L14608.
- 80 Bachman, R.W., J.E. Cloern, R.E. Hecky and D.W. Schindler (editors). 2006. Eutrophication of Freshwater and Marine Ecosystems. *Limnology and Oceanography*, Special Issue 51: 351-800.
- 81 Lopez, C.B., J.E. Cloern, T.S. Schraga, A.J. Little, L.V. Lucas, J.K. Thompson and J.R. Burau. 2006. Ecological values of shallow-water habitats: implications for restoration of a highly disturbed ecosystem. *Ecosystems* 9:422-440.
- Cloern, J.E., A.D. Jassby, T.S. Schraga and K.L. Dallas. 2006. What is causing the phytoplankton increase in San Francisco Bay? 2006 Pulse of the Estuary. San Francisco Estuary Institute, Oakland. Available online: <http://www.sfei.org/rmp/pulse/2006/index.html>
- Cloern, James (Lead Author) 2006. "Eutrophication." In: Encyclopedia of Earth. Eds. Cutler J. Cleveland (Washington, D.C.: Environmental Information Coalition, National Council for Science and the Environment) <http://www.eoearth.org/article/Eutrophication>
- 82 Cloern, J.E. 2007. Habitat Connectivity and Ecosystem Productivity: Implications from a Simple Model. *The American Naturalist* 169:E21-E33, <http://www.journals.uchicago.edu/AN/journal/issues/v169n1/41465/41465.web.pdf>

- 83 Monsen, N.E., J.E. Cloern, J.R. Burau. 2007. Effects of flow diversions on water and habitat quality: examples from the highly manipulated Sacramento-San Joaquin Delta (California). *San Francisco Estuary and Watershed Science*, Vol. 5, Issue 3, <http://repositories.cdlib.org/jmie/sfews/vol5/iss3/art2>
- 84 Cloern, J.E., A.D. Jassby, J.K. Thompson, K. Hieb. 2007. A cold phase of the East Pacific triggers new phytoplankton blooms in San Francisco Bay. *Proceedings of the National Academy of Sciences of the United States of America* 104(47):18561-18656, Open Access <http://www.pnas.org/cgi/reprint/104/47/18561> [highlighted in 07 December 2007 SCIENCE Magazine as an Editor's Choice Paper: <http://www.sciencemag.org/content/vol318/issue5856/twil.dtl#318/5856/1525d>] [recommended by Faculty1000 Biology Faculty of 1000 Biology: evaluations for Cloern JE et al *Proc Natl Acad Sci U S A* 2007 Nov 20 104 (47) :18561-5 <http://www.f1000biology.com/article/id/1098465/evaluation>]
- 85 Smetacek, V. and J.E. Cloern. 2008. Perspective: On phytoplankton trends. *Science* 319:1346-1348.
- 86 Thébault, J., T.S. Schraga, J.E. Cloern, E.G. Dunlavy. 2008. Primary production and carrying capacity of former salt ponds after reconnection to San Francisco Bay. *Wetlands* 28(3):841-851.
- 87 Cloern, J.E. and A.D. Jassby. 2008. Complex seasonal patterns of primary producers at the land-sea interface. *Ecology Letters*, 11:1294-1303.
- 88 Cloern, J.E. and A.D. Jassby. 2009. Patterns and scales of phytoplankton variability in estuarine-coastal ecosystems. *Estuaries and Coasts* DOI: 10.1007/s12237-009-9195-3 <http://www.springerlink.com/content/e52651606gg83868/fulltext.pdf> (this paper was highlighted in Coastal & Estuarine Science News, <http://www.erf.org/cesn/november-2009#article3>)
- 89 Winder, M. and J.E. Cloern. 2010. The annual cycles of phytoplankton biomass. *Philosophical Transactions of the Royal Society B*, **365**, 3215-3226 doi: 10.1098/rstb.2010.0125, <http://rstb.royalsocietypublishing.org/content/365/1555/3215.full.pdf?ijkey=zqBcAp7H4HlOmI&keytype=finite>
- Jassby, A.D. and J.E. Cloern. 2010. R package **wq0.3-3**: *Exploring water quality monitoring data*. (<http://cran.r-project.org/web/packages/wq/index.html>).
- 90 Cloern, J. E., et al. 2010. Biological communities in San Francisco Bay track large-scale climate forcing over the North Pacific, *Geophys. Res. Lett.*, **37**, L21602, doi:10.1029/2010GL044774.
- Cloern, J.E. and R. Dugdale. 2010. San Francisco Bay, pp. 117-126 in Glibert, P.M. et al. (eds), *Nutrients in Estuaries: A Summary Report of the national Estuarine Experts Workgroup 2005-2007*.
- 91 Guarini J.-M., L. Chauvaud, J. Cloern, J. Clavier, J. Coston-Guarini, Y. Patry. 2011. Seasonal variations in ectotherm growth rates: quantifying growth as an intermittent non steady state compensatory process. *Journal of Sea Research*, doi:10.1016/j.seares.2011.02.001
- Paerl, H., K. Yin, and J. Cloern. 2011. Meeting Report, Global Patterns of Phytoplankton Dynamics in Coastal Ecosystems: Comparative Analysis of Time Series Observations, *SCOR Working Group 137 Meeting; Hangzhou, China, 17-21 October 2010*. *Eos*, 92(10):85
- 92 Cloern JE, Knowles N, Brown LR, Cayan D, Dettinger MD, Morgan TL, Schoellhamer, DH, Stacey, MT, van der Wegen, M, Wagner, RW, Jassby, AD. 2011. Projected Evolution of California's San Francisco Bay-Delta-River System in a Century of Climate Change. *PLoS ONE* 6(9): <http://www.plosone.org/article/info%3Adoi%2F10.1371%2Fjournal.pone.0024465>
- 93 Cloern, J.E. et al. 2012. Perils of correlating CUSUM-transformed variables to infer ecological relationships (Breton et al. 2006, Glibert 2010). *Limnology and Oceanography*, 57(2): 665-668. http://www.aslo.org/lo/toc/vol_57/issue_2/0665.pdf
- Williams, B.K., Wingard, G.L. Brewer, G., Cloern, J., Gelfenbaum, G., Jacobson, R.B., Kershner, J.L., McGuire, A.D., Nichols, J.D., Shapiro, C.D., van Riper III, C., and White, R.P., 2012, The USGS Ecosystem Science Strategy 2012-2022: Advancing discovery and application through collaboration: U.S. Geological Survey Open-File Report 2012-XXXX, xx p., available at <http://pubs.usgs.gov/of/2012/XXXX/>.
- 94 Cloern, J.E. and A.D. Jassby, Drivers of change in estuarine-coastal ecosystems: discoveries from four decades of study in San Francisco Bay. *Reviews of Geophysics*, In press.

Drivers of Change in Estuarine-Coastal Ecosystems: Discoveries from Four Decades of Study in San Francisco Bay

James E. Cloern¹ and Alan D. Jassby²

¹ U. S. Geological Survey, 345 Middlefield Rd., Menlo Park, California, USA (tel 1-650-329-4594; e-mail jeclorn@usgs.gov)

² Department of Environmental Science and Policy, University of California, One Shields Avenue, Davis, California, USA (e-mail adjassby@ucdavis.edu)

1 **ABSTRACT**

2 Poised at the interface of rivers, ocean, atmosphere and dense human settlement, estuaries are
3 driven by a large array of natural and anthropogenic forces. San Francisco Bay exemplifies the
4 fast-paced change occurring in many of the world's estuaries, bays and inland seas in response to
5 these diverse forces. We use observations from this particularly well-studied estuary to illustrate
6 responses to six drivers that are common agents of change where land and sea meet: water
7 consumption and diversion; human modification of sediment supply; introduction of non-native
8 species; sewage input; environmental policy; and climate shifts. In San Francisco Bay, responses
9 to these drivers include, respectively, shifts in the timing and extent of freshwater inflow and
10 salinity intrusion; decreasing turbidity; restructuring of plankton communities; nutrient
11 enrichment; elimination of hypoxia and reduced metal contamination of biota; and food web
12 changes that decrease resistance of the estuary to nutrient pollution. Detection of these changes
13 and discovery of their causes through environmental monitoring have been essential for
14 establishing and measuring outcomes of environmental policies that aim to maintain high water
15 quality and sustain services provided by estuarine-coastal ecosystems. The wide range of
16 variability time scales and the multiplicity of interacting drivers place heavy demands on
17 estuarine monitoring programs. But the San Francisco Bay case study illustrates why the
18 imperative for monitoring has never been greater.

19 **1. INTRODUCTION**

20 Four decades ago, the study of ecosystems was emerging as a scientific discipline to understand
21 how biological communities and their physical environment are organized spatially and how they
22 change over time. A classic paper from that era was Eugene P. Odum's *The Strategy of*
23 *Ecosystem Development*, which depicts ecosystem change over time as an orderly process of
24 community development culminating in a stable system, and where stability is maintained by the
25 evolution of complex biological structure and its "increasing control of and homeostasis with the
26 physical system" [Odum, 1969]. Odum described ecosystem development as successional stages
27 from immature to mature communities, where the stages are *predictable*, include *subtle* changes
28 in food webs, and climax to a *steady state*. But Odum also pointed out that "[s]evere stress or
29 rapid changes brought about by outside forces can, of course, rob the system of these protective
30 mechanisms..." and "[m]ost physical stresses introduced by man are too sudden, too violent, or
31 too arrhythmic for adaptations to occur at the ecosystem level, so severe oscillation rather than
32 stability results". Odum's narrative description of unperturbed mature ecosystems provides a
33 benchmark from which we can judge the extent to which ecosystems are altered by severe
34 outside forces, including anthropogenic ones.

35 This review is about change in marine ecosystems connected to land, such as estuaries,
36 bays and lagoons. Odum and his contemporaries had access to long-term (multidecadal) records
37 of biological and environmental variability from terrestrial [Baltensweiler, 1964] and marine
38 [Southward, 1995] ecosystems, but in the late 1960s there were few observational records
39 documenting decadal-scale changes in estuaries. Numerous coastal research and monitoring
40 programs began in the 1970s and 1980s, however, and have been sustained long enough to
41 provide empirical bases for comparison against Odum's attributes of mature unperturbed

42 ecosystems. We show 10 examples (Figure 1) to illustrate changes in: sediment supply to the
43 Yangtze Estuary, China [*Li et al.*, 2012]; dissolved oxygen concentration and phosphorus input
44 to the Potomac Estuary, USA [*Jaworski et al.*, 2007]; diatom productivity (Si-uptake) in northern
45 San Francisco Bay, USA [*W Kimmerer*, 2005]; optical properties of Chesapeake Bay, USA
46 [*Gallegos et al.*, 2011]; areal extent of submerged aquatic vegetation in the lower Potomac River,
47 USA [*Orth et al.*, 2010]; biomass of benthic invertebrates in Denmark's Ringkøbing Fjord
48 [*Petersen et al.*, 2008]; mercury content of mussels in the Forth Estuary, UK [*Dobson*, 2000];
49 annual landings of pelagic fish in the Limfjorden, Denmark [*Riisgard*, 2012]; and abundance of
50 shorebirds (black-tailed godwits) in the Tagus Estuary, Portugal [*Catry et al.*, 2011].

51 These examples are representative of the substantial and rapid physical, biogeochemical
52 and biological changes that have occurred in many of the world's estuarine-coastal ecosystems in
53 recent decades. They reveal complex and diverse patterns of change as monotonic trends or
54 abrupt shifts (up or down), oscillations and multi-year peaks in plant and animal abundance, and
55 they show that estuarine-coastal environments and their biological communities are changing at a
56 fast pace. Many of these changes were surprises and did not occur as *predictable* successional
57 stages, none could be classified as *subtle*, and the large trends, step changes and high-amplitude
58 oscillations are not characteristic of *steady state*. Therefore, empirical observations from the
59 world's estuaries, bays and lagoons reveal dynamics distinctly different from Odum's description
60 of how ecosystems develop in the absence of unusual external forces.

61 Our purpose is to use a suite of observations collected in a particularly well-studied
62 estuary to review what has been learned in recent decades about the external forces that drive
63 diverse and rapid changes at the land-sea interface. We describe changes in hydrology, sediment
64 dynamics, biological communities and water quality that have been captured in sampling

65 programs sustained over multiple decades in San Francisco Bay, its watershed and the adjacent
66 coastal Pacific Ocean. We selected examples where change can be attributed to a specific driver
67 or pressure -- a human action or climatic forcing that brings about fundamental change at the
68 ecosystem scale. Identification of drivers is a key to understanding past changes, and it provides
69 the foundation for anticipating and adapting to future changes [*S T Jackson, 2007*]. Aided by rich
70 observational records, we use the San Francisco Bay case study to illustrate responses to six
71 drivers that are common agents of change in the world's estuarine-coastal systems: consumption
72 and diversion of fresh water, modification of sediment supply, introduction of non-native
73 species, sewage input, environmental policy and climate shifts.

74

75 **2. DESCRIPTION OF THE ESTUARY**

76 San Francisco Bay is an estuary, a coastal bay where seawater is measurably diluted with fresh
77 water from land drainage [*Pritchard, 1967*]. Seawater enters through the narrow deep channel at
78 the Golden Gate (Figure 2), and its chemical and biological constituents are influenced by
79 seasonal upwelling in the adjacent coastal boundary current. Fresh water is delivered primarily
80 by the Sacramento and San Joaquin rivers, which carry runoff produced in the 163,000-km²
81 watershed bounded by the Cascade and Sierra Nevada mountains. Annual runoff is highly
82 variable (Figure 3). During the last century, for example, annual runoff ranged from a low of 7.6
83 km³ in 1977 to a high of 65 km³ in 1983, both El Niño years. Runoff is also highly seasonal,
84 reflecting a climate of wet winters and dry summers.

85 San Francisco Bay is the defining geographic feature of the "Bay Area", home to 7.5
86 million people (Figure 4). California's urban population centers and agricultural production are
87 largely dependent upon water diverted from the estuary. The Bay moderates regional climate,

88 assimilates wastewater from 50 municipal sewage treatment plants [*van Geen and Luoma, 1999*],
89 is a center of commercial shipping, serves as both nursery and migration route for ocean-
90 harvested fish and crabs, and includes the largest tidal wetland restoration project in the western
91 USA [*Thebault et al., 2008*]. San Francisco Bay supports 30% of shorebird populations and up to
92 half of diving duck populations in the Pacific Flyway [*Takekawa et al., 2001*]. The Bay and its
93 tributary rivers and wetlands provide habitat for threatened and endangered species of fish
94 (Chinook salmon *Oncorhynchus tshawytscha*; steelhead trout *Oncorhynchus mykiss*; delta smelt
95 *Hypomesus transpacificus*; longfin smelt *Spirinchus thaleichthys*) and birds (western snowy
96 plover *Charadrius alexandrinus nivosus*; California clapper rail *Rallus longirostris obsoletus*).
97 This estuary has been radically transformed by human actions that began soon after the 1848
98 discovery of gold in California and included, for example, near-complete (95%) diking and
99 filling of tidal marsh habitat [*Nichols et al., 1986*]. As in all the world's estuarine-coastal
100 ecosystems, changes continue in response to human disturbances and climatic variability.
101 Understanding the drivers of these changes requires a broad landscape perspective from
102 mountains to ocean because, as we show, processes of change originate far into the watershed,
103 within the Bay, and in the Pacific Ocean.

104 The San Francisco Bay system (Figure 2) comprises the North Bay (including Suisun and
105 San Pablo Bays) -- a partially stratified estuary of the Sacramento-San Joaquin Rivers; and the
106 South Bay -- a marine lagoon situated in a densely populated urban setting. We use observations
107 from Suisun Bay, just downstream of the confluence of the Sacramento and San Joaquin rivers,
108 as an example of an estuarine system strongly influenced by seasonal and annual fluctuations of
109 runoff from an agricultural watershed. We use observations from South Bay as an example of a

110 marine lagoon strongly influenced both by inputs from an urban landscape and connectivity to a
111 coastal ocean.

112 Suisun Bay is a turbid, low-salinity embayment with high nutrient (N, P, Si)
113 concentrations but low phytoplankton biomass (chlorophyll *a*) and primary production (Table 1).
114 South Bay is a larger nutrient-enriched embayment with higher salinity, lower turbidity and
115 higher phytoplankton biomass and primary production. Both embayments are broad expanses of
116 intertidal and shallow subtidal habitat incised by a relict river channel. Tidal currents are strong
117 (peak velocity $\sim 175 \text{ cm s}^{-1}$ at the Golden Gate), and tidal amplitude is damped as the tidal wave
118 propagates into North Bay but amplified along the semi-enclosed South Bay [*Walters et al.*,
119 1985]. Water residence time in Suisun Bay ranges from less than a day during large floods to
120 about a month during the dry season, and from weeks to months in South Bay. San Francisco
121 Bay is turbid because of large river inputs of suspended particulate material (SPM), mostly
122 mineral sediments. Unlike Chesapeake Bay and many other nutrient-enriched estuaries, San
123 Francisco Bay is not currently impaired by harmful algal blooms, excessive phytoplankton
124 production or hypoxia (Table 1; but see section 8.2).

125 Water quality and biological communities are sampled regularly in San Francisco Bay by
126 the U.S. Geological Survey (USGS) and the Interagency Ecological Program (IEP), a consortium
127 of state and federal agencies. These research and monitoring programs are motivated by the
128 common needs of resource managers and policy makers around the world's coastlines to
129 understand how environmental changes are brought about by climate variability and human
130 disturbance. These sampling programs have been sustained over four decades, providing one of
131 the longest and most comprehensive records of environmental and biological variability in a US
132 coastal ecosystem. Combined sampling by USGS, IEP and other agencies has produced a

133 valuable observational record for capturing large environmental changes as trends over time or
134 abrupt shifts, and for identifying their underlying causes. We use these records to illustrate six
135 drivers of change that are common in marine systems influenced by connectivity to land. The
136 next six sections follow a common format of: background information about a specific driver of
137 change, measured responses to that driver in San Francisco Bay, and discussion of the
138 significance of those changes from both a local and global perspective.

139

140 **3. WATER CONSUMPTION AND DIVERSION**

141 **3.1 Background**

142 The explosive population growth and economic development that began in California after the
143 19th century gold rush [*Nichols et al.*, 1986] required a stable water supply. That supply was met
144 with construction of a massive infrastructure that includes reservoirs to capture water produced
145 by runoff during the wet winter-spring, and canals to carry that water from the humid northern
146 region of California to the drier south where 75% of water demand is concentrated. This
147 infrastructure provides flood protection and water for urban centers and California's agricultural
148 industry, which annually produces crops valued at \$36 billion [*USDA*, 2010]. The two major
149 systems of water infrastructure are the Central Valley Project (CVP) operated by the U.S. Bureau
150 of Reclamation and the State Water Project (SWP) operated by the California Department of
151 Water Resources (Figure 2 shows their delivery facilities exiting the Delta). Although the CVP
152 and SWP storage facilities account for more than half the upstream water storage capacity, many
153 other upstream facilities and numerous small water users also affect water supply to San
154 Francisco Bay [*Arthur et al.*, 1996].

155 Construction of the CVP began with Friant Dam on the San Joaquin River in 1942, and

156 the last major facility constructed was the New Melones Dam on the Stanislaus River, a tributary
157 of the San Joaquin, in 1979 (Figure 5). The largest CVP reservoir is Lake Shasta (5.62 km³) on
158 the Sacramento River, formed by Shasta Dam in 1945. While the CVP impounds water from five
159 major rivers (American, Sacramento, San Joaquin, Stanislaus and Trinity), the SWP draws
160 primarily from the Feather River, the main tributary of the Sacramento. The largest SWP
161 reservoir (4.36 km³) is Lake Oroville, formed by Oroville Dam in 1968. Every large river in the
162 Sierra Nevada (except the Cosumnes River) has a large terminal storage reservoir. The
163 cumulative capacity (27 km³) of the 10 largest reservoirs mediating flow into San Francisco Bay
164 is approximately the same as the (1906-2010) median annual runoff in the Sacramento-San
165 Joaquin drainage (Figure 5). The CVP and SWP are two of the largest water diversions in the
166 world, and we show how their operations have changed the quantity and seasonal pattern of
167 freshwater inflow to San Francisco Bay.

168

169 **3.2 Reduced amount and altered timing of freshwater inflow**

170 The net outflow from the Delta is the most important freshwater input to San Francisco Bay.
171 Delta outflow is what remains of Delta inflow after exports to various water projects and
172 depletions within the Delta. Measurements of outflow, inflow, exports and depletions are
173 available for water years 1956-2010 (Table A1e). In addition, we can estimate *unimpaired* Delta
174 inflow, which is runoff that would have occurred had water flow remained unaltered in rivers
175 and streams upstream of the Delta instead of being stored in reservoirs, imported, exported or
176 diverted [CADWR, 2007]. We used these particular estimates of unimpaired inflow, even though
177 they are available only through 2003, because they include accretions from the Sacramento and
178 San Joaquin valley floors in addition to runoff from higher elevations. The difference between

179 measured and unimpaired Delta inflow provides an estimate of the upstream effect on water
180 losses, which can then be compared to the Delta effect. During 1956-2003, a median 61% of
181 unimpaired inflow from the watershed flowed out of the Delta into Suisun Bay (Figure 6), while
182 upstream and Delta effects accounted for 21% and 13%, respectively.

183 Although the upstream effect exhibited no long-term trend for 1956-2003, both exports
184 and Delta outflow changed systematically during this period. The trend in the Delta effect ($+4.0$
185 $\text{m}^3 \text{s}^{-1} \text{yr}^{-1}$, $p < 0.001$) essentially mirrored that in outflow ($-3.6 \text{m}^3 \text{s}^{-1} \text{yr}^{-1}$, $p < 0.001$). The Delta
186 effect therefore increased over time, at the expense of outflow to San Francisco Bay. The trend in
187 Delta effect is due to a trend in water exports from the Delta (also $+4.0 \text{m}^3 \text{s}^{-1} \text{yr}^{-1}$, $p < 0.001$), as
188 opposed to within-Delta depletion that contributes a median of only 19% of the Delta effect and
189 has no long-term time trend. The long-term increase in exports, from approximately 5% to 30%
190 of Delta inflow, is obvious in Figure 7, and it appears to end by the 1990s.

191 Delta inflow for the months July and August increased significantly from 1956 to 2010
192 (Figure 8A). Presumably, this is the result of storage-and-release patterns from impoundments
193 upstream of the Delta, which store excess supply from spring runoff and postpone releases until
194 drier summer conditions [*Knowles*, 2002]; there was no change in the seasonal pattern of
195 unimpaired Delta inflow for 1956-2003. Monthly exports from the Delta also increased every
196 month except May (Figure 8B). Because of the high inflow typical of winter months there was
197 no detectable change in Delta outflow during January-April, and the enhanced supply from
198 upstream ameliorated any effect of exports on Delta outflow during July-August. During
199 September-December, however, changes in the upstream supply no longer compensated for
200 increased export losses and, as a result, outflow from the Delta to San Francisco Bay declined
201 (Figure 8C). As in the case of annual exports (Figure 7), the downward trend in Delta outflow for

202 September-December was nonlinear, and essentially over by about 1990. Stabilization of the
203 water export ratio reflects a 1994 policy agreement on Bay-Delta environmental protection (Bay-
204 Delta Accord) between government agencies and diverse stakeholders to set monthly quotas for
205 water export [*CALFED*, 2012].

206 In response to these September-December flow trends, salinity now moves further
207 upstream during the latter part of the calendar year. The salinity gradient of North San Francisco
208 Bay can be characterized by X_2 , the distance (km) from the Golden Gate where near-bottom
209 salinity is 2 [*Jassby et al.*, 1995]. We determined X_2 from Delta outflow using a steady-state
210 model [*Monismith et al.*, 2002]. As implied by the negative trend in September-December
211 outflow from the Delta (Figure 8C), there is a corresponding positive trend in September-
212 December X_2 , i.e., an increase in the autumn intrusion of salinity into the estuary. Table 2
213 illustrates this trend using decadal averages of actual X_2 based on Delta outflow and
214 “unimpaired” X_2 based on unimpaired Delta outflow [*CADWR*, 2007], for September-
215 December. The average difference ΔX_2 between them was negative through the 1970s,
216 indicating how reservoir operations initially shifted the water supply to San Francisco Bay from
217 the earlier wet to the later dry months. But exports from the Delta eventually dominated and
218 salinity intrusions during September-December have become greater than they would have been
219 under unimpaired conditions, i.e., $\Delta X_2 > 0$ (Table 2). The estuarine salinity gradient has thus
220 been displaced landward relative to unimpaired conditions. Despite these overall trends, there is
221 high variability from year to year within each decade. In fact, interannual outflow variability
222 may be larger now than in pre-European times when flows were dampened by large wetland and
223 floodplain areas [*Enright and Culberson*, 2010].

224

225 **3.3 Significance of the changes**

226 Flow management in the San Francisco Bay-Delta watershed is so pronounced that a median
227 39% of its unimpaired runoff is consumed upstream or diverted from the estuary (Figure 6), and
228 the Sacramento-San Joaquin River system is thus classified as “strongly affected” by
229 fragmentation [*Dynesius and Nilsson, 1994*]. Responses to this fragmentation include annual
230 exports sometimes exceeding 50% of inflow (Figure 7A), shifts in the seasonal hydrograph
231 (Figure 8), and a landward displacement of the estuarine salinity gradient during autumn (Table
232 2). The era of increasing water exports from the Sacramento-San Joaquin Delta (Figure 7) has
233 been marked by population declines of native aquatic biota across trophic levels from
234 phytoplankton [*Alpine and Cloern, 1992*] to zooplankton [*Winder et al., 2011*] to pelagic fish
235 [*Sommer et al., 2007*], and large shifts in biological communities [*Winder and Jassby, 2011*].
236 These signs of ecosystem disturbance are related, at least partly, to altered flow regimes from
237 water consumption and exports [*Bennett, 2005; NRC, 2010; Sommer et al., 2007*]. Attribution of
238 specific biological changes to flow modification is difficult because of data gaps (water exports
239 began before biological monitoring), and confounding effects of other drivers of change such as
240 climate variability, pollutant inputs, introductions of non-native species and landscape
241 modifications [*Mac Nally et al., 2010*]. However, modifications of inflow and salinity are
242 contributing factors to population declines of native species in low-salinity habitats of the San
243 Francisco Bay system [*Moyle et al., 2010*] and to the remarkably successful establishment of
244 non-native species [*Winder et al., 2011*], including species that have restructured food webs and
245 their productivity [*Winder and Jassby, 2011*]. Water export from the Sacramento-San Joaquin
246 Delta is a direct source of mortality to fish, including imperiled species such as delta smelt and
247 longfin smelt [*Grimaldo et al., 2009; NRC, 2010*], and export plus within-Delta depletion alters

248 system energetics of an already low-productivity ecosystem by removing phytoplankton biomass
249 equivalent to 30% of Delta primary production [*Jassby et al.*, 2002]. Reduced autumn inflows
250 and associated salinity increases (Table 2) have lowered habitat quantity and quality for species
251 endemic to the upper estuary, such as the endangered delta smelt [*Feyrer et al.*, 2011].

252 These linkages between water diversion and sustainability of native fishes and their
253 supporting food webs are now recognized in policy, first through creation of an ambitious
254 biological conservation plan having co-equal goals of water supply reliability and ecosystem
255 restoration [*BDCP*, 2010]. Second, California’s State Water Resources Control Board recently
256 determined that current flows to the San Francisco Bay-Delta “are insufficient to protect public
257 trust resources” and proposed flow criteria based on its conclusion that “[f]low modification is
258 one of the few immediate actions available to improve conditions to benefit native species”
259 [*SWRCB*, 2010].

260 Outcomes of this policy recommendation are uncertain, but the San Francisco Bay
261 example illustrates the extent to which humans have modified hydrologic systems and the global
262 challenge of measuring and balancing the societal benefits and environmental costs of different
263 water management actions and policies. At least 90% of total river discharge in the USA is
264 strongly affected by channel fragmentation from reservoir operations, interbasin diversions and
265 irrigation consumption [*R B Jackson et al.*, 2001]. Flow management has had particularly large
266 effects at middle latitudes, where the cumulative discharge of many rivers declined 60% [*J D*
267 *Milliman et al.*, 2008] in the 1951-2000 era of accelerated dam construction and irrigation, which
268 dominates US water use [*Gleick and Palaniappan*, 2010]. Large-scale fragmentation of river
269 systems has been a significant disturbance to estuarine-coastal ecosystems. Iconic examples
270 include: extensive losses of wetlands and bivalve mollusks in the Gulf of California after

271 completion of the Glen Canyon Dam on the Colorado River [*Baron et al.*, 2002]; collapse of
272 Egypt's coastal fishery when Nile flows to the Mediterranean Sea were reduced by 90% after
273 completion of the Aswan Dam [*Nixon*, 2003]; salinity increase and restructuring of Florida Bay's
274 seagrass and fish communities after freshwater inflows from the Everglades were reduced by
275 60% [*Herbert et al.*, 2011]; and greatly reduced nutrient supply, primary production, diversity
276 and biomass of fish communities in the Bohai Sea after Huanghe (Yellow) River discharge was
277 reduced 73% between the 1950s and 1990s [*Fan and Huang*, 2008].

278

279 **4. HUMAN MODIFICATION OF SEDIMENT SUPPLY**

280 **4.1 Background**

281 One of the first scientific investigations of San Francisco Bay was published as a USGS
282 Professional paper by G. K. Gilbert [*Gilbert*, 1917]. This remarkably detailed and comprehensive
283 study included measurements in San Francisco Bay, its tributary rivers and their watersheds to
284 assess impacts of hydraulic gold mining in the Sierra Nevada Mountains on sediment supply to
285 and deposition in San Francisco Bay. Gilbert's conclusion was startling and accurate:
286 mobilization of sediments by hydraulic mining during the period 1849-1914 delivered nearly a
287 billion cubic meters of sediments to San Francisco Bay. A comparison of the 1856 and 1887
288 bathymetric surveys of San Pablo Bay (Figure 2) confirmed that the estuary accumulated
289 sediments during this period, when some regions filled by more than 4 m and intertidal mudflats
290 expanded 60% [*Jaffe et al.*, 2007]. Hydraulic mining was prohibited in 1884, and the late 19th
291 century era of sediment deposition was followed by a gradual shift to the current state of San
292 Francisco Bay as an erosional system [*Jaffe et al.*, 2007]. This shift was driven by multiple
293 processes including erosion of the hydraulic-mining debris deposited in the river system, diking

294 the rivers and disconnecting them from floodplains, and retention of sediments behind large
295 dams constructed in the 20th century (Figure 5) [*Schoellhamer, 2011; Wright and Schoellhamer,*
296 2004]. As a response to these changes in the watershed, sediment supply to the estuary has been
297 halved since the mid 20th century [*Wright and Schoellhamer, 2004*]. Sediment supply peaked at
298 about 12 Mt yr⁻¹ in the late 19th century, then declined as the era of large-dam building
299 progressed and is now <1 Mt yr⁻¹ [*Schoellhamer, 2011*]. Changes in sediment supply of this
300 magnitude have reshaped San Francisco Bay's geomorphology [*Jaffe et al., 2007*]. We focus
301 here on another important consequence -- reduced concentrations of suspended particulate matter
302 (SPM) and turbidity in the upper estuary.

303

304 **4.2 Decreasing sediment concentrations**

305 We chose two sampling locations in Suisun Bay to illustrate SPM trends (Figure 9): D7 in the
306 center of Grizzly Bay, a shallow subembayment, and D8 in the deep channel (Figure 2). Both
307 sites have been sampled at about 1 m depth approximately monthly from 1975 through 2010. We
308 calculated trends in water-year mean SPM concentration, using a model to separate out the
309 variability due to year-to-year changes in flow. This effect of flow on water quality constituents
310 is often separated out with an additive model that includes terms for both flow and long-term
311 trend [*Cohn et al., 1992*]. For our application, net Delta outflow was used as the flow variable
312 and water year was used as the trend variable. Rating curves for SPM are often nonlinear, and
313 our preliminary exploration suggested a nonlinear transform for flow within the additive model.
314 Both SPM and outflow also required log-transformation to ensure normality of residuals. The
315 resulting model is:

316

317 (1) $\ln M = c_0 + c_1 T + s(\ln Q_{\text{out}}) + \varepsilon$

318

319 where for each water year T , M is the mean SPM concentration (mg L^{-1}), Q_{out} is the mean
320 outflow ($\text{m}^3 \text{s}^{-1}$), c_0 and c_1 are constant coefficients, s represents a natural spline, and ε is the
321 residual. We summarize the fit to Equation 1 graphically for site D8 (Figure 10).

322 Both water year (long-term trend) and outflow are significant sources of SPM variability,
323 and the effect sizes of trend and outflow are similar (Figure 10). The outflow effect on SPM is
324 positive and reaches a maximum between 500 and 1000 $\text{m}^3 \text{s}^{-1}$. There is typically an estuarine
325 turbidity maximum in this embayment, and gravitational circulation is strongest in this outflow
326 range. The trend effect is equivalent to an annual SPM loss of 2.0% yr^{-1} . With outflow set to its
327 long-term median, the model implies that SPM concentrations dropped from 54 mg L^{-1} in 1975
328 to 27 mg L^{-1} in 2010, consistent with regional trends of declining turbidity in the upper estuary
329 [*W Kimmerer, 2004*]. Results for D7 are qualitatively similar (71 to 46 mg L^{-1}), although the
330 annual SPM loss was only 1.2% yr^{-1} , perhaps reflecting the greater importance of resuspension at
331 this shallow site. Loss rates of SPM concentration in Suisun Bay compare to a 1.3% yr^{-1} decline
332 of sediment supply during the last half of the 20th century [*Wright and Schoellhamer, 2004*].

333

334 **4.3 Significance of the changes**

335 The reduced sediment supply to San Francisco Bay (Figure 10A) has important ecological
336 implications for this estuary, including changes in the transport of sediment-bound contaminants,
337 exposure of legacy contaminants (e.g., mercury; see Section 7.3) as surface sediments continue
338 to erode [*Jaffe et al., 2007*], and a “bleak prognosis” for long-term sustainability of tidal marshes
339 in this urban setting where marshes cannot migrate upland to accommodate anticipated sea level

340 rise and low sediment supply [Stralberg *et al.*, 2011]. Here we consider implications of
341 decreasing SPM concentrations on turbidity and light availability to primary producers, an
342 under-explored response to human modifications of sediment supply to estuaries. San Francisco
343 Bay has high nutrient concentrations (see section 6), but Suisun Bay has unusually low
344 phytoplankton biomass (Table 1) because of fast water filtration by clams (see section 7.2) and
345 turbidity from high SPM concentrations leading to light limitation of photosynthesis [Alpine and
346 Cloern, 1992]. What, then, are implications of a 50% reduction of SPM concentration (Figure 9)
347 for primary productivity? To address this question we developed a multiple regression model
348 relating the attenuation coefficient for photosynthetically active radiation, k (m^{-1}), to SPM
349 concentrations and salinity. The model was linear in SPM and salinity, the latter serving as a
350 proxy for dilution of terrestrial-derived, colored dissolved organic matter [Twardowski and
351 Donaghay, 2001]. Setting salinity to its median of 5.1 during 1975-2010, and substituting flow-
352 adjusted values for SPM in 1975 and 2010, the model implies a drop in flow-adjusted k from
353 3.67 to 2.35 m^{-1} in the deep channel of Suisun Bay (D8). This implies a corresponding increase
354 in photic-zone depth from 1.3 to 2.0 m. Similar calculations for the shallow subembayment (D7)
355 yield an increase in photic-zone depth from 1.1 to 1.5 m.

356 This decrease in light attenuation has a direct effect on primary production.
357 Phytoplankton primary productivity in San Francisco Bay [Cole and Cloern, 1984] and the
358 upstream Delta [Jassby *et al.*, 2002] are well described with a simple model of biomass and light
359 availability:

360

361 (2) $P_g = \psi B I_0 z_p$

362

363 where P_g ($\text{mg C m}^{-2} \text{d}^{-1}$) is gross primary productivity, i.e., the photosynthetic incorporation rate
364 of carbon by phytoplankton beneath a square meter of water surface; ψ ($\text{mg C [mg Chl a]}^{-1}$
365 [$\text{Einstein m}^{-2} \text{J}^{-1}$]) is an efficiency factor; B ($\mu\text{g L}^{-1}$) is chlorophyll *a* concentration; I_0 (Einstein
366 $\text{m}^{-2} \text{d}^{-1}$) is incident photosynthetically active radiation (PAR); and z_p is photic-zone depth (m),
367 the depth at which PAR falls to 1% of its surface value I_0 . Given that primary productivity is
368 proportional to z_p , the increases in z_p calculated here imply a 54% increase in phytoplankton
369 productivity per unit biomass (P_g/B) at the channel site D8 and a 38% increase at the shallow site
370 D7, apart from any changes in ψ (cf. [Parker *et al.*, 2012]).

371 Turbid estuaries are inherently low-productivity ecosystems [Cloern, 1987]. Annual net
372 primary production measured in the Suisun Bay channel during 1980 was only $80 \text{ g C m}^{-2} \text{yr}^{-1}$,
373 compared to $160 \text{ g C m}^{-2} \text{yr}^{-1}$ in the South Bay channel where SPM concentrations are lower
374 [Cloern *et al.*, 1985]. However, Suisun Bay waters have become more transparent as SPM
375 concentrations declined (Figure 9), so the water-column mean irradiance to support
376 photosynthesis has increased significantly since those measurements were made. This illustrates
377 an unintended consequence of river impoundment -- reduced sediment supply to estuaries,
378 leading to smaller SPM concentrations, deeper light penetration and increased light availability
379 to primary producers.

380 Changes in water clarity have effects on other biological communities. Abundance of
381 delta smelt has fallen to critically low values, and sustainability of this endemic species is a
382 priority management goal. Delta smelt are most abundant in turbid, low-salinity habitats and
383 their association with turbidity may be an adaptation to minimize predation risk [M.L. Nobriga *et*
384 *al.*, 2008]. An index of delta smelt habitat suitability declined 78% from 1967 through 2008 as a
385 response to trends of increasing salinity (Section 3.2) and water transparency. Reduced habitat

386 quantity and quality are important factors contributing to population declines of this endangered
387 species [Feyrer *et al.*, 2011]. On the other hand, increasing water clarity has expanded habitat
388 area for rooted macrophytes including the non-native *Egeria densa*, introduced in the 1940s and
389 a target for removal as an invasive pest [Santos *et al.*, 2009]. *Egeria* now dominates shallow
390 regions of the upstream Delta, and its expansion provides increasing habitat for non-native fish,
391 such as centrarchids (bluegill, sunfish), whose populations have grown since the 1980s [L.R.
392 Brown and Michniuk, 2007]. The trend of diminishing sediment supply therefore has important
393 ecological implications through its effect on turbidity, light availability and photosynthesis of
394 primary producers, and habitat suitability for native and non-native plant and fish species.

395 Over half of the world's large river systems (172 of 292) are affected by dams [Nilsson *et*
396 *al.*, 2005]. Many of the world's major rivers have experienced similar drops in sediment
397 discharge due to dams, followed by ecological effects that are long-lived and significant.
398 Sediment discharge of the Yangtze River dropped from 490 to 150 Mt yr⁻¹ after closure of Three
399 Gorges Dam (Figure 1A), and the estuary downstream has become sediment-starved with
400 corresponding submersion of salt marshes and erosion of the coastal delta [Yang *et al.*, 2011].
401 The largest coastal wetland loss in the US is the 25% loss of Mississippi Delta wetlands after
402 dam construction in the upper watershed reduced sediment supply to the lower Mississippi River
403 from 400-500 MT yr⁻¹ to 205 MT yr⁻¹ [Blum and Roberts, 2009]. Louisiana's coastal wetlands
404 are now sediment-starved and, with subsidence and accelerating sea level rise, a further 10,000-
405 13,500 km² are projected to be submerged by 2100 [Blum and Roberts, 2009]. On the Nile River,
406 the Aswan Dam has limited the amount of nutrient-rich sediments reaching the delta and
407 negatively affected both agriculture and the functioning of coastal ecosystems [Hamza, 2009].
408 Dams on the Ebro River in Spain trap almost all suspended sediment and bedload in reservoirs,

409 causing ongoing riverbed incision downstream [*Vericat and Batalla, 2006*]. Worldwide, nearly a
410 third of sediment moving from land to the world oceans is now trapped behind dams [*Syvitski,*
411 2003] and, as a result, coastal wetlands are subsiding and river deltas are eroding. For regulated
412 basins, more than half of the sediment is trapped [*Vörösmarty et al., 2003*]. However, worldwide
413 sediment discharge may actually have been only half its current level before widespread
414 agriculture and deforestation began two to three millenia ago [*J Milliman and Syvitski, 1992*]

415

416 **5. INTRODUCED SPECIES**

417 **5.1 Background**

418 Accelerating globalization of commerce has had the unintended consequence of translocating
419 species of microbes, plants and animals, and human redistribution of life forms on Earth is now
420 recognized as a powerful component of global environmental change [*Vitousek et al., 1996*].
421 Biological invasions challenge the integrity of natural plant and animal communities and
422 confound conservation plans to preserve endangered species. The most important vector for
423 transferring marine species is movement of ship ballast water that is usually taken from and
424 discharged into bays and estuaries. US ports alone receive > 79 million tons of ballast water
425 annually from foreign ports [*Ruiz et al., 1997*]. As a result, the world's bays, estuaries and inland
426 waters with deep-water ports are described as marine analogs of highly invaded oceanic islands
427 and among the most threatened ecosystems on the planet [*Carlton and Geller, 1993*]. San
428 Francisco Bay stands out as a coastal ecosystem transformed by introduced species that
429 contribute up to 97% of the individuals and 99% of the biomass of some communities. The rate
430 of biological invasions is accelerating, and estimated at one new species introduced to the San
431 Francisco Bay-Delta system every 14 weeks from 1961 through 1995 [*Cohen and Carlton,*

432 1998]. As a result, this “may be the most invaded estuary and possibly the most invaded aquatic
433 ecosystem in the world” [Cohen and Carlton, 1998]. We describe here one of the most far-
434 reaching of these invasions: a restructuring of the Suisun Bay planktonic food web following
435 introduction of the non-native clam *Corbula amurensis* that quickly established itself as a
436 ‘keystone’ species. Regular sampling provided early detection of the clam’s arrival and
437 measurement of its rapid dispersal, making this one of the best-documented invasions of any
438 estuary [Carlton *et al.*, 1990].

439

440 **5.2 Restructured planktonic food web**

441 *Corbula amurensis* is a small clam native to rivers and estuaries of east Asia. It was first
442 discovered in Suisun Bay in October 1986 and was probably introduced as larvae discharged in
443 ship ballast water [Carlton *et al.*, 1990; Nichols *et al.*, 1990]. By summer 1988 *Corbula*
444 dominated the benthic community, exceeding 95% of the total in both numbers and biomass, and
445 it has reached abundances as high as 16,000 individuals m⁻² and biomass (ash-free dry weight) as
446 high as 131 g m⁻² [Chauvaud *et al.*, 2003]. The remarkably fast colonization and dominance of
447 the Suisun Bay benthos by *C. amurensis* is attributed to its capacity to utilize a broad range of
448 food resources [Parchaso and Thompson, 2002] and its adaptability to a wide range of salinities,
449 including tolerance of salinity < 1 [Nichols *et al.*, 1990]. The annual freshening of Suisun Bay
450 during the wet season precludes colonization by marine bivalves, so *C. amurensis* rapidly
451 occupied and filled a vacant niche. Abundance of this clam has fluctuated markedly since its
452 establishment (Figure 11A), and the single most prominent reason is salinity variability in
453 response to inflow changes [H A Peterson and Vayssières, 2010]. In particular, clam abundance
454 in Suisun Bay tends to increase as X2 shifts upstream (Section 3.2) [Nichols, 1985; Nichols *et*

455 *al.*, 1990; *Winder et al.*, 2011].

456 *Corbula amurensis* is a suspension-feeding bivalve that efficiently assimilates
457 phytoplankton cells [*Cole et al.*, 1992; *Werner and Hollibaugh*, 1993]. Its annual mean
458 filtration rate of Suisun Bay ($\sim 0.1\text{-}0.25\text{ d}^{-1}$) is about twice the growth rate of phytoplankton
459 ($\sim 0.05\text{-}0.1\text{ d}^{-1}$), so clam consumption exceeds local production of phytoplankton biomass
460 [*Thompson*, 2005]. As a result, average chlorophyll *a* concentration decreased abruptly after the
461 clam introduction (Figure 11B), from $11\pm 2\ \mu\text{g L}^{-1}$ during 1975-1986 to $2.2\pm 0.2\ \mu\text{g L}^{-1}$ during
462 1987-2010. This biomass drop is ecologically significant because chlorophyll *a* concentrations of
463 about $10\ \mu\text{g L}^{-1}$ represent a threshold below which zooplankton reproduction can become food
464 limited [*W J Kimmerer et al.*, 2005; *Mueller-Solger et al.*, 2002]. Introduction of *C. amurensis*
465 changed the seasonal pattern of phytoplankton biomass because its grazing effect is strongest
466 during summer (Figure 12). Prior to the introduction, Suisun Bay sustained high phytoplankton
467 biomass, usually $>10\ \mu\text{g chlorophyll } a\text{ L}^{-1}$, during May-September when freshwater inflow is
468 low and residence time is long enough for biomass to accumulate [*Cloern et al.*, 1983]. Since the
469 introduction, chlorophyll *a* concentration is now regularly $< 3\ \mu\text{g L}^{-1}$, even during the low-flow
470 season (Figure 12).

471 Once established, *Corbula amurensis* quickly transformed Suisun Bay by reducing
472 phytoplankton biomass and primary production five-fold [*Alpine and Cloern*, 1992], redirecting
473 much of the remaining primary production from pelagic (zooplankton) to benthic (clam)
474 consumers [*Thompson*, 2005], and creating a persistent state of low phytoplankton biomass and
475 potential food limitation of herbivorous zooplankton. Other sources of organic matter are
476 available to fuel production in food webs. The largest source to San Francisco Bay is river input
477 of detritus [*Jassby et al.*, 1993], but this is largely refractory and the labile components must be

478 converted through the inefficient microbial loop into forms accessible to zooplankton [*Jassby*
479 *and Cloern, 2000; Mueller-Solger et al., 2002; Sobczak et al., 2002*]. This pathway has also been
480 disrupted because *C. amurensis* consumes all components of the microbial loop, including
481 bacteria [*Werner and Hollibaugh, 1993*], ciliates and flagellates [*Greene et al., 2011*].
482 Introduction of a non-native clam therefore reduced the microplankton food resource available to
483 zooplankton and forced a shift toward their greater reliance on low-quality detritus [*Jassby,*
484 2008].

485 These fundamental changes at the base of the food web provoked a cascade of responses,
486 beginning with abrupt population declines of some zooplankton species [*W Kimmerer et al.,*
487 1994]. Average abundance of the rotifer *Synchaeta bicornis* decreased from 23,500±6,700 to
488 1,600±360 individuals m⁻³ (Figure 11C). Average abundance of the calanoid copepod
489 *Eurytemora affinis* dropped from 700±140 to 35±11 individuals m⁻³ (Figure 11D), and
490 abundance of another calanoid copepod, *Acartia spp.*, also declined sharply after 1987 [*W*
491 *Kimmerer, 2004*]. Average abundance of the mysid shrimp *Neomysis mercedis* dropped from
492 32±10 to 2.5±2.1 individuals m⁻³ and this species has virtually disappeared after a temporary
493 population rebound in 1993 (Figure 11E). Near-extinction of these previously abundant
494 zooplankton species is attributed to depletion of the phytoplankton food resource and, in the case
495 of *Eurytemora affinis*, predation on its larvae by the introduced clam [*W Kimmerer, 2004*]. These
496 well-documented observations in Suisun Bay before and after colonization by *Corbula*
497 *amurensis* illustrate the power of bivalve mollusks to alter ecosystem production, pathways of
498 energy flow, and food web structure through their predation upon and competition with
499 zooplankton.

500 The abrupt population declines of *Synchaeta bicornis*, *Eurytemora affinis*, *Acartia spp.*

501 and *Neomysis mercedis* (Figure 11C-E) followed population declines of other zooplankton taxa
502 in Suisun Bay that began before the introduction of *Corbula* [W Kimmerer et al., 1994].
503 Comparing mean biomass from the 1970s with the period after 1990, calanoid copepods fell
504 from 14 to 4 $\mu\text{g C L}^{-1}$, rotifers from 10 to 1 $\mu\text{g C L}^{-1}$, and cladocerans from 1.2 to 0.2 $\mu\text{g C L}^{-1}$
505 [Winder and Jassby, 2011]. These taxa were replaced by 8 species of non-native copepods and 2
506 species of non-native mysids that became established in the upper estuary as a sequence of
507 introductions during periods of low freshwater inflow, particularly during the 1987-1992 drought
508 when salinity intrusion facilitated establishment of introduced species adapted to brackish
509 habitat, including *C. amurensis* [Winder et al., 2011]. The cumulative changes since the 1970s
510 have produced a remarkable and perhaps unprecedented transformation of a zooplankton
511 community from one having large components of mysid shrimp, rotifers and calanoid copepods
512 to one dominated by introduced copepods indigenous to east Asia [Winder and Jassby, 2011].
513 This transformation included emergence of smaller cyclopoid copepods that contributed less than
514 2% of zooplankton biomass before 1987 but more than 24% of biomass after 1994.

515

516 **5.3 Significance of the changes**

517 Losses of rotifers, calanoid copepods and mysid shrimp have contributed to the collapses of fish
518 populations in low-salinity regions of San Francisco Bay because these are essential dietary
519 components for resident fish. Rotifers are preferred prey of larval delta smelt [*M. L. Nobriga,*
520 2002], and many planktivorous fish including adult delta smelt, longfin smelt and early life
521 stages of other species selectively prey on calanoid copepods that are larger and have higher
522 nutritional quality than cyclopoid copepods [Winder and Jassby, 2011]. Other species such as
523 American shad, starry flounder and juvenile striped bass feed primarily on mysids when

524 available [Feyrer et al., 2003]. Losses of these zooplankton components provoked adaptations by
525 their fish predators. Fish reliant on mysids shifted their diets to other prey, and those with the
526 largest dietary shifts had the largest population declines in Suisun Bay marshes after 1987
527 [Feyrer et al., 2003]. Northern anchovy (*Engraulis mordax*) is the biomass-dominant pelagic fish
528 in San Francisco Bay; summer abundance of this species fell 94% in the low-salinity region of
529 the estuary as anchovies adapted to the decreased food supply in Suisun Bay by migrating
530 seaward [W J Kimmerer, 2006]. In addition to food web transformations, the five-fold decrease
531 in primary production implies a comparable decrease in the energetic carrying capacity for fish
532 in Suisun Bay based on its primary production [Nixon, 1988].

533 Ecologists struggle to understand how and why some indigenous communities are
534 displaced by non-native species, but the near-complete restructuring of the zooplankton
535 community in low-salinity regions of San Francisco Bay during the past four decades appears to
536 be the synergistic result of multiple drivers [Winder and Jassby, 2011; Winder et al., 2011]:
537 introduction of the clam *Corbula amurensis*; an extended period of low freshwater inflow and
538 salinity intrusion; and amplification of the drought effect by water diversions (Section 3.2).
539 Ecosystem disruptions by species introduced to San Francisco Bay by transoceanic shipping
540 have shaped policy by motivating passage of California's Marine Invasive Species Act,
541 considered the strictest regulation of ship ballast discharge in the USA to prevent or minimize
542 release of nonindigenous species from commercial vessels [Takata et al., 2011].

543 More than 500 non-native species have become established in US coastal waters,
544 and accelerating species introductions rank as one of the "most pervasive threats to native
545 ecosystems and human economies" [Groszholz, 2005]. Brackish estuarine waters are vulnerable
546 not only because so many ports are situated in estuaries and brackish-water species are tolerant

547 of ballast-water tank conditions, but also because brackish waters tend to have fewer indigenous
548 species so that aliens can establish more easily [Wolff, 1998]. Subsequently, the effects of
549 international shipping reverberate up and down coasts as species are introduced into more
550 isolated estuaries by intraregional transport [Wasson *et al.*, 2001]. Alien species can disrupt
551 ecosystems by altering biogeochemical processes (e.g, silica cycling in the Bay of Brest
552 [Chauvand *et al.*, 2000]), by amplifying bioaccumulation of toxic contaminants in food webs
553 (e.g., selenium in San Francisco Bay [Stewart *et al.*, 2004]), and through disruption of ecosystem
554 functions that support native populations. Ecological regime shifts similar to that in Suisun Bay
555 followed introductions of other bivalve mollusks, such as the soft-shell clam *Mya arenaria* to
556 Denmark's Ringkøbing Fjord [Petersen *et al.*, 2008], and zebra mussel (*Dreissena polymorpha*)
557 and quagga mussel (*D. rostriformis bugensis*) introductions to lakes and rivers [Higgins and
558 Vander Zanden, 2010]. In all cases, phytoplankton and zooplankton abundance declined
559 significantly, and biological communities were restructured across multiple trophic levels.

560 Species introductions are the most important cause of bird extinctions and second most
561 important cause of fish extinctions globally [Clavero and Garcia-Berthou, 2005], and as many
562 as 80% of the endangered species in some regions of the world are at risk because of pressures
563 from non-native organisms [Pimentel *et al.*, 2005]. These disruptions have economic costs
564 through loss or reduced productivity of species harvested from coastal waters. For example,
565 extensive losses of the oyster *Crassostrea virginica* along the mid-Atlantic coast of the USA in
566 the 1950s was the result of disease (MSX) caused by the parasite *Haplosporidium nelsoni*
567 introduced from Asia [Carnegie and Burreson, 2011]. Stocks of the most abundant fish species
568 in the Caspian Sea (anchovy kilka) virtually collapsed after 2001 because of predation and food
569 competition from the introduced ctenophore *Mnemiopsis leidyi* [Daskalov and Mamedov, 2007].

570 Given the scope of biodiversity and economic losses from species introductions, coastal
571 scientists now advocate national policies to manage introduced species with the same efforts
572 currently applied to reduce chemical pollution and restore wetlands and fisheries stocks
573 [*Williams and Grosholz, 2008*].

574

575 **6. SEWAGE INPUT**

576 **6.1 Background**

577 San Francisco Bay has been described as “The Urbanized Estuary” [*Conomos, 1979*], reflecting
578 the landscape setting of its South Bay between the cities of San Francisco, San Jose (Silicon
579 Valley) and Oakland (Figure 2). The rate of urbanization accelerated in the mid-20th century as
580 the regional population grew from 2.7 million in 1950 to 7.2 million in 2010 (Figure 4). Humans
581 and their industrial, commercial and agricultural enterprises generate wastes delivered to coastal
582 waters through atmospheric deposition, land runoff, ground water and point sources such as
583 discharges from municipal sewage treatment plants (STPs). Sewage effluent contains an array of
584 pollutants including nutrients (nitrogen and phosphorus), organic matter (and its biochemical
585 oxygen demand, BOD), toxic metals, pharmaceuticals and pathogens that pose risks to human
586 and ecosystem health [*NRC, 1993*]. Nutrient enrichment can provoke excessive production of
587 algal biomass and sustain harmful algal blooms [*Cloern, 2001*]. Metabolism of algal biomass and
588 organic matter from wastewater can deplete water of dissolved oxygen, and the severity and
589 occurrence of hypoxic dead zones are expanding across the world’s coastal waters as a response
590 to anthropogenic nutrient enrichment [*Diaz and Rosenberg, 2008*]. Toxic metals [*Luoma and*
591 *Rainbow, 2008*] and endocrine-disrupting chemicals (e.g., synthetic estrogen) are commonly
592 found in sewage effluent [*Duffy et al., 2009*] and can impair growth, reproduction and immune

593 systems of fish and invertebrates. South San Francisco Bay receives 500,000 m³ of municipal
594 wastewater annually from 12 STPs serving 4 million people [McKee and Gluchowski, 2011]. We
595 use nutrient concentrations as an indicator of how sewage inputs alter the chemistry and potential
596 productivity of this urbanized estuary.

597

598 **6.2 Nutrient Enrichment**

599 Sewage discharges deliver 11,200 tons of dissolved inorganic nitrogen (DIN) and 1860 tons of
600 dissolved inorganic phosphorus (DIP) to South San Francisco Bay annually [McKee and
601 Gluchowski, 2011]. These inputs are 92% and 96% of the combined land-based and atmospheric
602 loadings of DIN and DIP, respectively, and South San Francisco Bay apparently ranks as the US
603 estuary having the largest sewage component of DIN loading (Table 3). The spatial distributions
604 of DIN and DIP concentration show progressive N and P enrichment from the Golden Gate to
605 lower South Bay, reflecting mixing between lower-nutrient ocean water and higher-nutrient Bay
606 water (Figure 13). On an areal basis, STP loadings to South San Francisco Bay are 1860 mmol
607 DIN m⁻² yr⁻¹ and 140 mmol DIP m⁻² yr⁻¹. As an index of the urbanization effect on nutrient input,
608 we compare these with nutrient loads to Tomales Bay, a smaller estuary just north of San
609 Francisco Bay with a similar latitude and climate but situated in a rural watershed. Inputs to
610 Tomales Bay from atmospheric deposition, ground water and surface water inflows are 154
611 mmol DIN m⁻² yr⁻¹ and 6.8 mmol DIP m⁻² yr⁻¹ [S V Smith et al., 1996]. Therefore, sewage input
612 to South San Francisco Bay contributes more than 10 times the total DIN input and more than 20
613 times the total DIP input to Tomales Bay per unit estuary area. As a result, DIN and DIP
614 concentrations are highly elevated in South San Francisco Bay. We show the enrichment effect
615 of wastewater by comparing DIN and DIP concentrations in South San Francisco Bay with those

616 in Tomales Bay and Willapa Bay, another estuary on the US west coast situated in a rural
617 landscape (Figure 14). The median DIN concentration in South San Francisco Bay is enriched
618 7.6- and 10-fold compared to these estuaries not having direct sewage inputs. The median DIP
619 concentration in South San Francisco Bay is 3.8 and 8.8 times the median values in Tomales and
620 Willapa bays, respectively. As a result of its setting in an urban landscape, South San Francisco
621 Bay is highly enriched with sewage-derived nitrogen and phosphorus.

622

623 **6.3. Significance of the changes**

624 The nutrient concentrations in South San Francisco Bay are typically well above those that limit
625 the growth rate of algae. This is illustrated by comparing DIN and DIP concentrations to the
626 half-saturation constants (K_N , K_P) for phytoplankton growth as an index of potential nutrient
627 limitation (Figure 13). Of 4096 DIN measurements made in South San Francisco Bay from
628 1969-2010, only 126 (0.03%) were smaller than the mean K_N for marine diatoms (1.6 μM ;
629 [Sarthou *et al.*, 2005]). Only 1 of 4330 DIP measurements was smaller than the mean K_P (0.24
630 μM ; [Sarthou *et al.*, 2005]).

631 Based on these high N and P concentrations, South San Francisco Bay has the potential to
632 produce phytoplankton biomass at levels that severely impair other nutrient-enriched estuaries,
633 such as Chesapeake Bay, where occurrences of large algal blooms have led to summer hypoxia
634 in bottom waters, loss of submerged vascular plants and alteration of biogeochemical processes
635 such as denitrification [Kemp *et al.*, 2005]. The nitrogen input to South San Francisco Bay from
636 sewage disposal is almost twice the total N input from all sources to Chesapeake Bay and its
637 tributaries (Table 3). As a result, N and P concentrations are substantially higher in South San
638 Francisco Bay than in Chesapeake Bay (Figure 14). However, South San Francisco Bay

639 paradoxically has low phytoplankton biomass relative to other enriched estuaries. The median
640 chlorophyll *a* concentration in South Bay is only 4.1 $\mu\text{g L}^{-1}$ (Table 1), but the potential
641 chlorophyll *a* concentration -- that expected if the median DIN concentration were converted into
642 phytoplankton biomass -- is about 28 $\mu\text{g L}^{-1}$ (assuming a chl-*a*:N ratio of 1 [Gowan *et al.*,
643 1992]). This high-nutrient low-chlorophyll state implies that San Francisco Bay is inefficient at
644 converting nutrients into algal biomass and, therefore, resistant to the harmful consequences of
645 enrichment observed in other estuaries such as Chesapeake Bay (we show in Section 8, however,
646 that this resistance is weakening).

647 San Francisco Bay has (at least) three attributes that confer resistance to the harmful
648 consequences of nutrient enrichment. First, its strong tidal currents generate sufficient turbulence
649 to break down stratification caused by surface heating and freshwater inflow. Chesapeake Bay
650 has weaker tides, weaker turbulent mixing and stratification that persists long enough (months)
651 for bottom waters to become and remain hypoxic or anoxic. Salinity stratification can develop in
652 South San Francisco Bay during weak neap tides, and these stratification events promote fast
653 growth of phytoplankton biomass in the surface layer. But the surface blooms dissipate on the
654 subsequent spring tide when the water column is mixed [Cloern, 1996]. Second, San Francisco
655 Bay is more turbid than Chesapeake Bay because it receives large river inputs of sediments and
656 is shallow, so sediments are maintained in suspension by wind waves and tidal currents [May *et*
657 *al.*, 2003]. As a result, the median light attenuation coefficient in South San Francisco Bay (1.4
658 m^{-1} ; Table 1) corresponds to a photic depth of only 3.3 m and phytoplankton growth rate is
659 limited by low availability of sunlight energy [Cloern, 1999]. Third, accumulation of
660 phytoplankton biomass is controlled by bivalve mollusks (clams, mussels) that can filter a
661 volume of water equal to the South San Francisco Bay volume each day during summer [Cloern,

662 1982]. In Chesapeake Bay, this filter-feeding function was provided historically by an oyster
663 population that could filter that bay's volume in less than 4 d. That filtration time is now
664 hundreds of days because the oyster population has been decimated by overharvest, disease and
665 hypoxia [Kemp *et al.*, 2005]. Comparative analyses of Chesapeake Bay and San Francisco Bay
666 reveal that estuaries have inherent attributes, such as hydrodynamic, optical and biological
667 properties, that control the efficiency with which nutrients are converted into phytoplankton
668 biomass and, therefore, the expression of nutrient enrichment as a driver of environmental
669 change.

670 Nutrient pollution from municipal wastewater is a globally-significant problem that has
671 degraded water quality, reduced biological diversity, and altered biogeochemical functioning of
672 urban coastal areas such as Boston and New York harbors [NRC, 1993], Tampa Bay [Greening
673 *et al.*, 2011], Osaka Bay [Yasuhara *et al.*, 2007], Mersey Estuary [Jones, 2006], Hong Kong's
674 Tolo Harbor [Xu *et al.*, 2011], Rio de Janeiro's Guanabara Bay [Kjerfve *et al.*, 1997], Turkey's
675 Golden Horn Estuary [Tqs *et al.*, 2006] and Australia's Swan-Canning Estuary [Hamilton and
676 Turner, 2001]. Environmental degradation by nutrient over-enrichment has motivated local,
677 national and multinational policies to reduce nutrient inputs from urban and agricultural sources
678 to coastal ecosystems. For example, a goal of the Chesapeake 2000 Agreement is to reduce N
679 and P inputs to Chesapeake Bay by 48% and 53%, respectively [Kemp *et al.*, 2005]. These are
680 similar to goals of multinational agreements to halve nutrient inputs to the Baltic Sea and North
681 Sea [Conley *et al.*, 2002]. The Danish government has enacted even more aggressive plans to
682 reduce N inputs to its aquatic environments by 50% and point sources of P by 80% [Conley *et*
683 *al.*, 2002].

684 The establishment of such quantitative targets for nutrient reduction is a challenging

685 policy application of estuarine science. Early responses of the Dutch Wadden Sea, Chesapeake
686 Bay and Danish fjords to nutrient reduction strategies have not all met the expectations of policy
687 makers [Carstensen *et al.*, 2011]. The contrasting responses of San Francisco Bay and
688 Chesapeake Bay to N and P enrichment teach that nutrient loading rate alone is not a good
689 predictor of algal biomass or the impairments associated with high algal biomass, such as
690 hypoxia and harmful blooms. This lesson appears to be general because a broad range of
691 empirical relationships exists between nutrient (e.g., total N) and chl-*a* concentrations measured
692 in 28 coastal systems, providing “overwhelming evidence that system-specific attributes
693 modulate the response of phytoplankton to nutrient enrichment” [Carstensen *et al.*, 2011]. As
694 explained above, these system-specific attributes go far beyond just hydraulic retention time,
695 noted long ago as a factor differentiating water bodies with respect to nutrient loading
696 [Vollenweider, 1975]. Policies to remediate over-fertilized coastal waters therefore might be
697 most effective and cost-efficient if they are tailored to the attributes of individual estuaries and
698 bays. The urgency for place-based nutrient-reduction strategies will likely accelerate in step with
699 continued urbanization, population and economic growth: global sewage emissions are projected
700 to increase from 6.4 Tg of N and 1.3 Tg of P in 2000 to emissions as high as 15.5 Tg N and 3.1
701 Tg P by 2050, with fastest increases in southern Asia [Van Drecht *et al.*, 2009].

702

703 **7. ENVIRONMENTAL POLICY -- THE US CLEAN WATER ACT**

704 **7.1 Background**

705 In 1972 the US Congress unanimously passed Public Law 92-500, which we know as the Clean
706 Water Act (CWA), to “restore and maintain the chemical, physical and biological integrity of the
707 nation’s waters” and attain “fishable and swimmable waters” across the USA. This landmark

708 legislation established the first federal regulation of sewage disposal by requiring secondary
709 treatment of municipal wastewater to reduce inputs of solids, oxygen-consuming chemicals and
710 pathogens to the nation's waters. The CWA provided funding for construction and improvement
711 of STPs, and it established effluent standards for BOD, suspended solids, fecal coliform bacteria
712 and pH. Enactment of the CWA and similar policies in other countries reflected growing public
713 concern about the accelerating and increasingly visible degradation of water quality caused by
714 municipal and industrial sources of pollution. The Potomac Estuary of "the Nation's River" was
715 an iconic example of environmental degradation from sewage pollution, manifested as noxious
716 algal blooms, hypoxia (Figure 1F), fish kills, loss of water clarity and waters unsafe for
717 swimming because of high counts of fecal coliform bacteria [Jaworski, 1990]. By the 1970s,
718 regions of nutrient-enriched Tampa Bay lost complete benthic communities and half their
719 seagrasses because of low DO and turbidity caused by high algal biomass [Greening and Janicki,
720 2006]. Perhaps the most infamous sign of the state of US water quality in this era occurred in
721 June 1969 when Cleveland's Cuyahoga River ignited because of its flammable pollutants.

722 Pollution effects became increasingly visible in San Francisco Bay during the 20th
723 century as the surrounding population grew (Figure 4). The San Francisco Bay-Delta was once
724 the foremost fishing center on the US west coast, but its commercial fisheries for sturgeon,
725 salmon, striped bass, shad and clams all ended by the 1950s because of habitat degradation,
726 overharvest and poor water quality [S E Smith and Kato, 1979]. Harvest of oysters -- the Atlantic
727 species *Crassostrea virginica* reared primarily in South San Francisco Bay -- was the most
728 valuable California fishery in the late 19th century. But oyster culture ended by the 1930s
729 because of poor growth and condition attributed to urban pollution, and shellfish harvest from
730 San Francisco Bay was quarantined in 1932 to protect human health from water-borne

731 pathogens. At the time of passage of the CWA, fish kills and skin tumors on fish were common
732 in San Francisco Bay, extremes of contamination for toxic metals in clams were equal to or
733 higher than anywhere in the world, and regions of South Bay had seasonal anoxia [*Luoma and*
734 *Cloern, 1982*]. We use data from sampling programs that began before and continued after
735 passage of the CWA to show how these pollution indicators in South San Francisco Bay changed
736 after national and local policies mandated enhanced sewage treatment.

737

738 **7.2 Recovery from sewage-derived pollutants**

739 The first measurements of dissolved oxygen (DO) in San Francisco Bay were made in the late
740 1950s and they showed recurrent summer anoxia in the southernmost region of South Bay
741 (below Dumbarton Bridge, Figure 2). City and regional policies mandated secondary treatment
742 of sewage to reduce BOD inputs to this region even before passage of the CWA. Secondary
743 treatment was fully implemented by all STPs discharging to lower South Bay by 1973. Prior to
744 1973 municipalities discharged untreated or primary treated sewage, and inputs of oxygen-
745 consuming organic matter and ammonium overwhelmed the assimilation capacity of this region.
746 Summer anoxia was eliminated in the 1970s as secondary treatment was implemented, but DO
747 concentrations still fell below 5 mg L^{-1} (Figure 15C), a common standard to protect marine fish
748 sensitive to low oxygen. The Clean Water Act provided incentives for further improvements in
749 wastewater treatment, and by 1980 all STPs discharging into this region implemented processes
750 to remove 99% of BOD and nitrification to convert ammonium into nitrate. From 1978 to 1980,
751 BOD input from the largest STP (San Jose-Santa Clara) dropped from 3700 to 400 t yr^{-1} , and
752 ammonium-N input dropped from 2800 to 40 t yr^{-1} (Figure 15A). In response, hypoxia was

753 eliminated from San Francisco Bay and DO concentrations are now consistently near or above 5
754 mg L⁻¹ (Figure 15C).

755 Environmental effects of sewage-derived metals were detected in the 1970s when
756 sediments and clams (*Macoma balthica*) sampled on a mudflat near the Palo Alto Regional STP
757 discharge were highly contaminated with copper, silver and other metals [*Hornberger et al.*,
758 2000; *Luoma and Cloern*, 1982]. Copper (310 µg g⁻¹) and silver (103 µg g⁻¹) in clam tissues
759 (Figure 15D) reached levels that impaired reproduction; histological analyses confirmed that
760 clams were nonreproductive; and the invertebrate community had low diversity and was
761 dominated by small forms, diagnostics of disturbance by toxic contaminants [*Hornberger et al.*,
762 2000]. Although the primary target of advanced wastewater treatment was removal of BOD,
763 incremental additions of new treatment processes (e.g., biological nutrient removal in the 1990s),
764 combined with industrial pretreatment at the source, were also highly effective at reducing metal
765 inputs from STPs. Annual copper loading from the Palo Alto Regional STP was 5800 kg in 1979
766 but dropped continuously through the 1980s and has been < 300 kg since 1995 (Figure 15B).
767 Annual silver inputs declined from 92 kg in 1989 to < 10 kg since 1995. As loadings decreased,
768 metal contamination of sediments and biota decreased proportionately. By the 1990s copper and
769 silver concentrations in the clam *Macoma balthica* had dropped 10- and 30-fold, respectively,
770 from their peaks of the 1970s (Figure 15D). With greatly reduced metal contamination, clams
771 became reproductive and larger forms of invertebrates re-colonized mudflats near the STP
772 outfall, both evidence that environmental stresses from metals have been greatly reduced since
773 the 1970s. Monthly sampling that began in South San Francisco Bay in 1975 provided one of the
774 first observational records in the US to demonstrate (a) correlation between metal levels in
775 organisms and metal inputs from municipal wastewater, and (b) recovery of physiological

776 impairment and biological communities after metal inputs to an estuary were reduced through
777 advanced wastewater treatment [*Hornberger et al.*, 2000; *Luoma and Cloern*, 1982].

778

779 **7.3 Significance of the changes**

780 In 1950, 40% of US municipal sewage collection systems discharged untreated sewage. By 1996
781 virtually all of the nation's 16,000 STPs were using secondary or advanced treatment. As a
782 result, STP discharge of BOD declined nationwide from 6900 t d⁻¹ to 3800 t d⁻¹ between 1968
783 and 1996, an era when the population served by STPs increased from 140 to 190 million and the
784 BOD influent to STPs increased 35% [*USEPA*, 2000]. The responses documented in San
785 Francisco Bay from measurements of DO and metal contamination before and after
786 implementation of the CWA exemplify the measurable improvements in water quality seen in
787 other US urban estuaries. The first (National Pollutant Discharge Elimination System) permits
788 under the CWA were issued in 1974 to STPs discharging to the Potomac Estuary. Treatment
789 processes targeted removal of BOD and phosphorus. Between 1954 and 1985, BOD loadings
790 decreased from 91,000 to 5400 kg d⁻¹ and P loadings decreased from 10,900 to 270 kg d⁻¹
791 [*Jaworski*, 1990]. In response, phosphorus concentrations in the estuary fell by 80%, algal
792 biomass decreased 60%, DO increased to > 5 mg L⁻¹ (Figure 1F) and fish kills no longer
793 occurred. Elsewhere, coliform bacteria decreased tenfold in the lower Hudson-Raritan Estuary
794 from 1968 to 1993 [*Brosnan and O'Shea*, 1996], and metal (copper, cadmium, nickel)
795 concentrations in the Hudson River estuary decreased 36% to 90% after the 1970s [*Sanudo-*
796 *Wilhelmy and Gill*, 1999]. Seagrass cover expanded more than 2000 ha in Tampa Bay after
797 nitrogen inputs were reduced 60%, and seagrass recovery continues as algal biomass decreases
798 and water clarity increases [*Greening et al.*, 2011]. By these kinds of measures, the Clean Water

799 Act was a highly successful policy to reduce point-source pollution of estuaries through water
800 treatment technology.

801 Policies similar to the CWA were implemented elsewhere to reduce wastewater pollutant
802 loadings at the scale of municipalities (e.g., Hong Kong's Water Pollution Control Ordinance;
803 Perth's Swan Canning Cleanup Program [*Hamilton and Turner, 2001*]; Golden Horn
804 Rehabilitation Project [*Tqs et al., 2006*]), or through national policies (UK Clean Rivers
805 [Estuaries and Tidal Waters] Act [*Matthiessen and Law, 2002*]; Denmark's Action Plan on the
806 Aquatic Environment [*Conley et al., 2002*]), or through multinational agreements such as the
807 European Community's Urban Wastewater Directive [*Hering et al., 2010*]. Perhaps the most
808 publicized rehabilitation was of the river Thames, which historically supported runs of Atlantic
809 salmon but had regions devoid of oxygen and fish from 1920-1964, largely because of sewage
810 inputs from London. Fish, invertebrates and water birds returned to the Thames Estuary after
811 London's sewage works were upgraded in the 1960s, and by 1976 a cumulative total of 112 fish
812 species had returned, including adult salmon for the first time in 140 years [*Attrill, 1998*].
813 Similarly, dissolved oxygen concentrations increased in Victoria Harbor after Hong Kong
814 implemented secondary sewage treatment in 2001 [*Xu et al., 2011*], metal and PCB
815 concentrations in sponges decreased in Cortiou Cove on the French Mediterranean after
816 Marseille implemented primary sewage treatment [*Perez et al., 2005*], and benthic invertebrates
817 and fish returned to Spain's heavily polluted Nervión Estuary [*Borja et al., 2006*] and the UK's
818 Mersey Estuary [*Hawkins et al., 2002*] after metal and BOD inputs were reduced with sewage
819 treatment.

820 These case studies illustrate the success of policies to rehabilitate estuarine-coastal
821 ecosystems from the severe degradation of water quality and disruption of biological

822 communities caused by disposal of untreated municipal waste. However, the standards
823 prescribed in the Clean Water Act have not been fully met. For example, a 2004 assessment of
824 141 US estuaries determined that the majority have moderate or high symptoms of
825 eutrophication expressed as low DO, loss of submerged vascular plants, proliferation of
826 macroalgae, or harmful algal blooms [Bricker *et al.*, 2007]. Nutrient pollution remains the largest
827 pollution problem in US coastal rivers and bays [Howarth *et al.*, 2002]. In some estuaries, such
828 as those of the NE United States [Whitall *et al.*, 2007], sewage input remains the largest source
829 of nitrogen, reflecting the design of secondary sewage treatment to reduce BOD (not nutrients) in
830 effluent. For others, such as Australia's Sydney estuary, stormwater runoff from urban
831 watersheds is the primary source of nutrients and metals [Beck and Birch, 2012]. The largest
832 sources of nitrogen to most estuaries, however, are the diffuse nonpoint sources generated by
833 agriculture and fossil fuel combustion [Howarth *et al.*, 2002]. The risk for coastal eutrophication
834 will likely continue to grow in many world regions because anthropogenic activities will increase
835 river nutrient loading and shift nutrient ratios toward those favoring blooms of harmful algae
836 [Seitzinger *et al.*, 2010].

837 Certain contaminants produced in the past remain in watersheds and persist in estuarine
838 sediments. For example, mercury and PCBs remain priority pollutants in South San Francisco
839 Bay because they are persistent and accumulate in food webs to levels that are health risks to
840 birds, harbor seals and humans [Grenier and Davis, 2010]. Contemporary sources include
841 atmospheric deposition (mercury) and urban runoff (PCBs). Moreover, accelerating erosion -- a
842 consequence of reduced sediment supply (Section 4) -- is now exposing buried sediments having
843 high concentrations as legacy contaminants from the gold-mining era (mercury) and later era of
844 PCB manufacture. A second priority is new contaminants that persist and accumulate in food

845 webs, such as PBDEs (brominated flame retardants). These compounds were undetected in the
846 1980s, but residues are now common in water, sediments and biota of South San Francisco Bay
847 where concentrations in harbor seals, bird eggs and humans are among the highest recorded
848 [*Grenier and Davis, 2010*]. The sources and environmental effects of PBDEs are largely
849 unknown, but their presence illustrates the challenge of maintaining the chemical and biological
850 integrity of estuaries when new contaminants emerge faster than our capacity to identify their
851 sources and assess their effects.

852 The Clean Water Act and similar policies of other countries have greatly reduced inputs
853 of organic matter, pathogens and toxic contaminants to coastal waters, with demonstrable
854 improvements in water and habitat quality. However, standards prescribed in the CWA have not
855 been fully met. Many of the world's estuaries, bays and inland seas are still not fishable and
856 swimmable. Further rehabilitation, or even maintenance of the status quo, will require innovative
857 strategies to solve the much more difficult problems of nonpoint sources of nutrients and toxic
858 contaminants [*T C Brown and Froemke, 2012; R A Smith et al., 1987*], legacy contaminants from
859 the past, and new contaminants of the future.

860

861 **8. SHIFT IN THE OCEAN-ATMOSPHERE SYSTEM**

862 **8.1 Background**

863 Fishermen have known for centuries that fish abundance in the sea fluctuates between eras of
864 good and poor catch that are tied to climate variability. Four centuries of catch records from the
865 English Channel show oscillations of herring and sardine stocks that are synchronized with shifts
866 between cold and warm periods [*Southward et al., 1998*]. More recently, marine and atmospheric
867 scientists have discovered that population fluctuations of fish and their food resources are

868 synchronized with shifting patterns of atmospheric pressure over ocean basins that are
869 represented by climate indices. Recruitment of yellowfin tuna is highest in the tropical Pacific
870 after El Niño events [*Lehodey et al.*, 2006]; salmon stocks in Alaska and California fluctuate
871 inversely with the Pacific Decadal Oscillation (PDO) [*Mantua et al.*, 2002]; cod recruitment in
872 the North Sea is high when the North Atlantic Oscillation (NAO) is positive [*Stige et al.*, 2006].
873 Progress is advancing to understand the linkage mechanisms between fish abundance and these
874 climate patterns through their influence on ocean currents, temperature, primary and secondary
875 production [*Lehodey et al.*, 2006].

876 Discovery of climate-related variability of marine fish populations is rooted in historical
877 observations, including catch records from some fisheries that have been maintained for a
878 century or longer and therefore capture variability over multiple periods of the NAO, PDO and
879 other multidecadal climate patterns. Most observational records in the world's estuarine-coastal
880 systems are much shorter, but they are becoming long enough that we can begin to ask if and
881 how variability in estuaries, bays and lagoons is related to interdecadal shifts in atmospheric
882 forcing across ocean basins. Given the intense human modification of estuarine-coastal systems
883 through changes in freshwater and sediment input, introductions of alien species and nutrient
884 enrichment, there is uncertainty that ecological responses to oscillating climate patterns can be
885 detected in observations having the large and varied signals of human disturbance shown above
886 [*Cloern and Jassby*, 2008]. Here we summarize an ecological regime shift that occurred in South
887 San Francisco Bay after a shift in atmospheric pressure patterns across the North Pacific Ocean.
888 Detection of this regime shift and its attribution to a climatic process was possible because of
889 observations sustained more than two decades before and a decade after the climate shift.

890

891 **8.2 Ecosystem regime shift**

892 The largest observed El Niño event occurred in 1997-98, and it was followed by an equally
893 strong La Niña in 1999. This abrupt transition appears to demarcate a climatic regime shift in the
894 North Pacific manifested as a change in atmospheric pressure and wind patterns, ocean
895 temperature and biological productivity. This shift was “possibly the most dramatic and rapid
896 episode of climate change in modern times” [*W T Peterson and Schwing, 2003*]. It was expressed
897 as sign changes in the two prominent modes of sea surface temperature and sea level pressure
898 across the North Pacific [*Chenillat et al., 2012*] -- the Pacific Decadal Oscillation (PDO), which
899 became strongly negative, and the North Pacific Gyre Oscillation (NPGO), which became
900 strongly positive in 1999 (Figure 16A,B). Regional responses to these ocean-basin scale indices
901 were measured in the coastal ocean adjacent to San Francisco Bay as cooling of surface waters
902 and increase in the upwelling index (Figure 16C,D). Intensification of upwelling and cooling at
903 this latitude are responses to strengthened equatorward winds and equatorward transport in the
904 California Current that are most strongly correlated with the NPGO. Because of this, the NPGO
905 is a primary indicator of upwelling, nutrient supply to phytoplankton, and primary production in
906 the California Current System (CCS) [*Di Lorenzo et al., 2008*]. Regional ocean models simulate
907 higher coastal nitrate concentration, chlorophyll *a* and zooplankton biomass in the central CCS,
908 and the differences result from both earlier and stronger upwelling in NPGO⁺ compared to
909 NPGO⁻ regimes [*Chenillat et al., 2012*]. Therefore, large-scale processes of ocean-atmosphere
910 coupling captured in Pacific climate indices like the NPGO have important ramifications for
911 biological productivity in the coastal waters adjacent to San Francisco Bay.

912 Unexpectedly, major changes in biological communities inside San Francisco Bay
913 followed the shift of the NE Pacific to its cool phase [*Cloern et al., 2007*]. We show examples as

914 abundance indices of demersal marine fish, crabs and shrimp species (Figure 16E-G) that migrate
915 into estuaries, either as adults to reproduce (shrimp) or as juveniles produced in the coastal ocean
916 (e.g., English sole, Dungeness crab). Populations in each of these communities reached record-
917 high levels during or soon after the 1998-99 climate shift and abundances have remained above
918 their 1980-2010 means since, except for 2005 and 2006 when the NPGO was weak. Synchronous
919 with these changes was an increase of phytoplankton biomass (chlorophyll *a*) in South Bay that
920 has exceeded the long-term mean each year since 1999. Bivalve mollusks disappeared from
921 shallow regions of South Bay in 1999 (Figure 16H) and bivalves have remained scarce in these
922 regions during the past decade [J. Thompson, USGS, personal communication 2012].

923 These observations reveal an ecological regime shift in San Francisco Bay that was
924 coherent with a climate shift in the N Pacific, suggesting a previously unrecognized linkage
925 between interdecadal variability of the ocean-atmosphere system and biological communities
926 inside this estuary. We compare the two regimes as box plots of NPGO, PDO, upwelling and
927 SST in the coastal ocean, and abundances/biomass of organisms in marine-influenced regions of
928 San Francisco Bay for the periods 1980-1998 (warm) and 1999-2010 (cool). The arrows (Figure
929 17) depict a cascade of responses from the ocean-basin scale (shift to NPGO⁺/PDO⁻), to the
930 regional scale (cooling, strengthening of upwelling), and to San Francisco Bay as population
931 changes of organisms across a range of trophic levels. The demersal fish, crabs and shrimp are
932 all predators that feed on bivalve mollusks, so the absence of bivalves since 1999 may be a result
933 of increased predation mortality, as observed in other estuaries [*Beukema and Dekker, 2005*].
934 The phytoplankton increase is presumably, then, a response to decreased bivalve grazing [*Cloern*
935 *et al., 2007*]. Therefore, South San Francisco Bay's biological communities were reorganized
936 through a trophic cascade that was initiated by an abrupt increase in abundance of predators

937 whose populations most closely track the NPGO [*Cloern et al.*, 2010]. These predators include
938 juvenile forms that migrate into San Francisco Bay, so their high abundances since 1999 must
939 reflect changes in their production rate in the coastal ocean. A likely mechanism is the enhanced
940 plankton food supply to early life stages of flatfish and crabs by the shift from a warm, low-
941 production to a cool, high-production regime of the NE Pacific that amplifies abundances of
942 zooplankton, pelagic fish and seabirds in the CCS [*W T Peterson and Schwing*, 2003]. The last
943 climate shift in the N Pacific occurred in 1976, before regular biological sampling began in
944 South San Francisco Bay, so we have no observations (or even proxies) to determine if the
945 current ecosystem regime existed in earlier cool regimes of the NE Pacific (e.g., 1948-1976).
946 Validation of the linkages hypothesized in Figure 17 will require sampling through the next
947 warm regime, so we tell students that a single career is not long enough.

948 Every measured component of phytoplankton dynamics changed in South Bay after the
949 climate shift. The earliest sign of change was a surprising bloom in October 1999 (Figure 18A),
950 the first occurrence of an autumn bloom and a departure from the canonical 1978-1998 pattern of
951 one (spring) bloom each year. Other autumn-winter blooms have occurred since, including the
952 first observed dinoflagellate red tide in September 2004 [*Cloern et al.*, 2005]. Calculated gross
953 primary production (GPP) has increased during the past decade and it exceeded the long-term
954 mean every year since 1997, except one (Figure 18B). As a simple index of the seasonal
955 development of phytoplankton biomass, we calculated the day each year when cumulative
956 chlorophyll-*a* reached the midpoint (“fulcrum”) of annual cumulative chlorophyll-*a*. This index
957 shifted +31 days (from mid April to mid May) between 1978-1998 and 1999-2010 (Figure 18C),
958 reflecting the new occurrences of autumn-winter blooms and overall increases in summer
959 biomass.

960 We illustrate phytoplankton community changes as occurrence timelines of two marine
961 diatoms (*Thalassiosira rotula*, *T. punctigera*) and two heterotrophic (non-photosynthetic)
962 dinoflagellates (*Oxytoxum milneri*, *Polykrikos schwartzii*). *Thalassiosira rotula* occurred
963 commonly and was the biomass-dominant species in San Francisco Bay during 1992-2001
964 [Cloern and Dufford, 2005]. It occurred less frequently in the past decade (Figure 18D) and its
965 contribution to biomass during 1999-2010 was ranked only 54th. The biomass-dominant after
966 1999 was *T. punctigera*, a species not observed previously. *Oxytoxum milneri* occurred
967 commonly before but was never observed after 1997. Conversely, *Polykrikos schwartzii* first
968 appeared in 1999 and has been observed regularly since (Figure 18D). The synchrony of these
969 species appearances and disappearances with the 1998-99 climate shift suggests that they are
970 related to phytoplankton species changes in the coastal ocean, perhaps analogous to the switch
971 from warm- to cold-water copepod species in the northern CCS after 1998 [W T Peterson and
972 Schwing, 2003]. However, our knowledge of the life cycles, biogeography and physiological
973 ecology of marine phytoplankton is not sufficient to explain why one *Thalassiosira* species
974 would nearly completely replace another or why one heterotrophic dinoflagellate would
975 completely replace another at about the same time. We also know surprisingly little about the
976 ecological significance of these kinds of species changes, although they could be substantial
977 because of differences in cell size, behavior and nutritional quality among phytoplankton species
978 [Cloern, 1996]. The diminished biomass of *Thalassiosira rotula*, for example, is intriguing
979 because this marine diatom produces oxylipins that arrest hatching of copepod eggs in laboratory
980 experiments [Carotenuto et al., 2011]. The environmental significance of these experiments is
981 debated, but they highlight the critical need for increased knowledge of biological interactions at

982 the species level to understand the significance of climate-driven community changes at the
983 ecosystem level.

984

985 **8.3 Significance of the changes**

986 The San Francisco Bay response exemplifies an emerging principle: variability in coastal oceans
987 can be a powerful driver of variability inside estuaries and bays, and shifts in large-scale ocean-
988 atmosphere dynamics can induce ecological regime shifts in estuaries and bays through their
989 connectivity to coastal oceans. Other examples are accumulating. Species diversity, abundance
990 and growth rates of juvenile marine fish in the Thames Estuary are significantly correlated with
991 the NAO index [*Attrill and Power, 2002*]. Recruitment of Atlantic croaker (*Micropogonias*
992 *undulatus*) along the US east coast is highest during the warm (+) phase of the NAO when
993 estuarine water temperature and overwinter survival of juveniles are high [*Hare and Able, 2007*].
994 Shellfish toxicity in Puget Sound is associated with the warm (+) phase of the PDO because
995 warming promotes growth of the toxin-producing dinoflagellate *Alexandrium catenella* [*Moore*
996 *et al., 2010*]. Perhaps the most striking climate-driven transformation of an estuarine ecosystem
997 has occurred in Narragansett Bay, where a 1.7 °C winter warming of coastal waters since 1970
998 has been accompanied by loss of the traditional winter phytoplankton bloom, 40-50% reduction
999 in primary production, reduced supply of organic matter to sediments, decreased benthic
1000 metabolism and abundance of demersal fish, and a switch of N cycling from net denitrification to
1001 net N fixation [*Nixon et al., 2009*]. Historical management of estuarine ecosystems has been
1002 based from a landward-looking perspective because of human disturbances to surrounding
1003 landscapes (Sections 3-7). However, water quality, system production, biological communities

1004 and biogeochemical cycling in estuaries also respond to oceanographic processes influenced by
1005 large-scale climate patterns.

1006 The oceanic drivers of estuarine variability add to the already complex challenge of
1007 managing water quality and living resources of nearshore coastal ecosystems because they can
1008 confound outcomes of actions to reduce effects of human disturbance, such as nutrient
1009 enrichment. In the Hood Canal of Puget Sound, where scientific investigations are focused on the
1010 genesis of bottom-water hypoxia, the coastal ocean is a significant source of nutrients providing,
1011 for example, more than 90% of the nitrogen input [*Steinberg et al.*, 2010]. Recent studies in the
1012 Columbia River Estuary documented multiple intrusions of deep, low-DO coastal water brought
1013 to the surface by wind-driven upwelling and transported into the estuary by tidal advection and
1014 estuarine circulation [*Roegner et al.*, 2011]. Tidal dispersion drives inputs of phytoplankton
1015 biomass into San Francisco Bay during the upwelling season [*Martin et al.*, 2007], and
1016 metabolism (and therefore oxygen consumption) in Tomales Bay is fueled by inputs of
1017 phytoplankton biomass produced in the adjacent upwelling system [*S V Smith et al.*, 1996].
1018 Harmful algal blooms in coastal waters provide inocula for blooms of harmful species to develop
1019 in estuaries [*Cloern et al.*, 2005]. Therefore, inputs from the ocean and especially from upwelling
1020 systems can have similar consequences to those of anthropogenic nutrient enrichment: high
1021 nutrients and phytoplankton biomass, low DO and harmful algal blooms.

1022

1023 **9. THE MONITORING IMPERATIVE**

1024 **9.1 Patterns of temporal change**

1025 The study of temporal change has always been an important part of ecology. Diel, tidal, annual
1026 and longer cycles in population behavior and abundance are the most obvious and a matter of

1027 study since the early days of ‘modern’ ecology [Elton, 1927]. Non-cyclic change is also a
1028 fundamental component of ecological understanding, such as the continuous species colonization
1029 and extinction within communities revealed by island biogeographical studies [MacArthur and
1030 Wilson, 1963] and the maturation of undisturbed whole ecosystems so well summarized by
1031 Odum [1969]. In the last few decades, the possibility of chaotic dynamics in populations
1032 [Hastings *et al.*, 1993] and biotic responses to decadal ocean-atmosphere regimes (Section 8)
1033 have been a focus of investigation. Accordingly, stationarity of ecological time series -- which
1034 implies constant mean and variance -- has never been a dominant idea in ecology, even for
1035 ecosystems not subject to anthropogenic impacts. Although a characteristic average long-term
1036 state has traditionally been a useful principle in related fields such as hydrology, even that
1037 assumption is now recognized as untenable in the face of strong climate change [Milly *et al.*,
1038 2008].

1039 A key feature of many temporal patterns observed in nature – including both
1040 environmental [Steele, 1985] and biotic [Pimm and Redfearn, 1988] time series -- is that the
1041 general appearance of the pattern is more or less unchanged when observed at different time
1042 scales. Although known for a long time, this scale invariance became a subject of study only in
1043 the 1970s when the needed mathematical tools became available [Gisiger, 2001]. Now we know
1044 that ecological variability can often be described by a power-law spectrum proportional to $(1/f)^{\nu}$,
1045 where f is frequency (cycles yr^{-1}) and $0 \leq \nu \leq 2$ [Cyr and Cyr, 2003; JM Halley, 1996]. $\nu = 0$
1046 corresponds to white noise, which emphasizes short time scales; $\nu = 2$ corresponds to a random
1047 walk or brown noise, which emphasizes long time scales; and $\nu = 1$ corresponds to $1/f$ or pink
1048 noise, which is not biased towards any particular time scale.

1049 Pink noise -- as opposed to the traditional white noise -- appears to be the most suitable

1050 null model for ecological time series [*J Halley and Inchausti, 2004*]. The prevalence of pink
1051 noise has implications for estuarine monitoring: What distinguishes pink and even ‘redder’ noise
1052 with $\nu > 1$ is that variance continues increasing, no matter how long the time series. Given that
1053 marine environmental variables such as temperature tend to have reddened spectra, and
1054 terrestrial whitened, at least for time periods up to 50-100 years [*Vasseur and Yodzis, 2004*], the
1055 variability spectrum for estuaries may therefore depend on the relative importance of oceanic and
1056 terrestrial drivers. In particular, estuaries with strong ocean influence, such as San Francisco
1057 Bay, may have redder spectra and be more influenced by relatively slow or rare environmental
1058 fluctuations with long periods. The data requirements to reliably distinguish $1/f$ -noise from
1059 alternative noise models and to estimate ν [*Fleming, 2008*] are greater than currently available
1060 for San Francisco Bay biotic and water quality variables. Nonetheless, we should anticipate
1061 continuous and unexpected long-term changes and trends in the estuary's drivers.

1062

1063 **9.2 Implications for monitoring**

1064 These patterns of temporal change imply several lessons for monitoring, which we summarize
1065 briefly in the context of San Francisco Bay and Delta. First, discrete monitoring programs must
1066 take into account variability on scales shorter than the sampling interval because of potential
1067 uncertainty and bias due, for example, to tidal and diel cycles [*Jassby et al., 2005; Lucas and*
1068 *Cloern, 2002; Lucas et al., 2002*]. Moreover, the assumption of white or gaussian noise may not
1069 be appropriate to account for this uncertainty. Monitoring programs must, at some point, include
1070 focused, higher-frequency studies to understand the effects of shorter time scales. Milly et al.
1071 [*2008*] also caution that the nature of variability on these shorter scales may itself be changing,
1072 requiring more attention and adjustments to our current assumptions and sampling designs.

1073 Second, pink noise implies the presence of slow processes that cannot be identified
1074 without a commitment to ongoing sampling. Of our case studies, the gradual recovery of water
1075 clarity over decades after a short period of hydraulic mining was perhaps the slowest process.
1076 This recovery was a ‘noisy’ one (Figure 9), which could not be identified with a data set much
1077 shorter than the 35 years available. One can assume that there are many slow trends underway
1078 which we cannot yet see. Inclusion of higher-frequency automated monitoring cannot substitute
1079 for sites that already have a long record, even if at a much lower frequency. Commitments are
1080 therefore essential to continue sampling into the indefinite future at a small number of sentinel
1081 sites that are the most representative of their subregions and have the longest monitoring records
1082 [*Burt et al.*, 2010; *Jassby*, 1998]. Uninterrupted and long data records are key to the most
1083 informative monitoring programs [*Southward*, 1995].

1084 Many estuaries do have monitoring programs that are sustained because of mandates. In
1085 the San Francisco Bay-Delta, for example, some long-term monitoring programs have persisted
1086 through budget shortfalls because they are mandated in water rights decisions and biological
1087 opinions about the long-term operations of the Central Valley and State Water projects.
1088 Mandated monitoring thus improves the chances for long-term survival. Unfortunately it can also
1089 be difficult to modify a mandated program even if its design could be made more informative or
1090 if better sampling and analysis methods become available. The cost of long-term, historically
1091 mandated programs may also make it difficult to establish new, potentially more useful
1092 programs. Moreover, data management, analysis, synthesis and communication have been much
1093 less sustainable than the sampling itself, especially during budget shortfalls [*Hughes and Peck*,
1094 2008]. Accordingly, monitoring programs themselves need to be ‘adaptively managed’ while
1095 recognizing two essential principles: a corresponding capacity for useful data synthesis must

1096 accompany changes in sampling; and at least a small number of sentinel sites must survive
1097 changes in the programs.

1098 Third, some processes may be so rare that we have no experience with them, but they
1099 may be large, have a disproportionate effect and need immediate attention. In the world of
1100 finance statistics, the term ‘black swan’ has been coined for these events, which have had such
1101 widespread and devastating consequences [*Taleb, 2007*]. Perhaps the pelagic organism decline
1102 described below could be considered such an event. But there is evidence that such catastrophic
1103 ecosystem events have statistical early warning indicators [*Carpenter et al., 2011; Scheffer et al.,*
1104 *2009*] that could be observed in routine monitoring. Indeed, the word *monitoring* derives from
1105 the Latin *monere*, to warn. ‘Black swans’ require both continuity of long-term monitoring and a
1106 high enough sampling frequency to calculate these early-warning indicators.

1107 Finally, some processes are the result of multiple drivers that cannot be sorted out with
1108 short data sets. For example, a nonlinear regression model with three coefficients (in addition to
1109 the intercept) was required to disentangle the decline of suspended particulate matter from
1110 interannual variability in flow (Equation 1, Figure 10). Simulation studies suggest that 10-20
1111 observations are required per regression coefficient [*Harrell, 2001*], which means that 30-60
1112 years are needed just for this simple model.

1113 Yet multiple drivers and more complex interactions are probably the rule. Perhaps the
1114 most prominent example from this estuary is the marked decrease during the past decades of
1115 some estuarine-dependent fish, some to the point of near-extinction. Particular interest in the past
1116 decade has been given to four pelagic fish species whose populations all declined significantly in
1117 the early 2000s: delta smelt, longfin smelt, striped bass (*Morone saxatilis*), and threadfin shad
1118 (*Dorosoma petenense*) [*Sommer et al., 2007*]. Population declines of multiple species across

1119 multiple trophic levels are symptoms of an intensely disturbed ecosystem, and the single most
1120 urgent questions posed to the scientific community are: What are the drivers of these population
1121 declines, and what actions can be taken to promote recovery and then sustain populations of fish
1122 endemic to this estuary? These questions have social and economic significance because “the
1123 solutions under consideration include major investments in infrastructure, changes in water
1124 management, and rehabilitation of species’ habitats that collectively will cost billions of dollars”
1125 [Thomson *et al.*, 2010].

1126 Often implicit in these questions is the expectation (or hope) that stressors on the estuary
1127 originate from a single or small set of drivers whose effects can be mitigated. However, the fish
1128 and plankton species of concern have distinct life histories and varying patterns of population
1129 collapse over time, suggesting that stressors arise from multiple, interacting drivers of change
1130 associated with human disturbances and climate anomalies such as sustained drought. Results
1131 from two independent analyses (multivariate autoregressive modeling and change point analysis)
1132 are consistent with this interpretation. They reveal significant associations between population
1133 declines of individual species during the period 1967-2007 and each of the drivers described in
1134 Sections 3-5, e.g., negative associations between: longfin smelt abundance and salinity (X2);
1135 striped bass abundance and water clarity; delta smelt abundance and water exports during winter;
1136 and density of *Corbula* and calanoid copepods [Mac Nally *et al.*, 2010; Thomson *et al.*, 2010].
1137 Our preceding remarks about the suspended particulate matter decline should make it clear that
1138 many decades of observation may be required to understand the relative importance and modes
1139 of operation of so many drivers.

1140

1141 **9.3 Monitoring and environmental policy**

1142 A consensus has emerged from these and other analyses [BDCP, 2010; L. R. Brown and Moyle,
1143 2005; NRC, 2010] that the challenge of sustaining communities of native species will require an
1144 integrated strategy to mitigate the cumulative effects of water consumption and diversions, river
1145 impoundments, introductions of non-native species, and other human disturbances such as land-
1146 use change [Cloern, 2007] and inputs of toxic pollutants [Brooks et al., 2011]. The challenge will
1147 grow as the effects of global warming drive further changes in freshwater inflow, sea level, water
1148 temperature, salinity and sediment inputs [Cloern et al., 2011]. A general lesson, relevant to
1149 other damaged ecosystems, emerges from decades of sustained and careful observation in San
1150 Francisco Bay: “a holistic approach to managing the ecology of imperiled fishes in the delta will
1151 be required if species declines are to be reversed” [NRC, 2010].

1152 In the past, sustained monitoring gave rise to the Clean Water Act and then proved its
1153 efficacy. More recently, monitoring of the *Corbula* invasion has spurred rules regarding
1154 discharge of ballast water into San Francisco Bay and other estuaries. The same sustained
1155 monitoring is now enabling us to anticipate emerging problems, exemplified by the changing
1156 role of nutrients in South San Francisco Bay, as we now describe. Marine species use estuaries as
1157 nursery habitats, and recruitment from estuaries can strongly drive marine population dynamics
1158 of many commercial fish species [Attrill and Power, 2002]. Observations in San Francisco Bay
1159 revealed that the immigration of marine fish and invertebrates can also drive biological
1160 community changes inside estuaries (Section 8). These community changes have altered the
1161 balance between phytoplankton production and consumption, leading to increases in
1162 phytoplankton biomass and primary production during the past decade. The trend of increasing
1163 primary production from 1978-2010 is ecologically significant because it spans the ranges
1164 defining oligotrophic (low-production, $< 100 \text{ g C m}^{-2} \text{ yr}^{-1}$), mesotrophic (moderate-production,

1165 100-300 g C m⁻² yr⁻¹), and eutrophic (high-production, > 300 g C m⁻² yr⁻¹) estuaries [*Nixon*,
1166 1995]. Thus, South San Francisco Bay shifted from an oligo-mesotrophic estuary to a meso-
1167 eutrophic estuary after 1997 (Figure 18B). This upward shift signals an increased efficiency in
1168 the conversion of nutrients into algal biomass and a weakening of the estuary's resistance to the
1169 harmful consequences of nutrient enrichment. Nutrient enrichment of San Francisco Bay was not
1170 a concern to water-quality managers in the past, but it is now and they ask: Is South Bay on a
1171 trajectory toward the impairments seen in Chesapeake Bay, what standards are appropriate for
1172 protecting the ecological integrity of this estuary, and will policies be required to mandate
1173 additional wastewater treatment processes to remove nutrients?

1174 Policies to reduce nutrient loadings to other estuarine-coastal systems have had mixed
1175 and sometimes disappointing results. These policies are usually established from an assumed
1176 functional relationship between responses, such as amplified algal biomass, and nutrient loading
1177 rate [*Carstensen et al.*, 2011]. Target responses are selected, and appropriate nutrient loading
1178 rates are then prescribed from the functional relationship. For some estuaries, such as Tampa
1179 Bay, this approach has been highly successful and steady progress has been made to reduce algal
1180 biomass and recover seagrasses [*Greening et al.*, 2011]. In other cases, costly programs to reduce
1181 nutrient inputs have had unexpected results as algal responses have been muted and shown lags
1182 and hysteresis, patterns interpreted as manifestations of “shifting baselines” [*Carstensen et al.*,
1183 2011]. Observations in San Francisco Bay illustrate a shifting baseline after biological
1184 communities were restructured through a climate regime shift (Figure 17). These links between
1185 climate regimes, biological communities and water quality are newly emerging themes of
1186 estuarine-coastal research (e.g., [*Cloern et al.*, 2007], [*Nixon et al.*, 2009]), so they have not yet
1187 been considered in most nutrient-management strategies.

1188

1189 **9.4 Concluding perspective**

1190 In this review we used data sets collected in a well-studied estuary to illustrate how the coupling
1191 of regular sampling with ongoing analyses and retrospective syntheses has become a powerful
1192 research approach for understanding ecosystem dynamics at time scales longer than the duration
1193 of individual grants. This approach has been used broadly to reveal that the long-term behavior
1194 of estuarine habitats, biological communities and biogeochemical processes is nonstationary and
1195 includes abrupt shifts driven by local-scale processes such as species introductions (e.g., Figure
1196 11) and global-scale processes such as climate shifts (e.g., Figure 17). Empirical observations
1197 over the past half-century have documented fast, large-amplitude changes in the world's
1198 estuarine-coastal ecosystems that depart radically from Eugene P. Odum's depiction of the
1199 natural evolution of biological communities and their habitats in unperturbed environments.
1200 These changes reflect today's human domination of the Earth's ecosystems as "most aspects of the
1201 structure and functioning of Earth's ecosystems cannot be understood without accounting for the
1202 strong, often dominant influence of humanity" [Vitousek *et al.*, 1997].

1203 Monitoring is essential for managing the human dimension of ecosystem dynamics
1204 because it detects environmental changes, provides insights into their underlying causes, can
1205 provide early warning signs of impending state shifts [Carpenter *et al.*, 2011; Scheffer *et al.*,
1206 2009], prompts mitigation and adaptation policies (such as the Clean Water Act), and it measures
1207 outcomes of those policies. Yet, commitments to monitoring programs are difficult to secure and
1208 sustain because their value accrues over the long term and precise benefits cannot be prescribed
1209 in advance. The stakes are growing, however, as the cumulative effects of fast-paced change
1210 across all global ecosystems might be driving a trajectory toward a planetary state shift with
1211 large social and economic consequences [Barnosky *et al.*, 2012]. Given the scope and

1212 breathtaking pace of change occurring in the world's estuarine-coastal ecosystems (e.g., Figure
1213 1), the imperative for monitoring data and their analysis has never been greater.

1214

1215 **ACKNOWLEDGMENTS**

1216 The following people generously shared data used in this review: Heather Fuller, Karen Gehrts
1217 and Tariq Kadir (California Department of Water Resources); Kathy Hieb and April Hennessy
1218 (California Department of Fish and Game); James Ervin, Neal Van Keuren, Bob Wandro and
1219 Alo Kauravlla (City of San Jose, Environmental Services); Ken Torke (City of Palo Alto);
1220 Michelle Hornberger (USGS); Tim Hollibaugh (University of Georgia); Norbert Jaworski
1221 (USEPA, retired); David Wilcox (Virginia Institute of Marine Science); Judith Dobson (Scottish
1222 Environment Protection Agency); Teresa Catry (CESAM/Museu Nacional de História Natural,
1223 Portugal); Charles Gallegos (Smithsonian Environmental Research Center); Jens Würigler
1224 Hansen (Aarhus University, Denmark); Shilun Yang (East China Normal University); Wim
1225 Kimmerer (San Francisco State University); Hans Ulrik Riisgard (University of Southern
1226 Denmark). And we thank our colleagues Lisa Lucas, Noah Knowles, Anke Mueller-Solger, Dave
1227 Schoellhamer, Mike Brett and Erica Goldman for their helpful suggestions to improve this paper.
1228 Funding for the long-term USGS studies of San Francisco Bay biological communities and water
1229 quality has been provided by the USGS Toxic Substances Hydrology Program, USGS Priority
1230 Ecosystem Science, National Research Program of the USGS Water Resources Discipline, and
1231 by the San Francisco Estuary Institute -- Regional Monitoring Program. We are deeply
1232 appreciative of the characteristically insightful comments and supportive words from the late
1233 Scott Nixon.

1234

1235

1236 **APPENDIX**

1237 **Data sources**

1238 The water quality data used for analysis here were obtained primarily by the Interagency
1239 Ecological Program's (IEP) Environmental Monitoring Program (EMP) and the USGS Water
1240 Quality of San Francisco Bay Program. The IEP is a consortium of 10 member agencies
1241 cooperating on research in the Bay and Delta since 1970, and its EMP activities of primary
1242 interest here -- discrete water quality, zooplankton and fish monitoring -- are operated by the
1243 California Departments of Water Resources (DWR) and of Fish and Game (DFG). The EMP
1244 samples water quality and zooplankton mostly in the Delta and Suisun Bay, while the USGS
1245 program samples water quality mostly in San Francisco Bay (including Suisun Bay). The
1246 California DFG San Francisco Bay Study of fish and shellfish includes all of the Bay and much
1247 of the Delta. Many other types and sources of data are also used, all of which are summarized in
1248 Table A1.

1249

1250 **Data analysis**

1251 Except in the cases where individual observations were called for, we binned water quality data
1252 for each variable and station by month using the mean to form a collection of monthly time
1253 series. For zooplankton and *Corbula amurensis*, we imputed missing monthly data using the
1254 long-term mean for the month. Annual averages for zooplankton used March-November data
1255 only. The 1987 *Corbula* average is based on May-December values only.

1256 Fish counts for all five indicated species for all marine subembayments (South, Central,
1257 San Pablo bays) were summed for each tow, converted to CPUE = 1000 * count/tow-area, and

1258 then averaged over each year (February-October). A similar procedure was used for three crab
1259 and two shrimp species, except that CPUE = count/5-min-tow.

1260 When estimating trends in time series, we used a robust measure sometimes known as the
1261 Theil-Sen slope, unless otherwise noted. This is simply the median slope of the lines joining all
1262 pairs of points in the series. We assessed the significance of these trends using the Mann-Kendall
1263 test. Tests were conducted only if at least 50% of the total possible number of values in the
1264 beginning and ending fifths of the record were present [*Helsel and Hirsch, 2002*].

1265 In the graphs, boxplots are traditional boxplots, i.e., the line within the box represents the
1266 median, the boxes extend from the first through third quartiles, and the vertical lines extend to all
1267 points within 1.5 times the interquartile distance (box height). Smoothing lines in graphs are
1268 local polynomial regressions, in particular, loess smooths with span = 0.75 and degree = 2
1269 [*Helsel and Hirsch, 2002*].

1270 Restricted cubic, or natural, splines were used as transforms for predictors in regression
1271 relationships. To minimize the number of parameter estimates, a restricted cubic spline with only
1272 three knots was used, requiring only two parameters. The knot positions -- at the 0.1, 0.5, and 0.9
1273 quantiles -- were chosen based on general recommendations from simulation studies, and were
1274 not tailored in any way for these particular data sets. We conducted analyses of variance to
1275 determine if the nonlinear part of each predictor transform actually improved the regression
1276 model or if it could be replaced by a simple linear term [*Harrell, 2001*].

1277 Multivariate regression model results are illustrated as partial residual plots, which show
1278 the relationship between a given independent variable and the response variable, while
1279 accounting for the other independent variables in the model. Partial residuals for any predictor
1280 are formed by omitting that term from the model [*Chambers, 1992*].

1281 We used the **R** language [*RDevelopmentCoreTeam*, 2012] for all calculations and graphs,
1282 including extensive use of the **ggplot2** [*Wickham*, 2009] and **wq** [*Jassby and Cloern*, 2012]
1283 packages.

References

- Alpine, A. E., and J. E. Cloern (1992), Trophic interactions and direct physical effects control phytoplankton biomass and production in an estuary, *Limnol Oceanogr*, 37(5), 946-955.
- Arthur, J. F., M. D. Ball, and S. Y. Baughman (1996), Summary of federal and state water project environmental impacts in the San Francisco Bay-Delta estuary, California, in *San Francisco Bay; the Ecosystem*, edited by J. T. Hollibaugh, pp. 445-495, Pacific Division, American Association for the Advancement of Science.
- Attrill, M. J. (Ed.) (1998), *A Rehabilitated Estuarine Ecosystem. The Environment and Ecology of the Thames Estuary*, Kluwer Academic Publishers, Dordrecht.
- Attrill, M. J., and M. Power (2002), Climatic influence on a marine fish assemblage, *Nature*, 417(6886), 275-278.
- Baltensweiler, W. (1964), *Zeiraphera griseana* Hubner (Lepidoptera: Tortricidae) in the European Alps. A contribution to the problem of cycles, *Canad. Entomol.*, 96, 792-800.
- Barnosky, A. D., et al. (2012), Approaching a state shift in Earth's biosphere, *Nature*, 486(7401), 52-58.
- Baron, J. S., N. L. Poff, P. L. Angermeier, C. N. Dahm, P. H. Gleick, N. G. J. Hairston, R. B. Jackson, C. A. Hohnston, B. D. Richter, and A. D. Steinman (2002), Meeting the ecological and societal needs for freshwater, *Ecol Appl*, 12(5), 1247-1260.
- BDCP (2010), Highlights of the BDCP (Bay Delta Conservation Plan). Available at: http://bdcweb.com/Files/Highlights_of_the_BDCP_FINAL_03-17-11.pdf
- Beck, H. J., and G. F. Birch (2012), Metals, nutrients and total suspended solids discharged during different flow conditions in highly urbanised catchments, *Environmental Monitoring and Assessment*, 184(2), 637-653.
- Bennett, W. A. (2005), Critical assessment of the delta smelt population in the San Francisco Estuary, California, *San Francisco Estuary and Watershed Science*, 3(2), 1-73. Available at: <http://escholarship.org/uc/item/0725n0725vk>
- Beukema, J. J., and R. Dekker (2005), Decline of recruitment success in cockles and other bivalves in the Wadden Sea: possible role of climate change, predation on postlarvae and fisheries, *Mar Ecol-Prog Ser*, 287, 149-167.
- Blum, M. D., and H. H. Roberts (2009), Drowning of the Mississippi Delta due to insufficient sediment supply and global sea-level rise, *Nature Geoscience*, 12, 488-491.
- Borja, Á., I. Muxika, and J. Franco (2006), Long-term recovery of soft-bottom benthos following urban and industrial sewage treatment in the Nervión estuary (southern Bay of Biscay), *Marine Ecology Progress Series*, 313, 43-55.

Bricker, S., B. Longstaff, W. Dennison, A. Jones, K. Boicourt, C. Wicks, and J. Woerner (2007), Effects of Nutrient Enrichment In the Nation's Estuaries: A Decade of Change. NOAA Coastal Ocean Program Decision Analysis Series No. 26. , 328 pp.

Brooks, M. L., et al. (2011), Life histories, salinity zones, and sublethal contributions of contaminants to pelagic fish declines illustrated with a case study of San Francisco Estuary, California, USA, *Estuar Coast*, 35(2), 603-621.

Brosnan, T. M., and M. L. O'Shea (1996), Sewage abatement and coliform bacteria trends in the lower Hudson-Raritan Estuary since passage of the Clean Water Act, *Water Environment Research*, 68, 25-35.

Brown, L. R., and P. B. Moyle (2005), Native fishes of the Sacramento–San Joaquin drainage, California: A history of decline, *American Fisheries Society Symposium* 45, 75–98.

Brown, L. R., and D. Michniuk (2007), Littoral fish assemblages of the alien-dominated Sacramento–San Joaquin Delta, California, 1980–1983 and 2001–2003, *Estuaries and Coasts*, 30, 186-200.

Brown, T. C., and P. Froemke (2012), Nationwide assessment of nonpoint source threats to water quality, *BioScience*, 62(2), 136-146.

Burt, T. P., N. J. K. Howden, F. Worrall, and M. J. Whelan (2010), Long-term monitoring of river water nitrate: how much data do we need?, *Journal of Environmental Monitoring*, 12(1), 79.

CADWR (2007), California Central Valley unimpaired flow data, 4th edition. Bay-Delta Office, California Dept. of Water Resources, Sacramento.

CALFED (2012), History of CALFED Bay-Delta Program <http://calwater.ca.gov/calfed/about/History/Detailed.html>, Accessed: 13 June 2012., (13 June 2012).

Carlton, J. T., and J. B. Geller (1993), The global transport of nonindigenous marine organisms, *Science*, 261(5117), 78-82.

Carlton, J. T., J. K. Thompson, L. E. Schemel, and F. H. Nichols (1990), Remarkable invasion of San Francisco Bay (California, USA) by the Asian clam *Potamocorbula amurensis*. I. Introduction and dispersal, *Marine Ecology Progress Series*, 66, 81-94.

Carnegie, R. B., and E. M. Burreson (2011), Declining impact of an introduced pathogen: *Haplosporidium nelsoni* in the oyster *Crassostrea virginica* in Chesapeake Bay, *Mar Ecol-Prog Ser*, 432, 1-15.

Carotenuto, Y., A. Ianora, and A. Miralto (2011), Maternal and neonate diatom diets impair development and sex differentiation in the copepod *Temora stylifera*, *Journal of Experimental Marine Biology and Ecology*, 396(2), 99-107.

Carpenter, S. R., et al. (2011), Early warnings of regime shifts: a whole-ecosystem experiment, *Science*,

332(6033), 1079-1082.

Carstensen, J., M. Sanchez-Camacho, C. M. Duarte, D. Krause-Jensen, and N. Marba (2011), Connecting the dots: responses of coastal ecosystems to changing nutrient concentrations, *Environ Sci Technol*, 45(21), 9122-9132.

Catry, T., et al. (2011), Long-term declines of wader populations at the Tagus estuary, Portugal: a response to global or local factors?, *Bird Conservation International*, 21(04), 438-453.

Chambers, J. M. (1992), Linear models, in *Statistical Models in S*, edited by J. M. Chambers and T. J. Hastie, Wadsworth & Brooks/Cole.

Chauvand, L., F. Jean, O. Ragueneau, and G. Thouzeau (2000), Long-term variation of the Bay of Brest ecosystem: benthic-pelagic coupling revisited, *Mar Ecol-Prog Ser*, 200, 35-48.

Chauvaud, L., J. K. Thompson, J. E. Cloern, and G. Thouzeau (2003), Clams as CO₂ generators: The *Potamocorbula amurensis* example in San Francisco Bay, *Limnol Oceanogr*, 48(6), 2086-2092.

Chenillat, F., P. Rivière, X. Capet, E. Di Lorenzo, and B. Blanke (2012), North Pacific Gyre Oscillation modulates seasonal timing and ecosystem functioning in the California Current upwelling system, *Geophys Res Lett*, 39(1).

Clavero, M., and E. García-Berthou (2005), Invasive species are a leading cause of animal extinctions, *Trends Ecol Evol*, 20, 100.

Cloern, J. E. (1982), Does the benthos control phytoplankton biomass in South San Francisco Bay?, *Marine Ecology Progress Series*, 9, 191-202.

Cloern, J. E. (1987), Turbidity as a control on phytoplankton biomass and productivity in estuaries, *Cont Shelf Res*, 7(11-12), 1367-1381.

Cloern, J. E. (1996), Phytoplankton bloom dynamics in coastal ecosystems: A review with some general lessons from sustained investigation of San Francisco Bay, California, *Rev Geophys*, 34(2), 127-168.

Cloern, J. E. (1999), The relative importance of light and nutrient limitation of phytoplankton growth: a simple index of coastal ecosystem sensitivity to nutrient enrichment, *Aquatic Ecology*, 33, 3-16.

Cloern, J. E. (2001), Our evolving conceptual model of the coastal eutrophication problem, *Mar Ecol-Prog Ser*, 210, 223-253.

Cloern, J. E. (2007), Habitat connectivity and ecosystem productivity: Implications from a simple model, *Am Nat*, 169(1), E21-E33.

Cloern, J. E., and R. Dufford (2005), Phytoplankton community ecology: principles applied in San Francisco Bay, *Mar Ecol-Prog Ser*, 285, 11-28.

- Cloern, J. E., and A. D. Jassby (2008), Complex seasonal patterns of primary producers at the land-sea interface, *Ecol Lett*, 11(12), 1294-1303.
- Cloern, J. E., B. E. Cole, R. L. J. Wong, and A. E. Alpine (1985), Temporal dynamics of estuarine phytoplankton - a case study of San Francisco Bay, *Hydrobiologia*, 129(Oct), 153-176.
- Cloern, J. E., A. D. Jassby, J. K. Thompson, and K. A. Hieb (2007), A cold phase of the East Pacific triggers new phytoplankton blooms in San Francisco Bay, *P Natl Acad Sci USA*, 104(47), 18561-18565.
- Cloern, J. E., A. E. Alpine, B. E. Cole, R. L. J. Wong, J. F. Arthur, and M. D. Ball (1983), River discharge controls phytoplankton dynamics in the northern San Francisco Bay estuary, *Estuarine Coastal and Shelf Science*, 16, 415-429.
- Cloern, J. E., T. S. Schraga, C. B. Lopez, N. Knowles, R. G. Labiosa, and R. Dugdale (2005), Climate anomalies generate an exceptional dinoflagellate bloom in San Francisco Bay, *Geophys Res Lett*, 32(14).
- Cloern, J. E., et al. (2011), Projected evolution of California's San Francisco Bay-Delta-River System in a century of climate change, *PLoS One*, 6(9): e24465.
- Cloern, J. E., et al. (2010), Biological communities in San Francisco Bay track large-scale climate forcing over the North Pacific, *Geophys Res Lett*, 37, L21602.
- Cohen, A. N., and J. T. Carlton (1998), Accelerating invasion rate in a highly invaded estuary, *Science*, 279, 555-558.
- Cohn, T. A., D. L. Caulder, E. J. Gilroy, L. D. Zynjuk, and R. M. Summers (1992), The validity of a simple statistical model for estimating fluvial constituent loads: An empirical study involving nutrient loads entering Chesapeake Bay, *Water Resources Research*, 28(9), 2353-2363.
- Cole, B. E., and J. E. Cloern (1984), Significance of biomass and light availability to phytoplankton productivity in San Francisco Bay, *Marine Ecology Progress Series*, 17, 15-24.
- Cole, B. E., J. K. Thompson, and J. E. Cloern (1992), Measurement of filtration rates by infaunal bivalves in a recirculating flume, *Mar Biol*, 113(2), 219-225.
- Conley, D. J., S. Markager, J. Andersen, T. Ellermann, and L. M. Svendsen (2002), Coastal eutrophication and the Danish National Aquatic Monitoring and Assessment Program, *Estuaries*, 4b, 848-861.
- Conomos, T. J. (1979), *San Francisco Bay: The Urbanized Estuary*, Pacific Division American Association for the Advancement of Science, San Francisco.
- Cyr, H., and I. Cyr (2003), Temporal scaling of temperature variability from land to oceans, *Evolutionary Ecology Research*, 5(8), 1183-1197.

Daskalov, G. M., and E. V. Mamedov (2007), Integrated fisheries assessment and possible causes for the collapse of anchovy kilka in the Caspian Sea, *ICES Journal of Marine Science*, 64(3), 503-511.

Di Lorenzo, E., et al. (2008), North Pacific Gyre Oscillation links ocean climate and ecosystem change, *Geophys Res Lett*, 35(8).

Diaz, R. J., and R. Rosenberg (2008), Spreading dead zones and consequences for marine ecosystems, *Science*, 321(5891), 926-929.

Dobson, J. (2000), Long term trends in trace metals in biota in the Forth Estuary, Scotland, 1981-1999, *Marine Pollution Bulletin*, 40(12), 1214-1220.

Duffy, T. A., A. E. McElroy, and D. O. Conover (2009), Variable susceptibility and response to estrogenic chemicals in *Menidia menidia*, *Marine Ecology Progress Series*, 380, 245-254.

Dynesius, M., and C. Nilsson (1994), Fragmentation and flow regulation of river systems in the northern third the world, *Science*, 266(5186), 753-762.

Elton, C. S. (1927), *Animal Ecology*, Methuen pp., Sidgwick and Jackson, London.

Enright, C., and S. D. Culbertson (2010), Salinity trends, variability, and control in the northern reach of the San Francisco Estuary, *San Francisco Estuary and Watershed Science*, 7(2).

Fan, H., and H. Huang (2008), Response of coastal marine eco-environment to river fluxes into the sea: a case study of the Huanghe (Yellow) River mouth and adjacent waters, *Mar Environ Res*, 65(5), 378-387.

Feyrer, F., B. Herbold, S. A. Matern, and P. B. Moyle (2003), Dietary shifts in a stressed fish assemblage: Consequences of a bivalve invasion in the San Francisco Estuary, *Environmental Biology of Fishes* 67, 277-288.

Feyrer, F., K. Newman, M. Nobriga, and T. Sommer (2011), Modeling the effects of future outflow on the abiotic habitat of an imperiled estuarine fish, *Estuar Coast*, 34(1), 120-128.

Fleming, S. W. (2008), Approximate record length constraints for experimental identification of dynamical fractals, *Annalen der Physik*, 17(12), 955-969.

Gallegos, C. L., P. J. Werdell, and C. R. McClain (2011), Long-term changes in light scattering in Chesapeake Bay inferred from Secchi depth, light attenuation, and remote sensing measurements, *Journal of Geophysical Research*, 116.

Gilbert, G. K. (1917), Hydraulic mining debris in the Sierra Nevada. US Geological Survey Professional Paper 105.

Gisiger, T. (2001), Scale invariance in biology: coincidence or footprint of a universal mechanism?, *Biological Reviews*, 76, 161-209.

Gleick, P. H., and M. Palaniappan (2010), Peak water limits to freshwater withdrawal and use, *P Natl Acad Sci USA*, 107(25), 11155-11162.

Gowan, R. J., P. Tett, and K. J. Jones (1992), Predicting marine eutrophication: the yield of chlorophyll from nitrogen in Scottish coastal waters, *Marine Ecology Progress Series*, 85, 153-161.

Greene, V. E., L. J. Sullivan, J. K. Thompson, and W. J. Kimmerer (2011), Grazing impact of the invasive clam *Corbula amurensis* on the microplankton assemblage of the northern San Francisco Estuary, *Marine Ecology Progress Series*, 431, 183-193.

Greening, H. S., and A. Janicki (2006), Toward reversal of eutrophic conditions in a subtropical estuary: water quality and seagrass response to nitrogen loading reductions in Tampa Bay, Florida, USA, *Environmental Management*, 38(2), 163-178.

Greening, H. S., L. M. Cross, and E. T. Sherwood (2011), A multiscale approach to seagrass recovery in Tampa Bay, Florida, *Ecological Restoration*, 29, 82-93.

Grenier, J. L., and J. A. Davis (2010), Water quality in South San Francisco Bay, California: current condition and potential issues for the South Bay Salt Pond Restoration Project, in *Reviews of Environmental Contamination and Toxicology*, edited by D. M. Whitacre, pp. 115-147, Springer, New York.

Grimaldo, L., T. Sommer, N. Van Ark, G. Jones, E. Holland, P. Moyle, B. Herbold, and P. Smith (2009), Factors affecting fish entrainment into massive water diversions in a tidal freshwater estuary: Can fish losses be managed?, *North American Journal of Fisheries Management*, 29(5), 1253-1270.

Grosholz, E. D. (2005), Recent biological invasion may hasten invasional meltdown by accelerating historical introductions, *P Natl Acad Sci USA*, 102(4), 1088-1091.

Halley, J., and P. Inchausti (2004), The increasing importance of 1/f-noises as models of ecological variability, *Fluctuation and Noise Letters*, 4(2), R1-R26.

Halley, J. M. (1996), Ecology, evolution and 1/f-noise, *Trends Ecol Evol*, 11(1), 33-37.

Hamilton, D. P., and J. V. Turner (2001), Integrating research and management for an urban estuarine system: the Swan-Canning Estuary, Western Australia, *Hydrological Processes*, 15(13), 2383-2385.

Hamza, W. (2009), Nile: Origin, Environments, Limnology and Human Use, in *Monographiae Biologicae*, edited by H. J. Dumont, pp. 75-94, Springer.

Hare, J. A., and K. W. Able (2007), Mechanistic links between climate and fisheries along the east coast of the United States: explaining population outbursts of Atlantic croaker (*Micropogonias undulatus*), *Fish Oceanogr.*, 16(1), 31-45.

Harrell, F. E. (2001), *Regression modeling strategies: with applications to linear models, logistic*

regression, and survival analysis, Springer Verlag.

Hastings, A., C. L. Hom, S. Ellner, P. Turchin, and H. C. J. Godfray (1993), Chaos in ecology: is mother nature a strange attractor?, *Annu. Rev. Ecol. Syst.*, 24, 1-33.

Hawkins, S. J., et al. (2002), Recovery of polluted ecosystems: the case for long-term studies, *Mar Environ Res*, 54(3-5), 215-222.

Helsel, D. R., and R. M. Hirsch (2002), Statistical methods in water resources, in *Techniques of Water-Resources Investigations of the United States Geological Survey. Book 4, Hydrologic Analysis and Interpretation*, edited, pp. 1-510, U.S. Geological Survey.

Herbert, D. A., W. B. Perry, B. J. Cosby, and J. W. Fourqurean (2011), Projected reorganization of Florida Bay seagrass communities in response to the increased freshwater inflow of Everglades restoration, *Estuar Coast*, 34(5), 973-992.

Hering, D., et al. (2010), The European Water Framework Directive at the age of 10: a critical review of the achievements with recommendations for the future, *Sci Total Environ*, 408(19), 4007-4019.

Higgins, S. N., and M. J. Vander Zanden (2010), What a difference a species makes: a meta-analysis of dreissenid mussel impacts on freshwater ecosystems, *Ecological Monographs*, 80(2), 179-196.

Hornberger, M. I., S. N. Luoma, D. J. Cain, F. Parchaso, C. L. Brown, R. M. Bouse, C. Wellise, and J. K. Thompson (2000), Linkage of bioaccumulation and biological effects to changes in pollutant loads in south San Francisco Bay, *Environ Sci Technol*, 34(12), 2401-2409.

Howarth, R. W., A. Sharpley, and D. Walker (2002), Sources of nutrient pollution to coastal waters in the United States: Implications for achieving coastal water quality goals, *Estuaries*, 25, 656-676.

Hughes, R. M., and D. V. Peck (2008), Acquiring data for large aquatic resource surveys: the art of compromise among science, logistics, and reality, *Journal of the North American Benthological Society*, 27, 837-859.

Jackson, R. B., S. R. Carpenter, C. N. Dahm, D. M. McNight, R. J. Naiman, S. L. Postel, and S. W. Running (2001), Water in a changing world, *Ecol Appl*, 11(4), 1027-1045.

Jackson, S. T. (2007), Looking forward from the past: history, ecology, and conservation, *Frontiers in Ecology and the Environment*, 5, 455.

Jaffe, B. E., R. E. Smith, and A. C. Foxgrover (2007), Anthropogenic influence on sedimentation and intertidal mudflat change in San Pablo Bay, California: 1856-1983, *Estuar Coast Shelf S*, 73, 175-187.

Jassby, A. D. (1998), Interannual variability at three inland water sites: Implications for sentinel ecosystems, *Ecol Appl*, 8(2), 287.

Jassby, A. D. (2008), Phytoplankton in the upper San Francisco Estuary: Recent biomass trends, their causes and their trophic significance, *San Francisco Estuary and Watershed Science*, 6(1), Article 2.

Jassby, A. D., and J. E. Cloern (2000), Organic matter sources and rehabilitation of the Sacramento-San Joaquin Delta (California, USA), *Aquat Conserv*, 10(5), 323-352.

Jassby, A. D., and J. E. Cloern (2012), wq: Exploring water quality monitoring data, R package version 0.3-6. Available at: <http://cran.r-project.org/web/packages/wq/index.html>.

Jassby, A. D., J. E. Cloern, and T. M. Powell (1993), Organic-carbon sources and sinks in San-Francisco Bay - variability induced by river flow, *Mar Ecol-Prog Ser*, 95(1-2), 39-54.

Jassby, A. D., J. E. Cloern, and B. E. Cole (2002), Annual primary production: Patterns and mechanisms of change in a nutrient-rich tidal ecosystem, *Limnol Oceanogr*, 47(3), 698-712.

Jassby, A. D., A. B. Mueller-Solger, and M. Vayssières (2005), Short-term variability of chlorophyll and implications for sampling frequency in the San Joaquin River, *IEP [Interagency Ecological Program for the San Francisco Estuary] Newsletter*, 18(1), 28.

Jassby, A. D., W. J. Kimmerer, S. G. Monismith, C. Armor, J. E. Cloern, T. M. Powell, J. R. Schubel, and T. J. Vendlinski (1995), Isohaline position as a habitat indicator for estuarine populations, *Ecol Appl*, 5(1), 272-289.

Jaworski, N. A. (1990), Retrospective study of the water quality issues of the upper Potomac Estuary, *Reviews in Aquatic Sciences*, 3(1), 11-40.

Jaworski, N. A., B. Romano, and C. Buchanan (2007), The Potomac River Basin and its Estuary: Landscape Loadings and Water Quality Trends 1895-2005. Available at: http://www.potomacriver.org/cms/wildlifedocs/NAJ_09_Chap9.pdf.

Jones, P. D. (2006), Water quality and fisheries in the Mersey estuary, England: A historical perspective, *Marine Pollution Bulletin*, 53(1), 144-154.

Kemp, W. M., et al. (2005), Eutrophication of Chesapeake Bay: historical trends and ecological interactions, *Mar Ecol-Prog Ser*, 303, 1-29.

Kimmerer, W. (2004), Open water processes of the San Francisco Estuary: From physical forcing to biological responses, *San Francisco Estuary and Watershed Science*, 2(1). Available at: <http://escholarship.org/uc/item/9bp499mv>

Kimmerer, W. (2005), Long-term changes in apparent uptake of silica in the San Francisco estuary, *Limnol Oceanogr*, 50(3), 793-798.

Kimmerer, W., E. Gartside, and J. J. Orsi (1994), Predation by an introduced clam as the likely cause of substantial declines in zooplankton of San Francisco Bay, *Marine Ecology Progress Series*, 113, 81-94.

- Kimmerer, W. J. (2006), Response of anchovies dampens effects of the invasive bivalve *Corbula amurensis* on the San Francisco Estuary foodweb, *Marine Ecology Progress Series*, 324, 207-218.
- Kimmerer, W. J., N. Ferm, M. H. Nicolini, and C. Penalva (2005), Chronic food limitation of egg production in populations of copepods of the genus *Acartia* in the San Francisco Estuary, *Estuar Coast*, 28, 541-550.
- Kjerfve, B., C. H. A. Ribeiro, G. T. M. Dias, A. M. Filippo, and V. D. S. Quaresma (1997), Oceanographic characteristics of an impacted coastal bay: Baía de Guanahara, Rio de Janeiro, Brazil, *Cont Shelf Res*, 17(13), 1609-1643.
- Knowles, N. (2002), Natural and management influences on freshwater inflows and salinity in the San Francisco Estuary at monthly to interannual scales, *Water Resources Research*, 38(12), 1289, doi:1210.1029/2001WR000360.
- Lehodey, P., et al. (2006), Climate variability, fish, and fisheries, *Journal of Climate*, 19(20), 5009-5030.
- Li, P., S. L. Yang, J. D. Milliman, K. H. Xu, W. H. Qin, C. S. Wu, Y. P. Chen, and B. W. Shi (2012), Spatial, temporal, and human-induced variations in suspended sediment concentration in the surface waters of the Yangtze Estuary and adjacent coastal areas, *Estuaries and Coasts*, DOI 10.1007/s12237-011-9382-x.
- Lucas, L. V., and J. E. Cloern (2002), Effects of tidal shallowing and deepening on phytoplankton production dynamics: A modeling study, *Estuaries*, 25(4A), 497-507.
- Lucas, L. V., T. Schraga, C. B. Lopez, J. R. Burau, and A. D. Jassby (2002), Pulsey, patchy water quality in the Delta: implications for meaningful monitoring, *IEP [Interagency Ecological Program for the Sacramento-San Joaquin Estuary] Newsletter*, 15(3), 27.
- Luoma, S. N., and J. E. Cloern (1982), The impact of waste-water discharge on biological communities in San Francisco Bay, in *San Francisco Bay: Use and Protection*, edited by W. J. Kockelman, T. J. Conomos and A. E. Leviton, pp. 137-160, Pacific Division American Association for the Advancement of Science, San Francisco.
- Luoma, S. N., and P. S. Rainbow (2008), *Metal Contamination in Aquatic Environments: Science and Lateral Management*, Cambridge University Press, New York.
- Mac Nally, R., et al. (2010), Analysis of pelagic species decline in the upper San Francisco Estuary using multivariate autoregressive modeling (MAR), *Ecol Appl*, 20 (5), 1417-1430.
- MacArthur, R. H., and E. O. Wilson (1963), An equilibrium theory of insular zoogeography, *Evolution*, 17(4), 373-387.
- Mantua, N. J., S. R. Hare, Y. Zhang, J. M. Wallace, and R. C. Francis (2002), The Pacific Decadal Oscillation, *J. Oceanogr.*, 58, 35-44.

Martin, M. A., J. P. Fram, and M. T. Stacey (2007), Seasonal chlorophyll a fluxes between the coastal Pacific Ocean and San Francisco Bay, *Marine Ecology Progress Series*, 357, 51-61.

Matthiessen, P., and R. J. Law (2002), Contaminants and their effects on estuarine and coastal organisms in the United Kingdom in the late twentieth century, *Environmental Pollution*, 120(3), 739-757.

May, C. L., J. R. Koseff, L. V. Lucas, J. E. Cloern, and D. H. Schoellhamer (2003), Effects of spatial and temporal variability of turbidity on phytoplankton blooms, *Mar Ecol-Prog Ser*, 254, 111-128.

McKee, L. J., and D. C. Gluchowski (2011), Improved nutrient load estimates for wastewater, stormwater and atmospheric deposition to South San Francisco Bay (South of the Bay Bridge). A Watershed Program report prepared for the Bay Area Clean Water Agencies (BACWA). Available at: http://bayareanutrients.aquaticscience.org/sites/default/files/u23/Report_Nutrient_load_to_South_Bay_2011-8-31_revised.pdf.

Milliman, J., and J. Syvitski (1992), Geomorphic/tectonic control of sediment discharge to the ocean: the importance of small mountainous rivers, *Journal of Geology*, 100(5), 525-544.

Milliman, J. D., K. L. Farnsworth, P. D. Jones, K. H. Xu, and L. C. Smith (2008), Climatic and anthropogenic factors affecting river discharge to the global ocean, 1951–2000, *Global and Planetary Change*, 62(3-4), 187-194.

Milly, P. C. D., J. Betancourt, M. Falkenmark, R. M. Hirsch, Z. W. Kundzewicz, D. P. Lettenmaier, and R. J. Stouffer (2008), Climate change. Stationarity is dead: Whither water management?, *Science*, 319(5863), 573-574.

Monismith, S. G., W. Kimmerer, J. R. Burau, and M. T. Stacey (2002), Structure and flow-induced variability of the subtidal salinity field in northern San Francisco Bay, *Journal of Physical Oceanography*, 32(11), 3003-3019.

Moore, S. K., N. J. Mantua, B. M. Hickey, and V. L. Trainer (2010), The relative influences of El Niño-Southern Oscillation and Pacific Decadal Oscillation on paralytic shellfish toxin accumulation in northwest Pacific shellfish, *Limnol Oceanogr*, 55, 2262-2274.

Moyle, P. B., J. R. Lund, W. A. Bennett, and W. E. Fleenor (2010), Habitat variability and complexity in the upper San Francisco Estuary, *San Francisco Estuary and Watershed Science*, 8(3), 1-24. Available at: <http://escholarship.org/uc/item/20kf20d32x>

Mueller-Solger, A. B., A. D. Jassby, and D. C. Mueller-Navarra (2002), Nutritional quality of food resources for zooplankton (*Daphnia*) in a tidal freshwater system (Sacramento-San Joaquin River Delta), *Limnol Oceanogr*, 47, 1468-1476.

Nichols, F. H. (1985), Increased benthic grazing: an alternative explanation for low phytoplankton biomass in northern San Francisco Bay during the 1976-77 drought, *Estuarine Coastal and Shelf Science*, 21, 379-388.

Nichols, F. H., J. K. Thompson, and L. E. Schemel (1990), Remarkable invasion of San Francisco Bay (California, USA) by the Asian clam *Potamocorbula amurensis*. II. Displacement of a former community, *Marine Ecology Progress Series*, 66, 95-101.

Nichols, F. H., J. E. Cloern, S. N. Luoma, and D. H. Peterson (1986), The modification of an estuary, *Science*, 231(4738), 567-573.

Nilsson, C., C. Reidy, M. Dynesius, and C. Revenga (2005), Fragmentation and flow regulation of the world's large river systems, *Science*, 308(520), 405-408.

Nixon, S. W. (1988), Physical energy inputs and the comparative ecology of lake and marine ecosystems, *Limnol Oceanogr*, 33(4), 1005-1025.

Nixon, S. W. (1995), Coastal marine eutrophication - a definition, social causes, and future concerns, *Ophelia*, 41, 199-219.

Nixon, S. W. (2003), Replacing the Nile: are anthropogenic nutrients providing the fertility once brought to the Mediterranean by the great river?, *Ambio*, 32(1), 30-39.

Nixon, S. W., R. W. Fulweiler, B. A. Buckley, S. L. Granger, B. L. Nowicki, and K. M. Henry (2009), The impact of changing climate on phenology, productivity, and benthic-pelagic coupling in Narragansett Bay, *Estuarine, Coastal and Shelf Science*, 82(1), 1-18.

Nobriga, M. L. (2002), Larval delta smelt diet composition and feeding incidence: environmental and ontogenetic influences, *California Fish and Game*, 88, 149-164.

Nobriga, M. L., T. Sommer, F. Feyrer, and K. Fleming (2008), Long-term trends in summertime habitat suitability for delta smelt, *Hypomesus transpacificus*, *San Francisco Estuary and Watershed Science*, 6(1), Available at: <http://repositories.cdlib.org/jmie/sfews/vol6/iss1/art1>

NRC (1993), Managing Wastewater in Coastal Urban Areas. Committee on Wastewater Management for Coastal Urban Areas, Water Science and Technology Board, Commission on Engineering and Technical Systems, National Research Council. Available at: http://stantonrealtors.com/listing_thumbnails/Managing-Wastewater-in-Coastal-Urban-Areas/p107178/.

NRC (2010), *A Scientific Assessment of Alternatives for Reducing Water Management Effects on Threatened and Endangered Fishes in California's Bay Delta*. , The National Academies Press. Available at: http://www.nap.edu/openbook.php?record_id=12881.

Odum, E. P. (1969), The strategy of ecosystem development, *Science*, 164(3877), 262-270.

Orth, R. J., et al. (2010), Long-term trends in submersed aquatic vegetation (SAV) in Chesapeake Bay, USA, related to water quality, *Estuar Coast*, 33(5), 1144-1163.

Parchaso, F., and J. K. Thompson (2002), Influence of hydrologic processes on reproduction of the

introduced bivalve *Potamocorbula amurensis* in northern San Francisco Bay, California, *Pacific Science*, 56, 329-345.

Parker, A. E., W. J. Kimmerer, and U. U. Lidström (2012), Reevaluating the generality of an empirical model for light-limited primary production in the San Francisco Estuary, *Estuar Coast*, 35(4), 930-942.

Perez, T., D. Longet, T. Schembri, P. Rebouillon, and J. Vacelet (2005), Effects of 12 years' operation of a sewage treatment plant on trace metal occurrence within a Mediterranean commercial sponge (*Spongia officinalis*, Demospongiae), *Marine Pollution Bulletin*, 50, 301-309.

Petersen, J. K., J. W. Hansen, M. B. Laursen, P. Clausen, J. Carstensen, and D. J. Conley (2008), Regime shift in a coastal marine ecosystem, *Ecol Appl*, 18(2), 497-510.

Peterson, H. A., and M. Vayssieres (2010), Benthic assemblage variability in the upper San Francisco Estuary: A 27-year retrospective, *San Francisco Estuary and Watershed Science*, 8(1). Available at <http://www.escholarship.org/uc/item/4d0616c6>.

Peterson, W. T., and F. B. Schwing (2003), A new climate regime in northeast pacific ecosystems, *Geophys Res Lett*, 30(17).

Pimentel, D., R. Zuniga, and D. Morrison (2005), Update on the environmental and economic costs associated with alien-invasive species in the United States, *Ecol Econ*, 52, 273-288.

Pimm, S. L., and A. Redfearn (1988), The variability of population densities, *Nature*, 334(6183), 613-614.

Pritchard, D. W. (1967), What is an estuary: physical viewpoint, in *Estuaries*, edited, pp. 3-5, American Association for the Advancement of Science, Washington D.C.

RDevelopmentCoreTeam (2012), *R: A Language and Environment for Statistical Computing*, R Foundation for Statistical Computing, Vienna ISBN 3-900051-07-0.

Riisgard, H. U. (2012), From fish to jellyfish in the eutrophicated Limfjorden (Denmark), *Estuar Coast*, 35, 701-713.

Roegner, G. C., J. A. Needoba, and A. M. Baptista (2011), Coastal upwelling supplies oxygen-depleted water to the Columbia River Estuary, *PLoS One*, e18672. doi:10.1371/journal.pone.0018672.

Ruiz, G. M., J. T. Carlton, E. D. Grosholz, and A. H. Hines (1997), Global invasions of marine and estuarine habitats by non-indigenous species: mechanisms, extent, and consequences, *American Zoologist*, 37(6), 621-632.

Santos, M. J., S. Khanna, E. L. Hestir, M. E. Andrew, S. S. Rajapakse, J. A. Greenberg, L. W. J. Anderson, and S. L. Ustin (2009), Use of hyperspectral remote sensing to evaluate efficacy of aquatic plant management, *Invasive Plant Science and Management*, 2, 216-229.

- Sanudo-Wilhelmy, S. A., and G. A. Gill (1999), Impact of the Clean Water Act on the levels of toxic metals in urban estuaries: the Hudson River Estuary revisited, *Environ Sci Technol*, 33(20), 3447-3481.
- Sarthou, G., K. R. Timmermans, S. Blain, and P. Tréguer (2005), Growth physiology and fate of diatoms in the ocean: a review, *Journal of Sea Research*, 53(1-2), 25-42.
- Scheffer, M., J. Bascompte, W. A. Brock, V. Brovkin, S. R. Carpenter, V. Dakos, H. Held, E. H. van Nes, M. Rietkerk, and G. Sugihara (2009), Early-warning signals for critical transitions, *Nature*, 461(7260), 53-59.
- Schoellhamer, D. H. (2011), Sudden clearing of estuarine waters upon crossing the threshold from transport- to supply-regulation of sediment transport as an erodible sediment pool is depleted: San Francisco Bay, 1999, *Estuar Coast*, 34, 885-899.
- Seitzinger, S. P., et al. (2010), Global river nutrient export: A scenario analysis of past and future trends, *Global Biogeochemical Cycles*, 24, GB0A08, 1-16.
- Smith, R. A., R. B. Alexander, and M. G. Wolman (1987), Water-quality trends in the nation's rivers, *Science*, 235(4796), 1607-1615.
- Smith, S. E., and S. Kato (1979), The fisheries of San Francisco Bay: past, present and future, in *San Francisco Bay: The Urbanized Estuary*, edited by T. J. Conomos, pp. 445-468, Pacific Division American Association for the Advancement of Science, San Francisco.
- Smith, S. V., R. M. Chambers, and J. T. Hollibaugh (1996), Dissolved and particulate nutrient transport through a coastal watershed-estuary system, *Journal of Hydrology*, 176(1-4), 181-203.
- Sobczak, W. V., J. E. Cloern, A. D. Jassby, and A. B. Mueller-Solger (2002), Bioavailability of organic matter in a highly disturbed estuary: The role of detrital and algal resources, *P Natl Acad Sci USA*, 99(12), 8101-8105.
- Sommer, T., et al. (2007), The collapse of pelagic fishes in the upper San Francisco Estuary, *Fisheries*, 32, 270-277.
- Southward, A. J. (1995), The importance of long time-series in understanding the variability of natural systems, *Helgoländer Meeresunters*, 49, 329-333.
- Southward, A. J., G. T. Boalch, and L. Maddock (1998), Fluctuations in the herring and pilchard fisheries of Devon and Cornwall linked to change in climate since the 16th century, *Mar Biol Assoc UK*, 68, 423-445.
- Steele, J. H. (1985), A comparison of terrestrial and marine ecological systems, *Nature*, 313(6001), 355-358.
- Steinberg, P., M. Brett, J. Bechtold, J. Richey, L. Porensky, and S. Smith (2010), The influence of

watershed characteristics on nitrogen export to and marine fate in Hood Canal, Washington, USA, *Biogeochemistry*, 106, 415-433.

Stewart, A. R., S. N. Luoma, C. E. Schlekat, M. A. Doblin, and K. A. Hieb (2004), Food web pathway determines how selenium affects aquatic ecosystems: A San Francisco Bay case study, *Environmental Science and Technology*, 38(17), 4519-4526.

Stige, L. C., G. Ottersen, K. Brander, K. S. Chan, and N. C. Stenseth (2006), Cod and climate: effect of the North Atlantic Oscillation on recruitment in the North Atlantic, *Mar. Ecol-Progr. Ser.*, 325, 227-241.

Stralberg, D., M. Brennan, J. C. Callaway, J. K. Wood, L. M. Schile, D. Jongsomjit, M. Kelly, V. T. Parker, and S. Crooks (2011), Evaluating tidal marsh sustainability in the face of sea-level rise: a hybrid modeling approach applied to San Francisco Bay, *PLoS One*, 6(11), e27388.

SWRCB (2010), Development of Flow Criteria for the Sacramento-San Joaquin Delta Ecosystem. Available at http://www.swrcb.ca.gov/waterrights/water_issues/programs/bay_delta/deltaflow/docs/final_rpt080310.pdf, Sacramento.

Syvitski, J. P. M. (2003), Supply and flux of sediment along hydrological pathways: research for the 21st century, *Global and Planetary Change*, 39, 1-11.

Takata, L., N. Dobroski, C. Scianni, and M. Falkner (2011), 2011 Biennial Report on the California Marine Invasive Species Program, edited, California State Lands Commission, Marine Facilities Division. Available at http://www.slc.ca.gov/spec_pub/mfd/ballast_water/Documents/2011_BiennialRpt_Final.pdf

Takekawa, J. Y., C. T. Lu, and R. T. Pratt (2001), Avian communities in baylands and artificial salt evaporation ponds of the San Francisco Bay estuary, *Hydrobiologia*, 466, 317-328.

Taleb, N. N. (2007), Black swans and the domains of statistics, *The American Statistician*, 61(3), 1-3.

Taş, S., E. Okus, and A. Aslan-Yilmaz (2006), The blooms of a cyanobacterium, *Microcystis cf. aeruginosa* in a severely polluted estuary, the Golden Horn, Turkey, *Estuar Coast Shelf S*, 68(3-4), 593-599.

Thebault, J., T. S. Schraga, J. E. Cloern, and E. G. Dunlavy (2008), Primary production and carrying capacity of former salt ponds after reconnection to San Francisco Bay, *Wetlands*, 28(3), 841-851.

Thompson, J. K. (2005), One estuary, one invasion, two responses: phytoplankton and benthic community dynamics determine the effect of an estuarine invasive suspension feeder, in *The comparative roles of suspension feeders in ecosystems*, edited by S. Olenin and R. Dame, pp. 291-316, Springer, Amsterdam.

Thomson, J. R., W. J. Kimmerer, L. R. Brown, K. B. Newman, R. M. Nally, W. A. Bennett, F. Feyrer, and E. Fleishman (2010), Bayesian change point analysis of abundance trends for pelagic fishes in the

upper San Francisco Estuary, *Ecol Appl*, 20(5), 1431-1448.

Twardowski, M. S., and P. L. Donaghay (2001), Separating in situ and terrigenous sources of absorption by dissolved materials in coastal waters, *J. Geophys. Res.*, 106, 2545-2560.

USDA (2010), California Agricultural Statistics 2008 Crop Year. Available at: http://www.nass.usda.gov/Statistics_by_State/California/Publications/California_Ag_Statistics/Reports/2009cas-all.pdf

USEPA (2000), Progress in Water Quality. An Evaluation of the National Investment in Municipal Wastewater Treatment. *Rep.*, United States Environmental Protection Agency, Washington, D.C.

Van Drecht, G., A. F. Bouwman, J. Harrison, and J. M. Knoop (2009), Global nitrogen and phosphate in urban wastewater for the period 1970 to 2050, *Global Biogeochemical Cycles*, 23, GBA03, 1-19.

van Geen, A., and S. N. Luoma (1999), The impact of human activities on sediments of San Francisco Bay, California: an overview, *Marine Chemistry*, 64, 1-6.

Vasseur, D. A., and P. Yodzis (2004), The color of environmental noise, *Ecology*, 85(4), 1146-1152.

Vericat, D., and R. Batalla (2006), Sediment transport in a large impounded river: The lower Ebro, NE Iberian peninsula, *Geomorphology*, 79(1), 72-92.

Vitousek, P. M., C. M. D'Antonio, L. L. Loope, and R. Westbrooks (1996), Biological invasions as global environmental change, *American Scientist*, 84(5), 468-478.

Vitousek, P. M., H. A. Mooney, J. Lubchenco, and J. M. Melillo (1997), Human domination of the Earth's ecosystems, *Science*, 277, 494-499.

Vollenweider, R. A. (1975), Input-output models with special reference to the phosphorus loading concept in limnology, *Schweizerische Zeitschrift für Hydrologie*, 37, 53-84.

Vörösmarty, C., M. Meybeck, B. Fekete, K. Sharma, P. Green, and J. Syvitski (2003), Anthropogenic sediment retention: major global impact from registered river impoundments, *Global and Planetary Change*, 39(1), 169-190.

Walters, R. A., R. T. Cheng, and T. J. Conomos (1985), Time scales of circulation and mixing processes of San Francisco Bay waters, *Hydrobiologia*, 129, 13-36.

Wasson, K., C. Zabin, L. Bedinger, M. Cristina Diaz, and J. Pearse (2001), Biological invasions of estuaries without international shipping: the importance of intraregional transport, *Biological Conservation*, 102(2), 143-153.

Werner, I., and J. T. Hollibaugh (1993), *Potamocorbula amurensis*: Comparison of clearance rates and assimilation efficiencies for phytoplankton and bacterioplankton, *Limnol Oceanogr*, 1993, 949-964.

Whitall, D., S. Bricker, J. Ferreira, A. M. Nobre, T. Simas, and M. Silva (2007), Assessment of eutrophication in estuaries: pressure-state-response and nitrogen source apportionment, *Environmental Management*, 40(4), 678-690.

Wickham, H. (2009), *ggplot2: Elegant Graphics for Data Analysis*, Springer, New York.

Williams, S. L., and E. D. Grosholz (2008), The invasive species challenge in estuarine and coastal environments: Marrying management and science, *Estuar Coast*, 31(1), 3-20.

Winder, M., and A. D. Jassby (2011), Shifts in zooplankton community structure: Implications for food-web processes in the upper San Francisco Estuary, *Estuar Coast*, 34(4), 675-690.

Winder, M., A. D. Jassby, and R. Mac Nally (2011), Synergies between climate anomalies and hydrological modifications facilitate estuarine biotic invasions, *Ecol Lett*, 14, 749-757.

Wolff, W. (1998), Exotic invaders of the meso-oligohaline zone of estuaries in the Netherlands: why are there so many?, *Helgoland Marine Research*, 52(3), 393-400.

Wright, S. A., and D. H. Schoellhamer (2004), Trends in the sediment yield of the Sacramento River, California, 1957-2001, *San Francisco Estuary and Watershed Science* 2(2), 1-14. Available at: <http://escholarship.ucop.edu/uc/item/891144f891144 - page-891141>.

Xu, J., J. H. W. Lee, K. Yin, H. Liu, and P. J. Harrison (2011), Environmental response to sewage treatment strategies: Hong Kong's experience in long term water quality monitoring, *Marine Pollution Bulletin*, 62(11), 2275-2287.

Yang, S. L., J. D. Milliman, P. Li, and K. Xu (2011), 50,000 dams later: Erosion of the Yangtze River and its delta, *Global and Planetary Change*, 75(1-2), 14-20.

Yasuhara, M., H. Yamazaki, T. A., and H. K. (2007), The effect of long-term spatiotemporal variations in urbanization-induced eutrophication on a benthic ecosystem, Osaka Bay, Japan, *Limnol Oceanogr*, 52, 1633-1644.

Figure Captions

Figure 1. Examples of change in estuaries, as altered: *A.* sediment supply to the Yangtze Estuary; *B.* total phosphorus loading to the Potomac Estuary; *C.* index of light scattering in Chesapeake Bay; *D.* biomass of benthic invertebrates in Ringkøbing Fjord; *E.* landings of pelagic fish in Limfjorden; *F.* dissolved oxygen concentration in bottom waters of the Potomac Estuary during summer; *G.* uptake of silicate as an index of diatom primary productivity in northern San Francisco Bay; *H.* area of submerged vascular plants (SAV) in the lower Potomac River; *I.* concentrations of mercury in mussels collected in the Forth Estuary; and *J.* abundance of black-tailed godwits in the Tagus Estuary. Data provided by: *A.* Shilun Yang (East China Normal University); *B., F.* Norbert Jaworski (USEPA, retired); *C.* Charles Gallegos (Smithsonian Environmental Research Center); *D.* Jens Würgler Hansen (Aarhus University, Denmark); *E.* Hans Ulrik Riisgard (University of Southern Denmark); *G.* Wim Kimmerer (San Francisco State University); *H.* David Wilcox (Virginia Institute of Marine Science); *I.* Judith Dobson (Scottish Environment Protection Agency); *J.* Teresa Catry (CESAM/Museu Nacional de História Natural, Portugal).

Figure 2. San Francisco Bay, fed by the waters of the Sacramento-San Joaquin River Delta and connected to the Pacific Ocean at the Golden Gate. The boundary between Bay and Delta is specified to be at Chipps Island. Water is exported from the southern Delta via state (SWP) and federal (CVP) water project canals. Numbers labeled "X2" represent distances along the axis of the estuary from the Golden Gate. We use observations at sampling sites shown in South Bay and Suisun Bay to illustrate drivers of change detected over the past four decades.

Figure 3. Unimpaired runoff for the water year (October-September), based on measured flows in the major tributaries to San Francisco Bay upstream of storage and diversion points. *Blue line*, a loess smoother with 0.95 confidence interval.

Figure 4. Growth of the total population of San Francisco Bay Area counties, 1860-2010.

Figure 5. Growth in California reservoir capacity since 1900. The 10 largest reservoirs are labeled next to the corresponding step increase in capacity. *Green line*, mean unimpaired runoff

for water years 1906-2010 (Figure 3).

Figure 6. The fate of tributary water to San Francisco Bay as a percentage of total unimpaired inflow to the Delta during 1956-2003, i.e., the inflow to the Delta that would have occurred in the absence of upstream human activities. *Left panel:* The major fates include net upstream use (including consumption, reservoir storage or release, and import or diversion); Delta use; and outflow from the Delta to the Bay. *Right panel:* Uses in the Delta can be classified as exports to state and federal water projects and depletions within the Delta (the net result of consumption, precipitation and evapotranspiration).

Figure 7. *A.* Water-year mean exports from the Delta. *B.* Exports as a percent of total inflow to the Delta. *Blue lines,* loess smoothers with 0.95 confidence intervals.

Figure 8. Long-term trends (1956-2010) in three important flow variables for San Francisco Bay. *A.* Total measured inflow to the Delta. *B.* Exports from the Delta to state, federal and local water projects. *C.* Net outflow from the Delta past Chipps Island (see Figure 2) to San Francisco Bay. *Blue shading,* significant trends ($p < 0.05$).

Figure 9. Monthly mean suspended particulate matter concentrations measured in surface waters at two locations in Suisun Bay. *A.* Grizzly Bay, a shallow subembayment location near Suisun Slough (Figure 2, site D7). *B.* In the deep channel off of Middle Point (site D8). *Blue lines,* loess smoothers with 0.95 confidence intervals.

Figure 10. Partial residual plots for a regression model that accounts for variability in annual mean suspended particulate matter in Suisun Bay (D8) as a result of a long-term trend plus variability in annual mean outflow from the Delta. *A.* The linear effect of trend. *B.* The nonlinear effect of outflow. *Blue lines,* loess smoothers with 0.95 confidence intervals.

Figure 11. Response of the planktonic food web in Suisun Bay to an introduced clam, *Corbula amurensis*. *A.* *Corbula* abundance. *B.* Phytoplankton biomass as chlorophyll *a* concentration. *C.* Density of the rotifer *Synchaeta bicornis*. *D.* Density of the copepod *Eurytemora affinis*. *E.*

Density of the mysid shrimp *Neomysis mercedis*. The horizontal line spanning each panel *B* through *E* is the long-term mean.

Figure 12. Boxplot summaries of monthly mean chlorophyll *a* in a shallow subembayment of Suisun Bay (Figure 2, site D7) before and after introduction of the clam *Corbula amurensis* in 1987. *Green line*, characteristic threshold concentration at which zooplankton growth or reproduction can become food-limited.

Figure 13. Boxplots showing spatial distributions of dissolved inorganic nitrogen (DIN) and phosphate (dissolved inorganic phosphorus, DIP) in surface waters (0-3 m) of South San Francisco Bay, 1969-2010 (sampling locations shown in Figure 2). Five extreme DIN values > 200 or $< 1 \mu\text{mol L}^{-1}$ are omitted. *Green lines*, characteristic half-saturation constants for DIN and phosphate uptake, respectively, as indices of nutrient levels that potentially limit phytoplankton growth.

Figure 14. Boxplots of dissolved inorganic nitrogen (DIN) and phosphate in South San Francisco Bay (1969-2010), Tomales Bay (1987-1995), Willapa Bay (1991-2006), and the deep channel of Chesapeake Bay (2006-2010). The data are from all available depths. *Green line*, characteristic half-saturation constant for phytoplankton growth rate.

Figure 15. Responses in South San Francisco Bay to improvements in sewage treatment efficiency. *A.* Annual loads of BOD and ammonium from the San Jose-Santa Clara wastewater treatment plant. *B.* Annual loads of copper and silver from the Palo Alto Regional wastewater treatment plant. *C.* Dissolved oxygen south of the Dumbarton Bridge. *Green line*, a common standard to protect marine fish sensitive to low oxygen. *D.* Copper (red) and silver (blue) concentrations in clam (*Macoma balthica*) tissue from a Palo Alto mudflat.

Figure 16. Time series of annual mean climate indices, ocean conditions near the mouth of San Francisco Bay, and annual mean abundances of various biota within San Francisco Bay, shown as anomalies about the long-term means. *A.* North Pacific Gyre Oscillation. *B.* Pacific Decadal Oscillation. *C.* Sea surface temperature at Farallon Islands. *D.* Upwelling index at 39°N. *E.* Sum

of catches per unit effort in the marine subembayments (South, Central and San Pablo bays) for five species of demersal fish (age-0 English sole, speckled sanddab, plainfin midshipman, bay goby, Pacific staghorn sculpin), *F.* three species of crabs (age-0 Dungeness, slender, and brown rock crab), and *G.* two species of shrimp (blacktail bay shrimp, Stimpson coastal shrimp). *H.* Dry weight of clams from all available shallow sampling sites in South Bay. *I.* Annual mean phytoplankton biomass (chlorophyll *a*) in surface waters of the three South Bay stations sampled most frequently (24, 27 and 30).

Figure 17. Ecosystem regime shift, depicted in boxplot distributions of the time series in Figure 16, divided into years before and after the 1998-1999 climate shift.

Figure 18. Behavior of the South San Francisco Bay phytoplankton community around the time of a Pacific Ocean regime shift in 1998-1999. *A.* Monthly mean phytoplankton biomass (chlorophyll *a*) in surface waters of the three South Bay stations sampled most frequently (24, 27 and 30). Arrow shows the appearance of the first autumn-winter bloom in October 1999. *B.* Calculated annual gross primary production averaged for the same three stations. *C.* Boxplot distributions of the fulcrum, i.e., the timing of the center of gravity of the annual chlorophyll pattern, for the two eras. *D.* Occurrence of four phytoplankton species in South and Central bays. Circles are plotted at each date when the indicated species was detected. *Thalassiosira rotula* and *Thalassiosira punctigera*, both centric diatoms, were the dominant species before and after the shift, respectively. *Oxytoxum milneri* gave way to *Polykrikos schwartzii*, both heterotrophic dinoflagellates.

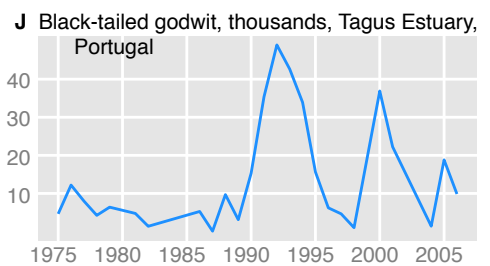
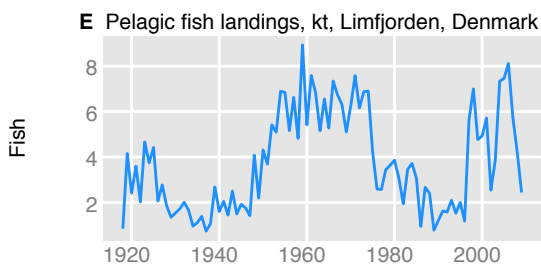
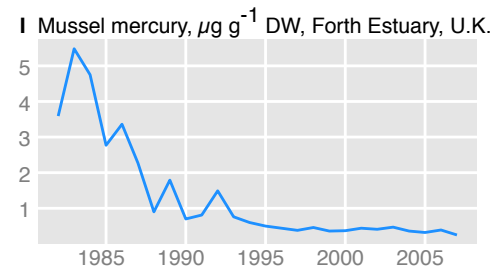
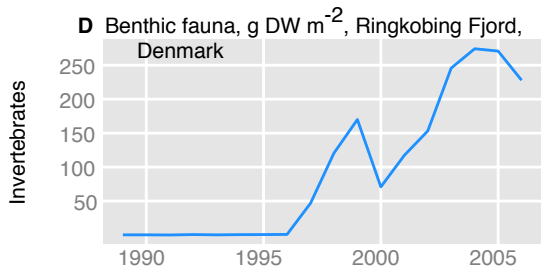
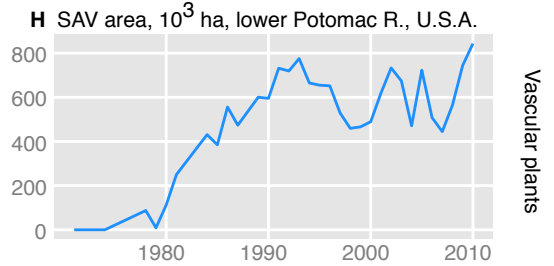
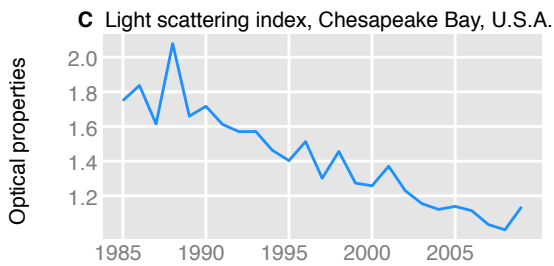
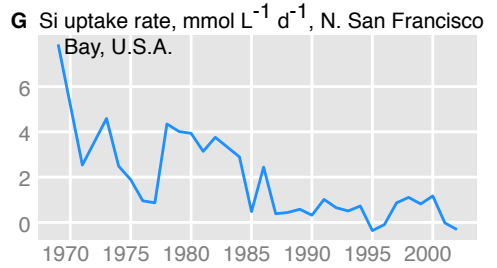
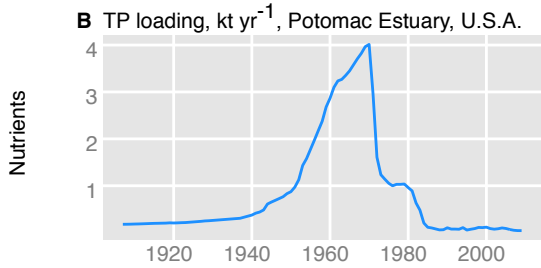
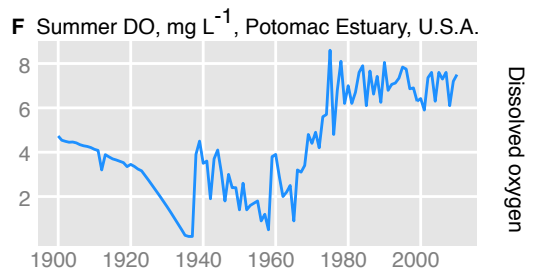
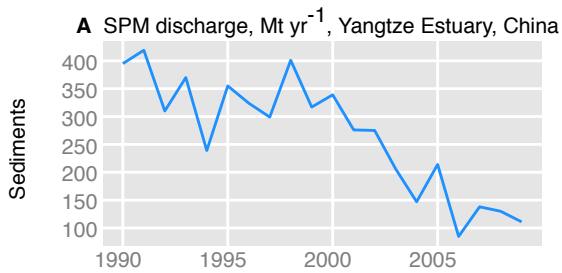
Table Captions

Table 1. Attributes of Suisun Bay and South Bay as contrasting estuarine habitats of San Francisco Bay, including their: dimensions, water residence time, phytoplankton primary production, and quartile values of water-quality constituents from sampling by the USGS (stations 4-7 [Suisun Bay] and 20-36 [South Bay]) and IEP (stations D6, D7, D8, D10 [Suisun Bay]) from 1969 through 2010.

Table 2. Decadal averages of X_2 (km) for September-December. X_2 , estimated from outflow; X_2^* , estimated from unimpaired outflow; ΔX_2 , difference between them; SD, standard deviation of difference.

Table 3. Annual inputs of dissolved inorganic nitrogen (DIN) to US estuaries and bays from treated sewage, and the sewage contribution to total DIN input.

Table A1. Sources of data used in this review.

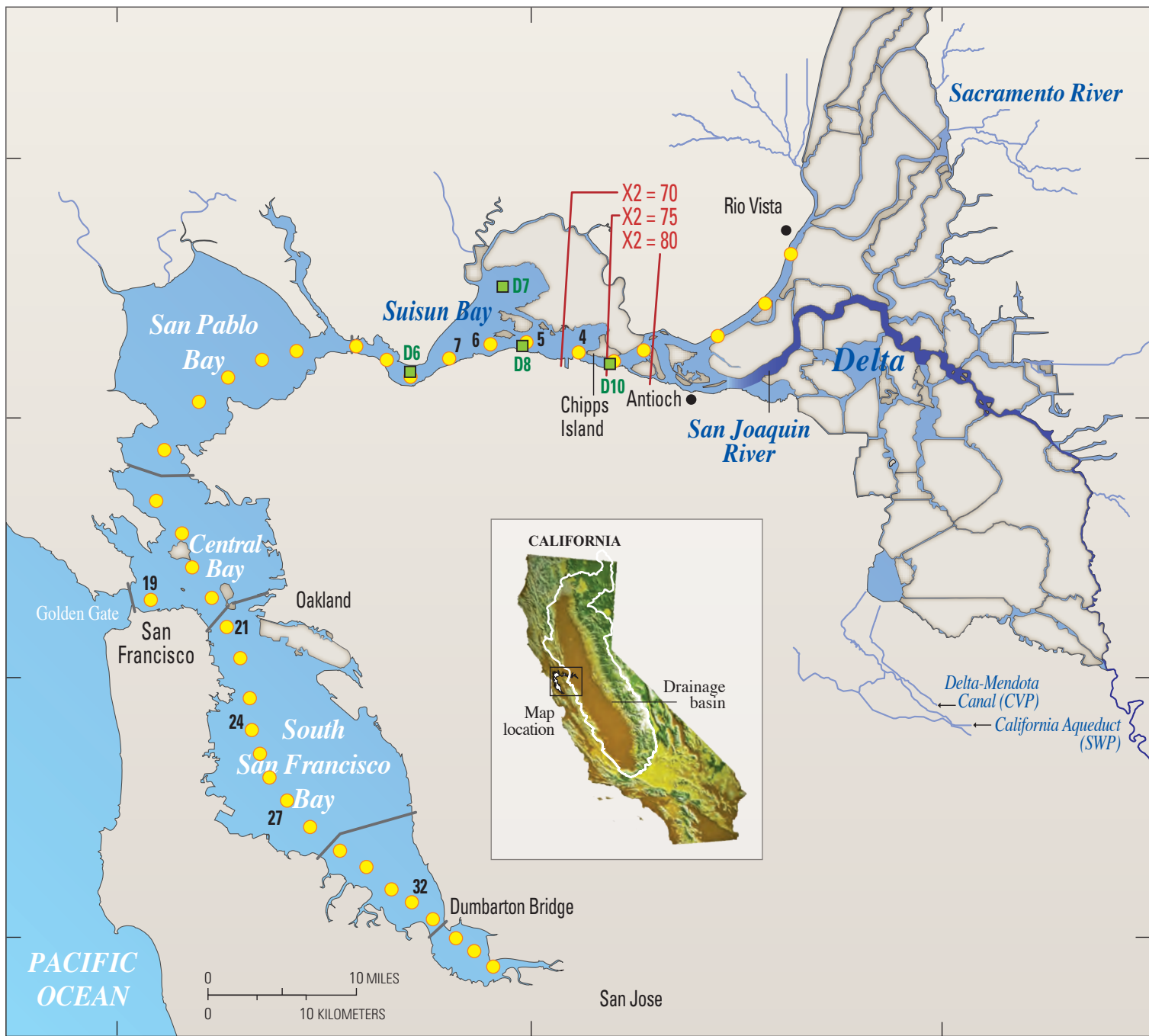


122°30'

122°00'

121°30'

38°00'



X2 = 70
 X2 = 75
 X2 = 80



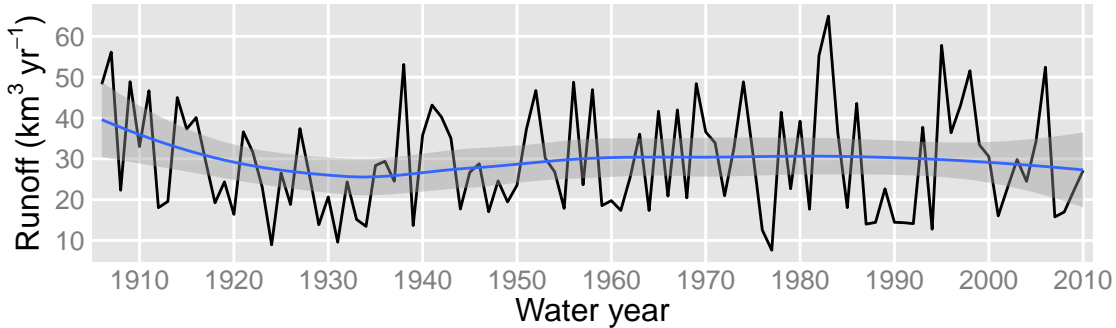
Delta-Mendota Canal (CVP)

California Aqueduct (SWP)

0 10 MILES

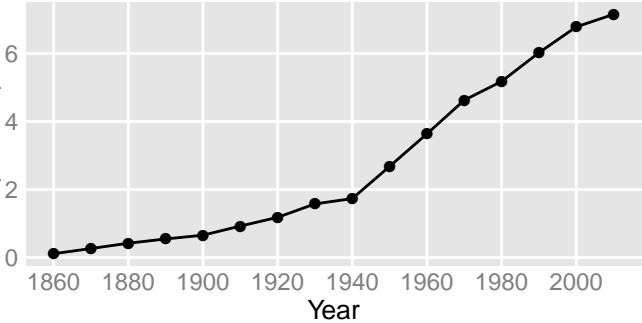
0 10 KILOMETERS

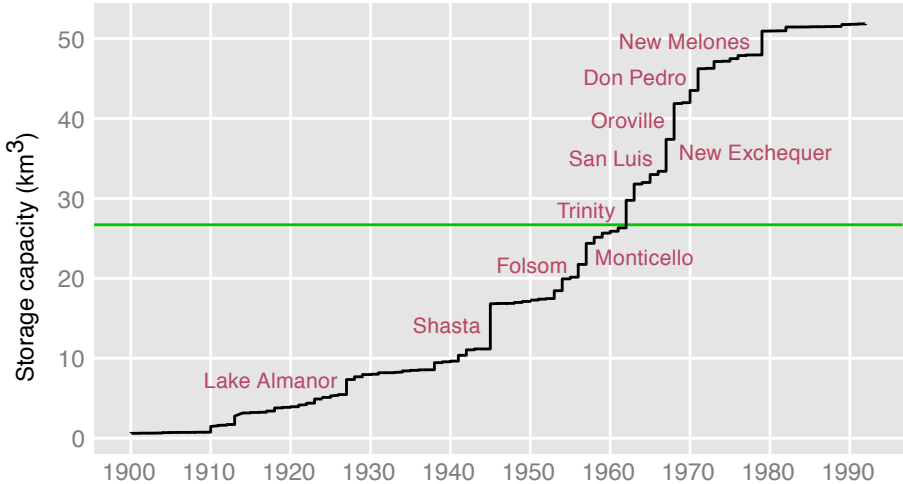
San Jose



Bay Area Population

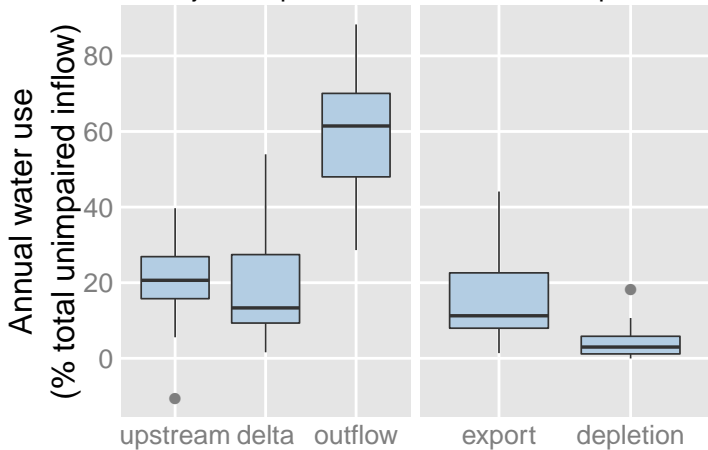
(millions)

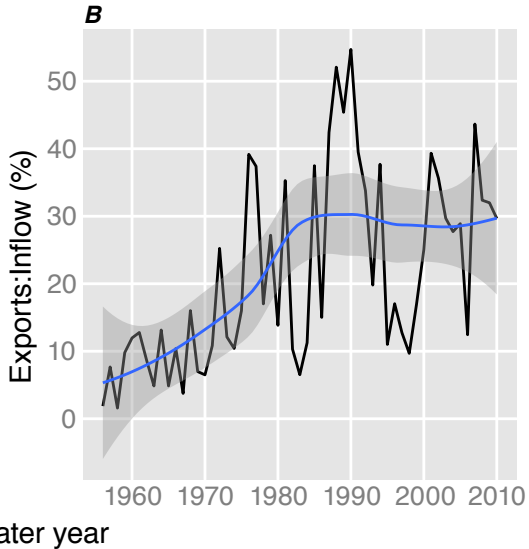
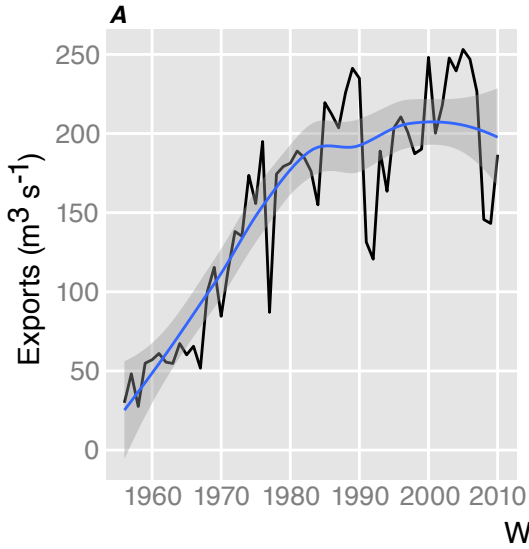


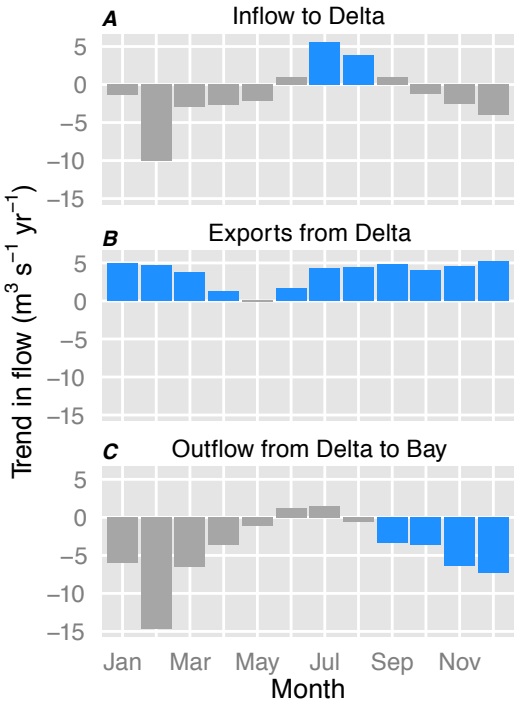


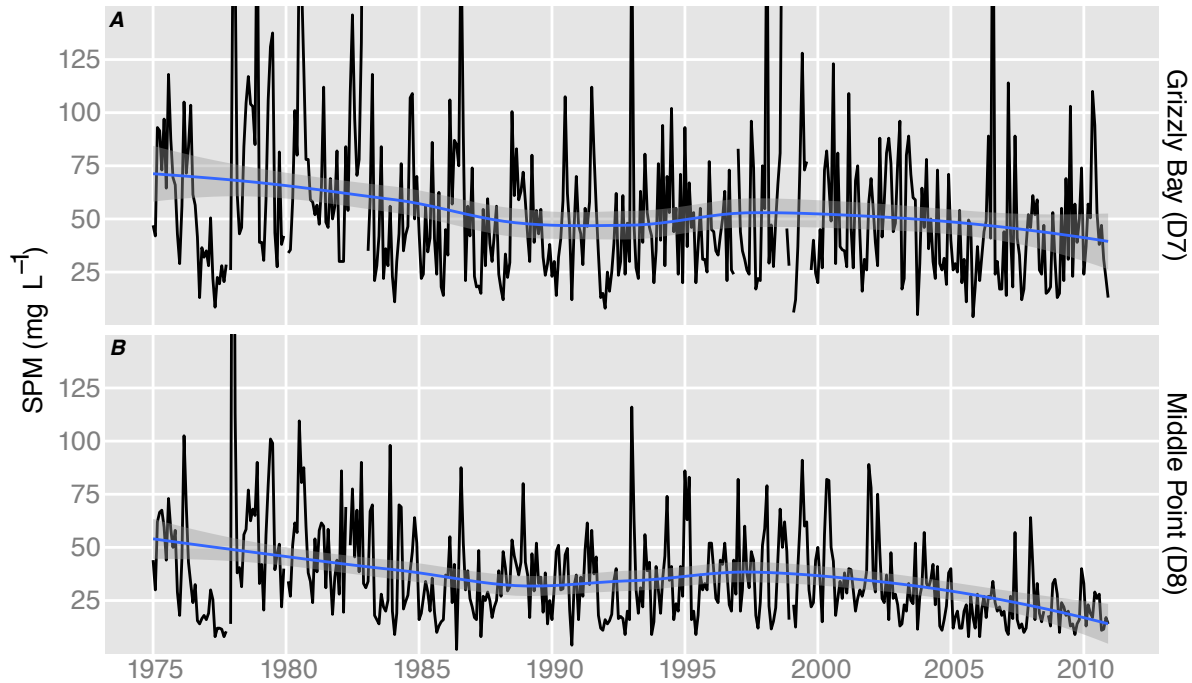
Major components

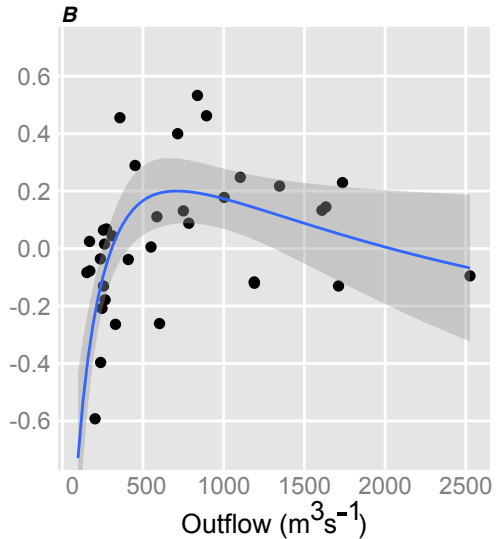
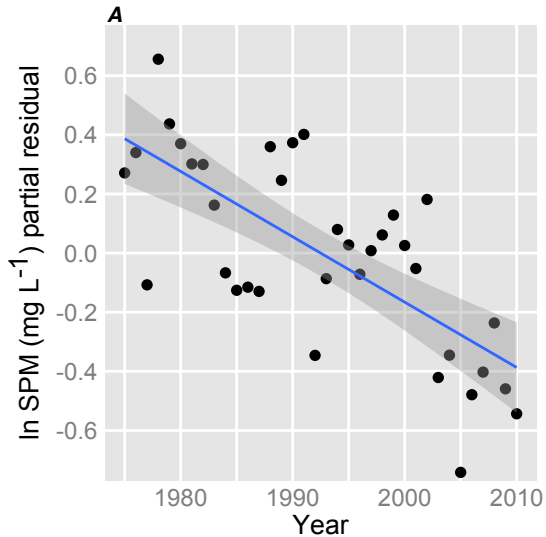
Delta subcomponents

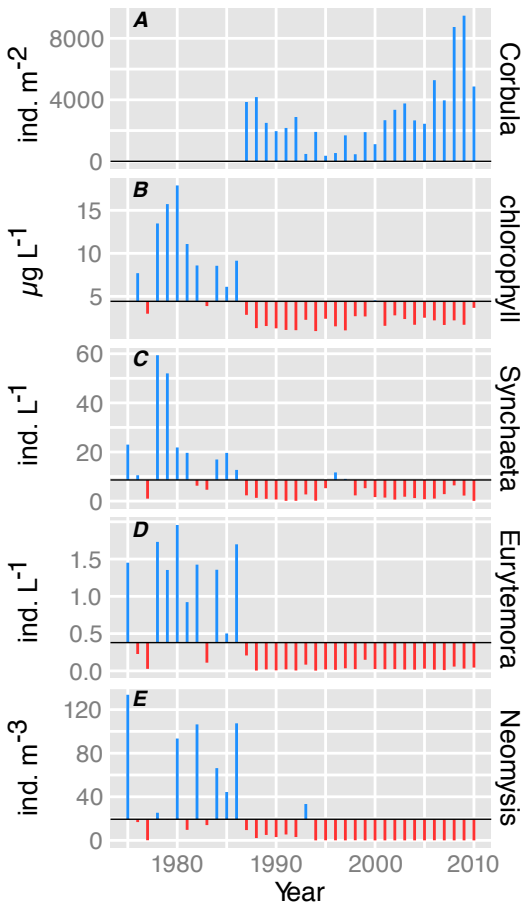




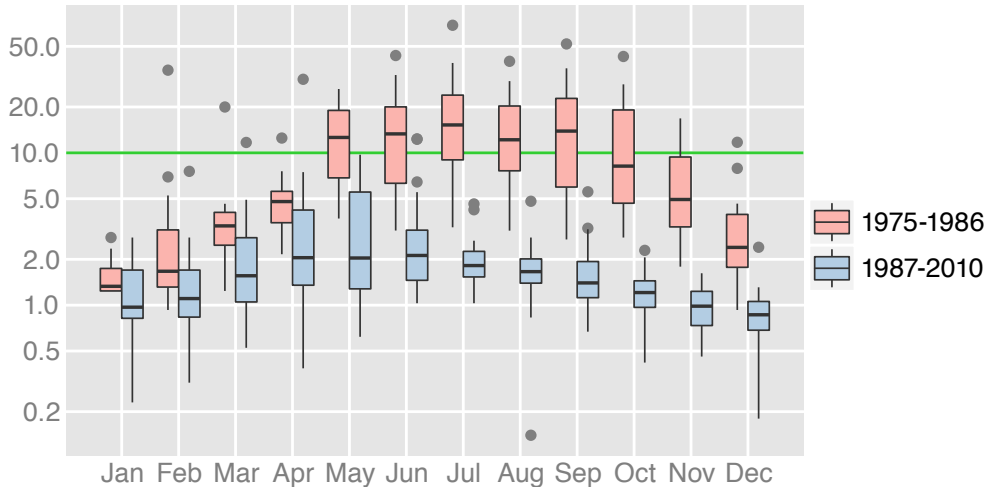




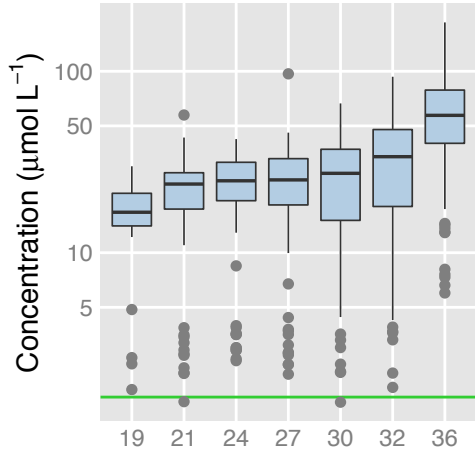




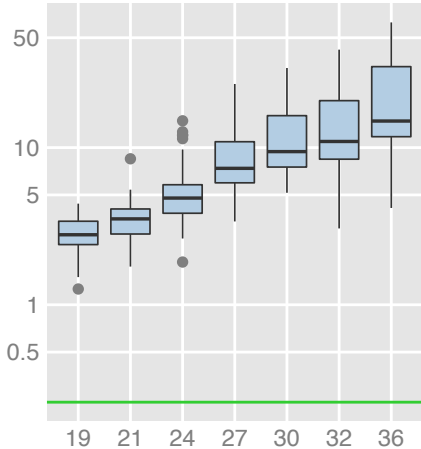
Chlorophyll *a* ($\mu\text{g L}^{-1}$)



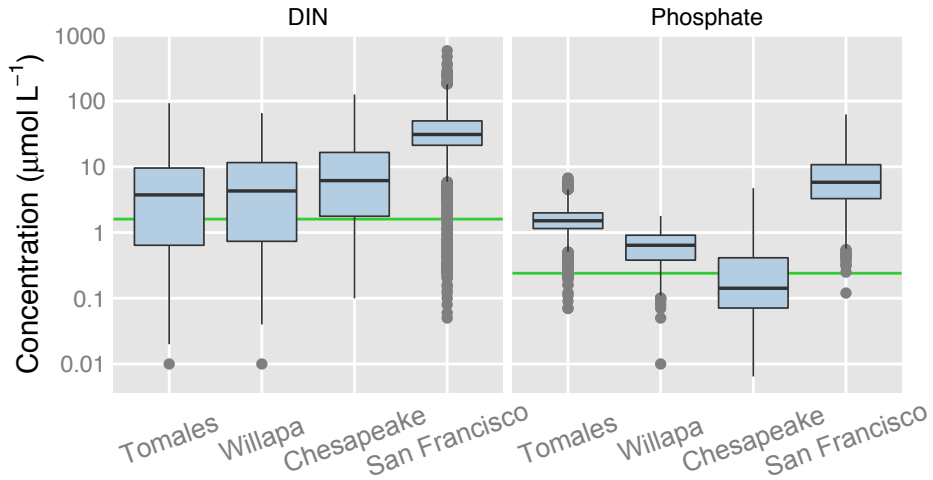
DIN

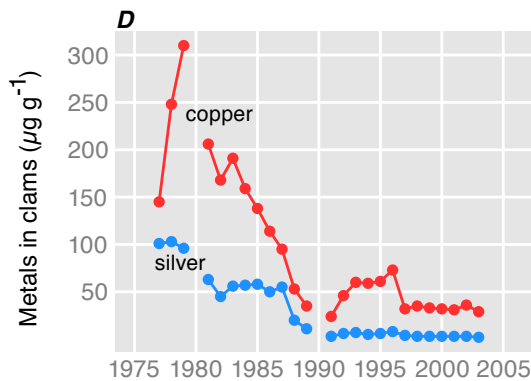
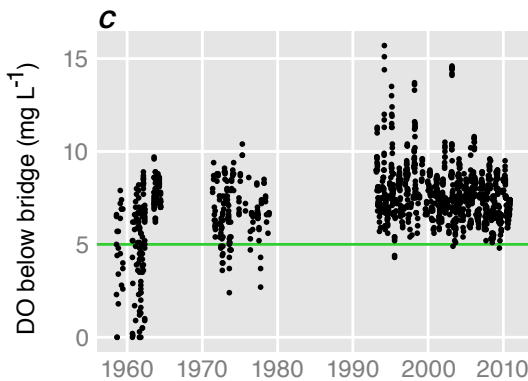
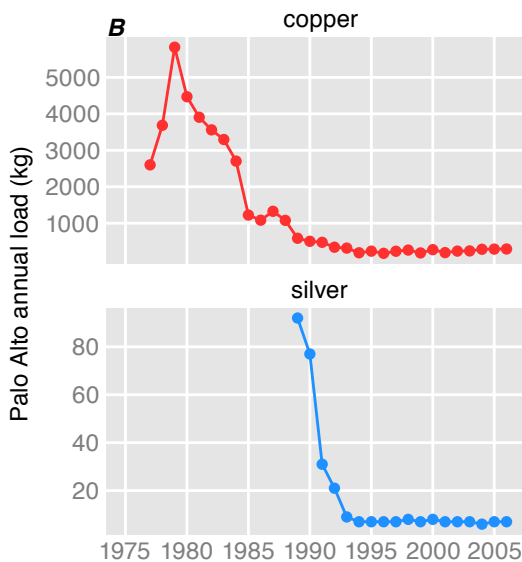
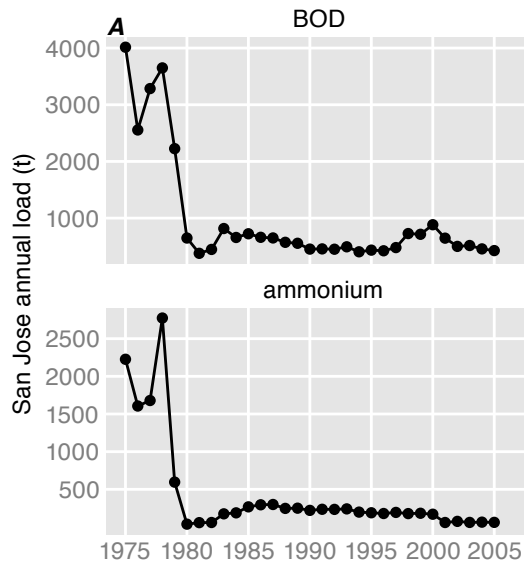


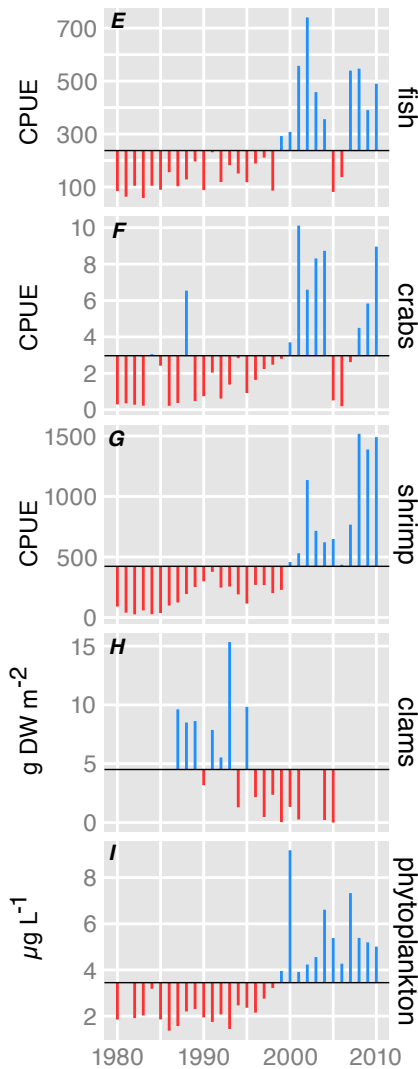
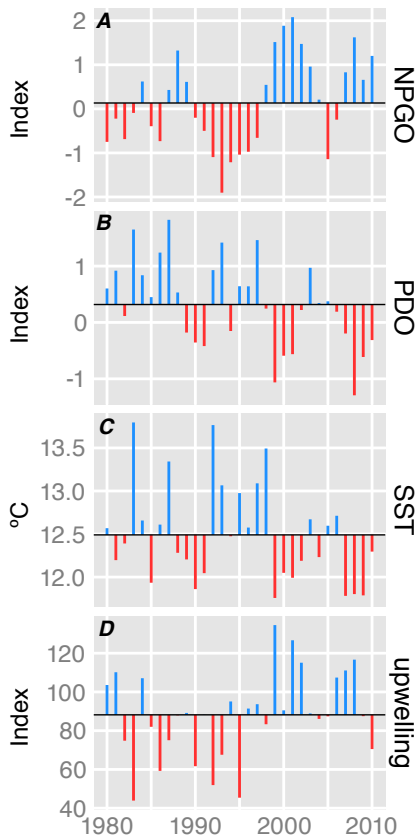
Phosphate

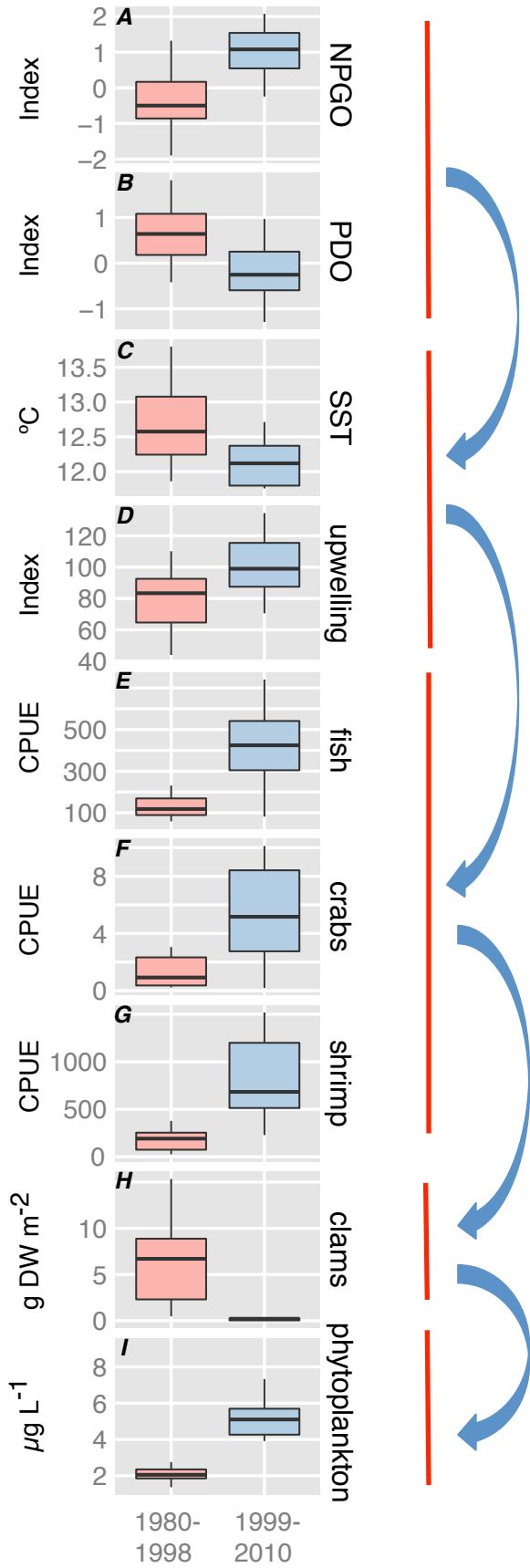


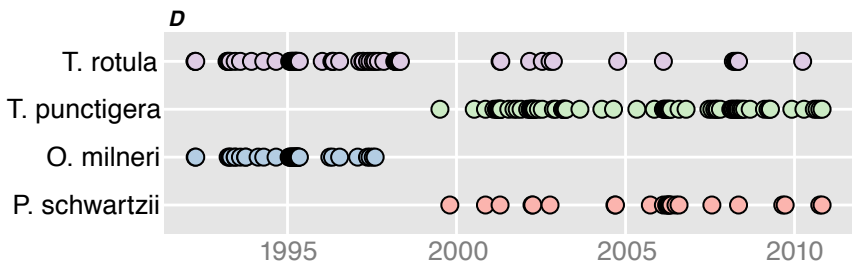
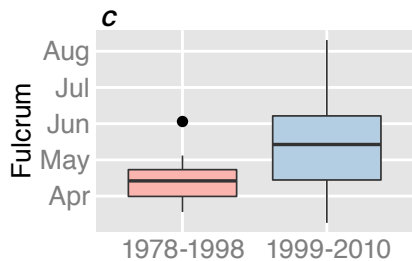
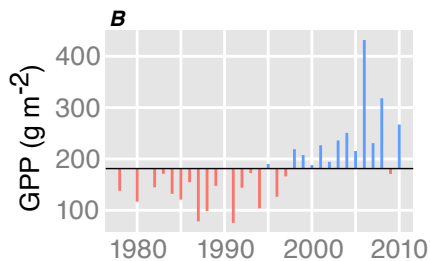
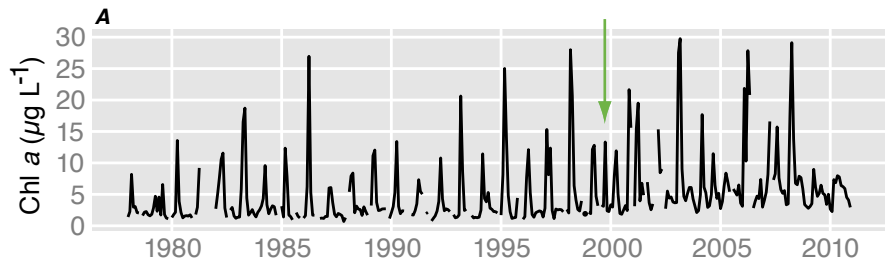
Station











	Median	25th Percentile	75th Percentile	Median	25th Percentile	75th Percentile
	<i>Suisun Bay</i>			<i>South Bay</i>		
Surface Area (km ²), MSL ^a	170			430		
Mean Depth (m), MSL ^a	4.6			5.8		
Mean Tidal Range (m) ^b	0.9 - 1.3			1.3 - 2.3		
Residence Time (d) ^c	0.5 - 35			14 - 160		
Primary Production (g C m ⁻² yr ⁻¹) ^d	20 - 130			130 - 210		
Salinity	5.8	1.0	10.7	27.4	23.8	30.0
Temperature (°C)	17.1	12.4	20.0	15.1	12.8	17.7
Chlorophyll <i>a</i> (µg L ⁻¹)	2.0	1.4	3.0	4.1	2.4	7.3
NO ₃ + NO ₂ (µM)	23.6	15.7	31.4	21.7	14.7	32.1
NH ₄ (µM)	4.3	2.3	7.1	6.1	3.4	8.7
PO ₄ (µM)	2.3	1.8	2.9	4.8	2.9	8.6
SiO ₄ (µM)	201	163	240	83	60	109
Suspended Particulate Matter SPM (mg L ⁻¹)	39	25	63	17	10	31
Attenuation coefficient <i>k</i> (m ⁻¹)	2.7	1.9	3.8	1.4	1.0	2.0
Dissolved Oxygen (mg L ⁻¹)	8.7	8.2	9.4	7.9	7.1	8.6

^a [USGS, 2007]; ^b [NOAA, 2006]; ^c [Walters *et al.*, 1985]; ^d [Alpine and Cloern, 1992; Cloern, 1987; Cloern *et al.*, 1985]

Decade	X2	X2*	$\Delta X2$	SD
1950-1959		73.7		
1956-1959	73.2	75.9	-2.7	2.1
1960-1969	71.3	73.3	-2.0	2.8
1970-1979	73.3	73.7	-0.5	5.2
1980-1989	75.1	72.5	2.6	4.7
1990-1999	78.6	75.9	2.7	2.9
2000-2003	79.9	74.2	5.6	0.7
2000-2010	80.5			

Estuary	Sewage DIN Input mmol N m ⁻² yr ⁻¹	Sewage %	Total DIN Input g N m ⁻² yr ⁻¹
Tomales Bay ^a	0	0	2
Apalachicola Bay ^b	10	2	8
Mobile Bay ^b	80	7	18
Chesapeake Bay ^c	NA	NA	14
Narragansett Bay ^b	390	41	13
Potomac Estuary ^b	390	48	11
Delaware Bay ^b	650	50	18
Long Island Sound ^b	270	67	6
New York Bay ^b	27,230	82	447
South San Francisco Bay ^d	1,860	92	28

^a [S V Smith et al., 1996]; ^b [Nixon and Pilson, 1983]; ^c [Kemp et al., 2005]; ^d [McKee and Gluchowski, 2011]

Description	Source	Date Accessed
a. Census populations of Bay Area counties and incorporated cities	http://www.dof.ca.gov/research/demographic/state_census_data_center/historical_census_1850-2010/view.php	2012-01-12
b. Measured unimpaired runoff to the Sacramento and San Joaquin valleys	http://cdec.water.ca.gov/cgi-progs/iodir/wsihist	2011-10-28
c. Reservoir storage volume in California	[CADWR, 1993]	-
d. Estimated unimpaired inflow to the Delta	[CADWR, 2007]	-
e. Dayflow Program flow data	http://www.water.ca.gov/dayflow/output/Output.cfm	2011-02-01
f. IEP Environmental Monitoring Program discrete water quality data	http://www.water.ca.gov/bdma/meta/Discrete/data.cfm	2011-03-20
g. USGS Water Quality of San Francisco Bay discrete water quality data	http://sfbay.wr.usgs.gov/access/wqdata/query/index.html	2011-06-03
h. IEP Environmental Monitoring Program discrete benthic data	http://www.water.ca.gov/bdma/meta/benthic/data.cfm	2011-10-10
i. IEP Environmental Monitoring Program discrete zooplankton data	http://www.water.ca.gov/bdma/meta/zooplankton.cfm	2011-09-07
j. Chesapeake Bay nutrients	http://www.chesapeakebay.net/data/downloads/cbp_water_quality_database_1984_present	2012-02-17
k. Tomales Bay nutrients	http://lmer.marsci.uga.edu/tomales/	2011-12-29

Description	Source	Date Accessed
l. Willapa Bay nutrients	http://www.ecy.wa.gov/programs/eap/mar_wat/mwm_intr.html	2012-01-27
m. SERL DO in South Bay	[Harris <i>et al.</i> , 1961; McCarty <i>et al.</i> , 1962]	-
n. San Jose/Santa Clara Water Pollution Control Plant performance summary	Neal Van Keuren, City of San Jose, Environmental Services	-
o. Palo Alto Regional Water Quality Control Plant metal loadings	Michelle Hornberger, USGS, personal communication, January 2012	-
p. USGS Ecology and Contaminants Program, metals in Palo Alto clams	http://www.rcamnl.wr.usgs.gov/tracel/	2012-02-09
q. PDO index	http://jisao.washington.edu/pdo/	2012-01-05
r. NPGO index	http://www.o3d.org/npgo/data/NPGO.txt	2012-01-05
s. Sea surface temperature, SE Farallon Island	http://www.sccoos.org/query/	2012-01-05
t. Upwelling index	http://www.pfeg.noaa.gov/products/pfel/modeled/indices/upwelling/upwelling.html	2012-01-10
u. California Department of Fish and Game, San Francisco Bay Study	Kathy Hieb, DFG, personal communication, 24 January 2012 and earlier	-