INTERAGENCY ECOLOGICAL PROGRAM, MANAGEMENT, ANALYSIS, AND SYNTHESIS TEAM

An updated conceptual model of Delta Smelt biology: our evolving understanding of an estuarine fish

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Fall Midwater Trawl survey crew deploying net, circa 2005. Photo from CDFW.

Cover photo by Steven Culberson, USFWS
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**Table 8.** Summary of relationships between the log-transformed 20 mm abundance index for Delta Smelt (response variable) and one or more predictor variables. All relationships modeled with simple least-squares linear models (LM). For explanation of column headings see Table 6.  

**Table 9.** Summary of relationships of larval recruitment indices (abundance index ratios) for Delta Smelt (response variable) and spring X2 (predictor variable; spring: February-June): n, number of observations (years); SE/Mean, model standard error (square root of mean squared residual) as proportion of mean response, P, statistical significance level for the model; R$^2$, coefficient of determination. All relationships modeled with least-squares linear models (LM).  

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An updated conceptual model of Delta Smelt biology: our evolving understanding of an estuarine fish

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**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>CCF</td>
<td>Clifton Court Forebay</td>
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<tr>
<td>CVP</td>
<td>Central Valley Project</td>
</tr>
<tr>
<td>Delta</td>
<td>Sacramento-San Joaquin River Delta</td>
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<tr>
<td>DRERIP</td>
<td>Delta Regional Ecosystem Restoration Implementation Program</td>
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<td>Delta Stewardship Council</td>
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<td>EMP</td>
<td>Environmental Monitoring Program</td>
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<td>FLaSH</td>
<td>Fall Low Salinity Habitat</td>
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<td>FMWT</td>
<td>Fall Midwater Trawl Survey</td>
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<td>IEP</td>
<td>Interagency Ecological Program</td>
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<tr>
<td>LSZ</td>
<td>low salinity zone</td>
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<tr>
<td>MAST</td>
<td>Management, Analysis, and Synthesis Team</td>
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<tr>
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<td>National Research Council</td>
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<tr>
<td>OMR</td>
<td>Old and Middle River</td>
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<td>POD</td>
<td>Pelagic organism decline</td>
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<tr>
<td>SFE</td>
<td>San Francisco Estuary</td>
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<tr>
<td>SKT</td>
<td>Spring Kodiak Trawl Survey</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<td>--------------</td>
<td>-----------</td>
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<tr>
<td>SFPF</td>
<td>Skinner Fish Protection Facility</td>
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<tr>
<td>SRWTP</td>
<td>Sacramento Regional Water Treatment Plant</td>
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<tr>
<td>SSC</td>
<td>suspended sediment concentration</td>
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<tr>
<td>SWP</td>
<td>State Water Project</td>
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<td>TFCF</td>
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An updated conceptual model of Delta Smelt biology: our evolving understanding of an estuarine fish

By Management, Analysis, and Synthesis Team

Executive Summary

The main purpose of this report is to provide an up-to-date assessment and conceptual model of factors affecting Delta Smelt (Hypomesus transpacificus) throughout its primarily annual life cycle and to demonstrate how this conceptual model can be used for scientific and management purposes. The Delta Smelt is a small estuarine fish that only occurs in the San Francisco Estuary. Once abundant, it is now rare and has been protected under the federal and California Endangered Species Acts since 1993. The Delta Smelt listing was related to a step decline in the early 1980s; however, population abundance decreased even further with the onset of the “pelagic organism decline” (POD) around 2002. A substantial, albeit short-lived, increase in abundance of all life stages in 2011 showed that the Delta Smelt population can still rebound when conditions are favorable for spawning, growth, and survival. In this report, we update previous conceptual models for Delta Smelt to reflect new data and information since the release of the last synthesis report about the POD by the Interagency Ecological Program for the San Francisco Estuary (IEP) in 2010. Specific objectives include:

1. Provide decision makers with a practical tool for evaluating difficult trade-offs associated with management and policy decisions.

2. Provide scientists with a framework from which they can formulate and evaluate hypotheses using qualitative or quantitative models.

3. Provide the general public with a new way of learning about Delta Smelt and their habitat.

Our updated conceptual model describes the habitat conditions and ecosystem drivers affecting each Delta Smelt life stage, across seasons and how the seasonal effects contribute to the annual success of the species. The conceptual model consists of two nested and linked levels of increasing specificity. The general life cycle conceptual model for four Delta Smelt life stages (adults, eggs and larvae, juveniles, and subadults) includes stationary ecosystem components and dynamic environmental drivers, habitat attributes, and Delta Smelt responses. The more detailed life stage transition conceptual models for each of the four Delta Smelt life stages describe relationships between environmental drivers, key habitat attributes, and the responses of Delta Smelt to habitat attributes as they transition from one life stage to the next.

Our analyses and conceptual model show that good larval recruitment is essential for setting the stage for a strong year class; however, increased growth and survival through subsequent life stages are also needed to achieve and sustain higher population abundance. We used our conceptual model to generate 16 hypotheses about the factors that may have contributed to the 2011 increase in Delta Smelt relative abundance. We then evaluated these hypotheses by comparing habitat conditions and Delta Smelt responses in the wet year 2011 to those in the
prior wet year 2006 and in the drier years 2005 and 2010. Larval recruitment was similarly high in both wet years and lower in the drier antecedent years, but juvenile and adult abundance increased only in 2011. In 2005 and 2006, the population was limited by very poor survival from the larval to the juvenile life stage. We found that in 2011, Delta Smelt may have benefitted from a combination of favorable habitat conditions throughout the year, including:

1. Adults and larvae benefitted from prolonged cool spring water temperatures, high 2011 winter and spring outflows which reduced entrainment risk and possibly improved other habitat conditions, and possibly enhanced food availability in late spring.

2. Juveniles benefitted from cool water temperatures in late spring and early summer as well as from improved food availability and low levels of harmful *Microcystis*.

3. Subadults also benefitted from improved food availability and from favorable habitat conditions in the large, low salinity zone (salinity 1-6) located more toward Suisun Bay in 2005-2006 and 2010.

Our comparisons of other habitat attributes either produced inconclusive results or were limited by a lack of suitable data or other necessary information. This was especially true for predation risk and toxicity, and other contaminant effects. Clearly more monitoring and studies are needed on these two topics, but we also found many other data and information gaps. Overall, we did not entirely reject any of our hypotheses. Together with the large amount of published information used to construct our conceptual model, this gives us some confidence that the majority of the elements and linkages of our conceptual model are relevant and (qualitatively) correct. However, the mechanisms they describe are likely variable in the degree to which they drive population outcomes, depending on the conditions in any given year and prior Delta Smelt abundance levels. In addition, the scientific merit of some linkages for which data are sparse (e.g., predation and contaminants effects) is impossible to evaluate without additional information.

Importantly, while this report identifies many data and information gaps that must be filled before some hypotheses can be objectively evaluated, the report includes a very large amount of pertinent data and information that is currently available. The San Francisco Estuary is clearly an intensely monitored and studied ecosystem and Delta Smelt may well be one of the most thoroughly studied endangered fish species in the world. The most critical data for this report came from four long-term Interagency Ecological Program fish monitoring surveys. These surveys provide sound, high-quality data about the annual distribution and relative abundance of Delta Smelt for time periods ranging from one to more than five decades. These four surveys, other monitoring surveys, and numerous research studies provide data about many habitat attributes and ecosystem drivers.

The report ends with key conclusions, a discussion of our hypothesis testing approach, and recommendations for future work and adaptive management applications. The final report Chapter contains many concrete examples of studies, modeling approaches, and management applications that are directly derived from the conceptual model. These examples are not meant to be exhaustive lists. Rather, they are primarily intended to illustrate science and management applications of our conceptual model.

We strongly recommend that analysis, synthesis and modeling efforts, such as this report, be a high priority for the management and science organizations that oversee monitoring and research in the estuary. Without these types of integrative efforts, ongoing and proposed adaptive
management processes must conduct such efforts in an *ad hoc* manner, often driven by unrealistic schedules that are unlikely to be fulfilled. Such adaptive management processes in the estuary include the ongoing adaptive management of fall outflow for Delta Smelt, the new “Collaborative Science and Adaptive Management Program,” the California Delta Stewardship Council’s Delta Plan, and the multi-agency Bay Delta Conservation Plan. On a more basic level, such synthesis efforts identify data gaps that serve to focus research and management efforts on scientifically relevant topics rather than the “crisis of the day.”

The 2011 increase in the Delta Smelt abundance index demonstrated that the species still has the ability to rebound to higher abundance levels. Delta Smelt has often been called an indicator – or canary in the coalmine – for overall ecosystem conditions in the estuary. The 2011 increase suggests that the system has not yet irreversibly shifted into an altered state that will no longer support native species. Given the profound habitat alterations in the San Francisco Estuary, continued study of the environmental drivers and habitat attributes and the subsequent responses of the Delta Smelt population seem critical to the wise management of the species. Some possible topics for future synthesis groups include:

1. Reviews and updates to existing conceptual and mathematical models.

2. Further development of mathematical models of Delta Smelt population abundance drawn specifically from the conceptual models described in this report; applications and extensions of recently published models to help make management decisions and guide new modeling efforts; additional modeling efforts and future research projects to improve resolution and understanding of the particular factors identified as critical to reproduction, recruitment, survival, and growth.

3. Review and refinement of new models such as the emerging comprehensive state-space population model (K. Newman, U.S. Fish and Wildlife Service, personal communication); development of additional models or modules of models specifically aimed at estimating effects of inadequately monitored or difficult to measure and evaluate habitat attributes such as predation risk and toxicity; development of new “nested” and/or “linked” mathematical modeling approaches that can accommodate multiple drivers and their interactive effects across temporal and spatial scales.

4. Interdisciplinary collaboration among scientists, managers, and stakeholders to develop and model management scenarios and strategies based on principles of integrative ecosystem and landscape-based management rather than relatively crude distinctions among categorical “water year types.”

Continued growth of California’s human population, climate change, new species invasions, and other changes will increase management challenges. Science and management have to go hand in hand to constantly identify, implement, evaluate, and refine the best management options for this ever-changing system. We hope that the conceptual model and information in this report will be useful for achieving these goals.
Chapter 1: Introduction

The San Francisco Estuary

Estuarine ecosystems are among the most complex ecosystems on earth (Wilson 1998). They are constantly changing ecosystems that respond to dynamic “drivers” of change (Healey et al. 2008, Baxter et al. 2010). Natural drivers include the geological and geographic setting, climatic and oceanic variability, dynamic hydrological and nutrient regimes, weather and disturbance regimes, biogeochemical processes, species assemblages, and many other biotic and abiotic features. Estuaries also respond to a broad range of human activities. Some of these “human drivers” have negative impacts on ecosystems. These negative human drivers are often called “stressors.” Human stressors on estuarine ecosystems include water and land use, pollutant discharges, species introductions, and fishing (Townend 2004, Lotze et al. 2006, Cloern and Jassby 2012). The interplay of natural and human drivers and their effects on the San Francisco Estuary and in particular on the Delta Smelt (*Hypomesus transpacificus*), an endemic fish species, is the subject of this report.

The San Francisco Estuary (SFE; Fig. 1) is comprised of an upstream region consisting of channels and islands associated with the confluence of the Sacramento and San Joaquin Rivers known as the “Delta” and a series of downstream bays and marshes that are separated from the Pacific Ocean by the “Golden Gate,” the sea passage between the San Francisco and Marin peninsulas. Because of California’s Mediterranean climate, the SFE experiences large interannual and seasonal flow variations, which are modulated by tides and human management of the rivers within the Delta watershed (Moyle et al. 2010). These hydrological variations lead to a dynamic estuarine salinity gradient. In the winter and spring fresh water often extends into San Pablo Bay, while in the summer and fall brackish water can intrude into the western Delta. These seasonal differences are exacerbated by pronounced interannual differences in precipitation in the watershed. Extremely dry years with little precipitation and very wet years with widespread flooding do not occur in predictable patterns (Dettinger 2011).

The SFE has undergone dramatic morphological, hydrological, chemical, and biological alterations since the onset of the California Gold Rush in the middle of the 19th century (Nichols et al. 1986, Arthur et al. 1996, Baxter et al. 2010, Brooks et al. 2012, NRC 2012, Whipple et al. 2012, Cloern and Jassby 2012). These alterations include five human activities that have changed ecological functions and habitats in many riverine and estuarine systems with increasingly dense human populations: diking, draining, dredging, diverting, and discharging. Specifically, diking and dredging have reduced the vast wetlands that once covered and surrounded the SFE to small remnants. There has been an 80-fold decrease in the ratio of wetland to open water area in the Delta, from a historical ratio of 14:1 to a current ratio of 1:6 (Whipple et al. 2012, Herbold et al. 2014). Diking and dredging have led to a substantial reconfiguration of the bays, sloughs, and channels, while large-scale water diversions, and discharge of contaminants have altered water quantity and quality. Small water diversions occur throughout the freshwater portion of the estuary, but the largest water diversions are at the pumping facilities of the federal Central Valley Project (CVP) and the State Water Project (SWP) that export water from the southwestern Delta to agricultural and urban areas to the south (Fig. 2). In addition, a wide variety of non-native plants and animals have been introduced and have become established in the SFE (Cohen and Carlton 1998, Light et al. 2005, Winder et al. 2011).
Figure 1. Map of the San Francisco estuary. The inset shows various values of X2, the distance in kilometers from the Golden Gate to the near bottom salinity 2 isohaline.
Figure 2. Map of the upper San Francisco estuary. The upper estuary includes the Suisun Bay region and the Sacramento-San Joaquin Delta, which are west and east of Chipps Island respectively. The area from approximately Chipps Island to the west end of Sherman Island is referred to as the “confluence.”
Many of the more recent ecological changes in the SFE have been documented by long-term monitoring surveys. Most of these surveys are conducted under the auspices of the Interagency Ecological Program (IEP), an interagency science consortium with three State and six federal member agencies (http://www.water.ca.gov/iep/). Together with monitoring conducted by others, these monitoring surveys provide one of the longest and most comprehensive environmental and biological data records in a U.S. coastal ecosystem. With each additional year of monitoring, this data record serves as an increasingly valuable tool for observing gradual changes or abrupt shifts in ecological conditions and for identifying their underlying causes (Cloern and Jassby 2012).

The modern SFE continues to be a dynamic and complex ecosystem that supports many important ecosystem services (Millennium Ecosystem Assessment 2005), including the provision of fresh water, agricultural crops, commercial and recreational fisheries, and other recreational opportunities. However, it no longer provides adequate habitat for many of its native species as evidenced by severe declines in several of its native fish populations (e.g., Bennett and Moyle 1996, Brown and Moyle 2005, Sommer et al. 2007).

**Pelagic fish declines**

Among the native fishes of the upper SFE (Fig. 2), the endemic Delta Smelt is of high management concern because of a decline of its annual abundance indices (see Chapter 3 for details of fish surveys and indices), particularly longer term indices for juveniles and subadults, to persistent low levels (Fig. 3). This decline led to its listing under the federal Endangered Species Act in 1993. The Delta Smelt is a slender-bodied pelagic fish with a maximum size of about 120 mm standard length (length from snout to end of vertebral column) and a maximum age of two years. It is the most estuary-dependent of the native fish species in the SFE (Moyle et al. 1992, Bennett 2005). The continued existence of the species is dependent upon its ability to successfully grow, develop, and survive in the SFE.

Delta Smelt is not the only fish species currently in decline in the Delta. Abundance indices of Longfin Smelt (*Spirinchus thaleichthys*), age-0 Striped Bass (*Morone saxatilis*), and Threadfin Shad (*Dorosoma petenense*) declined simultaneously with those of Delta Smelt in about 2002. This simultaneous decline has become known as the pelagic organism decline (POD) (Sommer et al. 2007, Baxter et al. 2008, 2010) (Fig. 4). Given the very different life histories of these four pelagic species, it is unlikely that a single environmental variable could account for the POD declines. In general, researchers have suggested that the POD declines were likely multi-causal (Sommer et al. 2007, Baxter et al. 2008, 2010, Mac Nally et al. 2010, Cloern and Jassby 2012). Several researchers have suggested that the SFE has undergone an ecological regime shift (Moyle and Bennett 2008, Baxter et al. 2010, Glibert et al. 2011, Cloern and Jassby 2012). In the present system, an invasive aquatic macrophyte (*Egeria densa*) dominates the littoral zone of many areas of the Delta and provides favorable habitat for many invasive fishes (e.g., Largemouth Bass *Micropterus salmoides*; Brown and Michniuk 2007); invasive clams (*Potamocorbula amurensis* and *Corbicula fluminea*) consume a large portion of the available pelagic phytoplankton (Alpine and Cloern 1992, Lopez et al. 2006, Lucas et al. 2002, Lucas and Thompson 2012); agricultural, industrial, and urban discharges transport large quantities of nutrients and a plethora of contaminants into many regions of the estuary; and current management of water for agricultural, industrial and urban purposes is focused on optimizing the reliability of water exports by the CVP and SWP.
Figure 3. Delta Smelt abundance index for life stages of Delta Smelt including the larvae-juveniles (20 mm Survey), juveniles (Summer Townet Survey), subadults (Fall Midwater Trawl), and adults (Spring Kodiak Trawl). The initiation of each individual survey is indicated by the initial bar with subsequent missing bars indicating when an index could not be calculated. See Chapter 3 for details of sampling programs, including geographic coverage, and Appendix B for details of calculating abundance indices.
Figure 4. Abundance indices from Fall Midwater Trawl for Delta Smelt, Longfin Smelt, age-0 Striped Bass, and Threadfin Shad. Missing bars indicate when an index could not be calculated. See Chapter 3 for details of sampling programs, including geographic coverage, and Appendix B for details of calculation indices of abundance indices.
Changes in Delta Smelt distribution and abundance

Long-term monitoring surveys conducted by the IEP have documented substantial changes in the distribution and abundance of Delta Smelt in its small native geographic range which extends from the upstream boundaries of tidal influence in the northern, eastern and southern Delta region of the estuary to Suisun and San Pablo Bays in the north-western region of the estuary. The geographic range of Delta Smelt also includes some of the larger tidal sloughs and tributaries adjacent to Suisun and San Pablo Bays, including some Suisun Marsh sloughs and the lower Napa River (Bennett 2005, Hobbs et al. 2007, Sommer et al. 2011, Merz et al. 2011, Sommer and Mejia 2013, Murphy and Hamilton 2013). Delta Smelt are generally considered a pelagic species. While they are commonly found in shallow shoal areas such as Honker and Grizzly Bays in the Suisun Bay region of the estuary and larger marsh sloughs such as Suisun and Montezuma Sloughs in Suisun Marsh and the lower reaches of Cache and Lindsey Sloughs in the northern Delta, they are less commonly encountered in near-shore areas and only rarely in smaller marsh sloughs (Bennett 2005, Merz et al. 2011, Sommer and Mejia 2013).

The Delta Smelt has been characterized as a “semi-anadromous” fish species that spawns in fresh water and rears in fresh to brackish water (Fig. 5; Dege and Brown 2004, Bennett 2005, Sommer et al. 2011, Merz et al. 2011). While Delta Smelt have been documented throughout their geographic range during most months of the year (Sommer et al. 2011, Merz et al. 2011, Murphy and Hamilton 2013), their distribution varies seasonally in response to dynamic abiotic and biotic habitat attributes such as salinity, temperature, turbidity, and presumably food supplies (Bennett et al. 2005, Sommer et al. 2013, Brown et al. 2014). In years with high freshwater discharge in winter and spring, spawning and rearing of larval and early post-larval fish can temporarily extend seaward into San Pablo Bay, while in years with less discharge it usually occurs in the Delta, Suisun Bay and Suisun Marsh. Juveniles and adults are distributed across a broader salinity range (0 to about 18) than larval and post-larval fishes which tend to be most abundant in the low salinity zone (salinity 1-6). Dege and Brown (2004) and Sommer et al. (2011) found that the center of the Delta Smelt distribution is associated with salinities of about 2 during most months and moves with the estuarine salinity gradient as the salinity gradient responds to flow.

Historically, Delta Smelt were commonly observed throughout the fresh and low salinity portions of their geographic range (Erkkila et al. 1950, Radke 1966). Over the last two decades, their geographic distribution has become more constricted during the summer and fall. At present, Delta Smelt are less commonly found in the southern and eastern Delta during the winter and spring and are largely absent from this region in the summer and fall (Nobriga et al. 2008, Sommer et al. 2011). While Delta Smelt continue to be found in the northern Delta year-round and individual catches in this region are sometimes large, particularly during winter and spring, the majority of the population is usually observed in the region near to and west of the Sacramento-San Joaquin River confluence, especially in the summer and fall (Sweetnam 1999, Feyrer et al. 2007, Nobriga et al. 2008, Merz et al. 2011, Sommer et al. 2011, Sommer and Mejia 2013).

In addition to documenting changes in distribution, long-term IEP surveys also reveal that the annual abundance indices of Delta Smelt have greatly declined since the first long-term pelagic fish monitoring survey began in summer 1959 (Fig. 3). Both a gradual, long-term decline and step changes, most recently around 2002, have been described using a variety of qualitative and statistical approaches for subadult Delta Smelt caught in the fall (e.g., Bennett and Moyle 1996, Bennett 2005, Manly and Chotkowski 2006, Thomson et al. 2010). These declines have not been smooth or entirely unidirectional and also include a great deal of interannual variability (Fig. 3).
Since the beginning of the POD in 2002, the Delta Smelt abundance indices have often been at record low levels, leading to concerns about declines in effective population size (Fisch et al. 2011) and a loss of population-level resilience, meaning the ability of the population to recover to higher population abundances when conditions are suitable. For example, population sizes might become too small to produce enough eggs or larvae to outpace predation on eggs and larvae.

Delta Smelt previously rebounded from low population abundances, most recently in the wet years of the late 1990s (Fig. 3). The lack of increase in Delta Smelt in the wet year of 2006 combined with new evidence for genetic bottlenecks and a significant decline in effective population size from 2003 to 2007 (Fisch et al. 2011) were thus a source of great concern. However, during 2011, the next wet year after 2006, the species did increase in abundance (Fig. 3). Unfortunately, the increase in Delta Smelt abundance was short-lived and did not carry over into the following year-class in 2012, a drier year. Nevertheless, the temporary increase gave some cause for renewed optimism about the resilience of the species and its potential recovery. In addition, the contrasts between habitat conditions and Delta Smelt responses in 2006 and 2011 provided an opportunity to gain new insights into the Delta Smelt habitat requirements that might help better manage this species and its habitat.

**Protecting Delta Smelt**

Delta Smelt are currently protected under both California and federal endangered species legislation. The protection and recovery of Delta Smelt and its estuarine habitat in the SFE will
likely require the human population of California to reduce its dependence on some of the natural resources provided by the SFE. This will become even more challenging in the future because of climate change and the continued growth of California’s human population. California’s population has increased by approximately 38 million people compared to the population when California became a state in 1850 and has increased by about 22.5 million compared to 1959 when Delta Smelt monitoring started 55 years ago (U.S. Census Bureau data). More than three quarters of today’s 38 million Californians live south of the SFE, and the majority of these Californians and millions of acres of farmland rely on fresh water diverted from the Delta for all or part of their water supply. The conflicts and trade-offs between species protection measures and actions to provide water and other natural resources to California’s growing human population have resulted in repeated attempts to reconcile these seemingly irreconcilable objectives through regulatory requirements, new institutional arrangements, and management plans.

Among the regulatory requirements are the State water right decisions issued by the California State Water Resources Control Board, which grant SWP and CVP water rights permits, but also include requirements to protect fish. State regulations also include increasingly more stringent waste discharge permits. For example, the new permit recently issued to the Sacramento Regional County Wastewater Treatment Plant includes new requirements for major treatment upgrades to better protect downstream water uses and the health of the estuary. Federal regulations include water quality requirements under the Clean Water Act and Biological Opinions (BiOps) issued under the federal Endangered Species Act. Two BiOps assess the effects of the coordinated operations of the SWP and CVP on Delta Smelt, Green Sturgeon, and salmonid fish populations, and their designated critical habitat. These BiOps include “reasonable and prudent alternatives” to lessen negative impacts of SWP and CVP operations and avoid jeopardy to the species, while at the same time trying to avoid major reductions in water exports from the Delta.

Recent institutional reconciliation attempts include the multiagency, State and federal CALFED Bay-Delta Program and Authority (CALFED) and the California Delta Stewardship Council (DSC), a new State agency. From 1994 to 2010, CALFED attempted to reconcile water allocation and ecosystem restoration efforts in the estuary in a way that would allow them to “get better together” (Doremus 2009). After the demise of CALFED, the State of California created the DSC to address what the legislature termed the “co-equal goals” of providing a more reliable water supply for California and protecting, restoring, and enhancing the Delta ecosystem (CA Water Code §85054, [http://deltacouncil.ca.gov/](http://deltacouncil.ca.gov/)).

Among the many management plans aimed at reconciling species protection and human water and land use objectives are plans by the DSC, SWRCB, and new groupings of multiple agencies and stakeholders. The DSC recently completed and is now starting to implement its comprehensive “Delta Plan” ([http://deltacouncil.ca.gov/delta-plan-0](http://deltacouncil.ca.gov/delta-plan-0)) to achieve the co-equal goals, while the SWRCB is on track to complete a major update to its “Bay-Delta Plan” which may result in changes to water right permits ([http://www.waterboards.ca.gov/waterrights/water_issues/programs/bay_delta/](http://www.waterboards.ca.gov/waterrights/water_issues/programs/bay_delta/)). Three California State agencies recently completed a new California Water Action Plan that includes actions to help achieve the co-equal goals ([http://resources.ca.gov/california_water_action_plan/](http://resources.ca.gov/california_water_action_plan/)). A multi-agency planning effort that includes State and federal agencies as well as local Public Water Agencies (water contractors) is working to complete the “Bay-Delta Conservation Plan” (BDCP, [http://baydeltaconservationplan.com](http://baydeltaconservationplan.com)). The BDCP is a proposed Habitat Conservation Plan under the federal Endangered Species Act and a Natural Community Conservation Plan under the California Natural Community Conservation Planning Act. It proposes to implement habitat restoration measures, stressor reduction activities,
improved water project operations criteria, and new water conveyance infrastructure. If approved by the regulatory agencies, this plan would provide long-term permits for the various projects and water operations to proceed over a 50-year time frame.

Management actions, regulatory requirements, and institutional arrangements in the SFE have undergone substantial and complex changes over the last 150 years. Hanak et al. (2011) describe a progression from an early disorganized “laissez-faire” era of California and SFE water management followed by increasingly organized and large-scale management schemes, from local water use to state-wide water projects, which led to a current “era of conflict” and the hope for a new “era of reconciliation.” A complete review of these changes is outside the scope of this report and the reader is referred to Hanak et al. (2011) and other existing reports on this topic. It is important to note, however, that increasingly, these changes have been “adaptations” based on the results of monitoring, studies, and other scientific activities in the SFE. Many of these scientific activities have been conducted under the auspices of the IEP (Herrgesell 2013). It can be argued that some of the activities preceding and ultimately leading to the creation of the IEP in 1970 ushered in an era of increasingly intense and formalized “adaptive management” before the term itself was coined.

Adaptive management is a formal approach to natural resource management that closely connects science with management to devise, track, and improve management outcomes. This connection started to become an important aspect of fisheries management in the 1950s (e.g., Beverton and Holt 1957), although the term itself was not coined until 1978 when Holling (1978) and Walters and Hilborn (1978) provided a conceptual framework for adaptive resources management. This framework was later refined to distinguish between “passive” and “active” adaptive management. According to Williams (2011), “active adaptive management actively pursues the reduction of uncertainty through management interventions, whereas passive adaptive management focuses on resource objectives, with learning a useful but unintended byproduct of decision making […]. In practice this means that a key difference between passive and active adaptive management is the degree to which the objectives that guide decision making emphasize the reduction of uncertainty.” In active adaptive management, management actions are designed as “experimental treatments” with clear hypotheses about outcomes that are tested through rigorous data collection and analyses. This accelerates learning, but can come at the expense of achieving resource objectives because potentially less effective management actions may be included in the experimental design. Moreover, the more intense science efforts needed for active adaptive management can be costly over the short term (Williams 2011). This may explain why passive adaptive management, while not always referred to by this name or implemented in the formal and rigorous way now advocated by the DSC’s Delta Plan (DSC 2013), has been and continues to be common in the SFE, but active adaptive management – viewed by some as the only “real” adaptive management – is still rare.

Of all current management actions and requirements affecting Delta Smelt, the actions required in the 2005 and 2008 BiOps issued by the U.S. Fish and Wildlife Service (FWS) are most directly aimed at the protection of Delta Smelt. The 2008 BiOp takes a life cycle approach to protecting Delta Smelt and includes an explicit requirement for adaptive management of fall outflow. After initial steps to design a passive adaptive management program, the U.S. Bureau of Reclamation (Reclamation) decided to take a more active approach aimed at more rapidly reducing uncertainties about the underlying mechanisms and effects of fall outflow management on Delta Smelt (Reclamation 2011, 2012, Brown et al. 2014). The study component of the fall outflow adaptive management plan, also known as the “fall low salinity habitat” (FLaSH) studies, was developed with the help of a new conceptual model (FLaSH conceptual model, Brown et
al. 2014) and has been implemented by the IEP starting in 2011. The FLaSH studies provided an opportunity to intensely study the increase in the Delta Smelt abundance index observed in 2011. At this initial stage of the adaptive management program and the FLaSH studies, the 2011 data were compared to data gathered in the previous wet year, 2006, during which fall outflow was lower. The initial data analysis effort also considered antecedent conditions in 2010 and 2005, resulting in a simple comparative approach focusing on four years (Brown et al. 2014).

**Report Purpose and Organization**

It is clear that the recovery of Delta Smelt and other listed and unlisted native species will be a key requirement of any plan to manage the resources of the SFE. Understanding the factors driving Delta Smelt population dynamics is a major goal of resource management agencies. The main purpose of this report is to provide an up to date assessment of factors affecting Delta Smelt throughout its primarily annual life cycle. Specific goals are to provide decision makers with scientific information for evaluating difficult trade-offs associated with management and policy decisions, provide scientists with a resource for formulating and testing hypotheses and mathematical models, and provide the general public with a new way for learning about Delta Smelt and their habitat.

We address these goals through a synthesis of scientific information about Delta Smelt with an emphasis on new information since the release of the last POD synthesis report in 2010 (Baxter et al. 2010). As in previous reports, conceptual models play a key role in this report. Conceptual models are useful tools for organizing and synthesizing information, designing research and modeling studies, and for evaluating potential outcomes of management actions. Here, we revisit previously developed conceptual models for Delta Smelt, and synthesize new information about factors affecting Delta Smelt and Delta Smelt responses to those factors. This comprehensive body of information is then used to construct and populate a Delta Smelt conceptual model, within a new framework.

Numerous conceptual models have been developed to describe the relationships and linkages among environmental drivers of ecosystem change, ecosystem and habitat attributes, and Delta Smelt responses. In Chapter 2 of this report, we provide a brief introduction to conceptual models and review some of the conceptual models developed for the SFE and for Delta Smelt. In Chapter 3, we introduce a new conceptual model framework for Delta Smelt and describe our approach to updating the previously developed Delta Smelt conceptual models. We also describe the data sources and analytical approaches used in this report. In Chapter 4, we review and synthesize recent information about drivers and habitat attributes affecting Delta Smelt and Delta Smelt responses to habitat attributes. In Chapter 5, we present an updated conceptual model for Delta Smelt that include key drivers, habitat attributes, interactions between them, and Delta Smelt responses discussed in Chapter 4. In Chapter 6, we review and synthesize recent information about Delta Smelt population dynamics, life history, and population trends. In Chapter 7, we use the updated conceptual model to formulate hypotheses about Delta Smelt responses and changing habitat conditions and test them using a simple comparative approach similar to the FLaSH approach (Brown et al. 2014), but for all life stages of Delta Smelt. The purpose of Chapter 7 is to put the new conceptual model along with the comparative approach to an immediate test that is of high relevance to the management of Delta Smelt. Chapter 8 presents key results and conclusions from the preceding Chapters. In Chapter 9, we discuss next steps for future conceptual, qualitative, and quantitative modeling as well as the science and management implications of the information contained in this report.
Chapter 2: Conceptual Models

Overview

We learn and think about the world we live in through mental models of how the world looks and how it works. Our mental models guide all our conscious decisions and actions. They are never static; we constantly update them with new information gained by observing the world around us and by assessing the outcomes of our decisions and actions. In our minds, we compare the new information against our existing mental models. Observations that agree with our mental models strengthen them, observations that don’t agree with our mental models force us to modify, adjust, and update them.

Conceptual models are formalized versions of mental models that are communicated to others verbally and graphically. Ecologists and environmental managers use them to communicate hypotheses about “how ecosystems work” and to explore how human actions and other drivers change ecosystems. They usually use a combination of narrative text and graphical illustrations about ecosystem components and the relationships among them. More informal narrative conceptual models verbally describe cause-effect relationships, while more formal conceptual models may express them through scientific hypotheses or mathematical equations.

Conceptual model illustrations often take the form of pictures, plots, schematic images or diagrams, matrices, or tables (Fischenich 2008). For example, the IEP Estuarine Ecology Team used elaborate matrices to illustrate and assess the likely mechanisms underlying the statistically determined relationships between SFE fishes and “X2,” an indicator of estuarine salinity dynamics (Estuarine Ecology Team 1997), while Reclamation (2011, 2012) used a table format to illustrate how fall outflow interacts with other features of Delta Smelt habitat and affects Delta Smelt. Schoellhamer et al. (2012) used a series of conceptual X-Y plots to illustrate a conceptual model of sediment supply reduction and downstream propagation in the SFE. Gibert (2012) and Glibert et al. (2011) used schematic images to conceptualize changes in nutrients, flows, biogeochemical processes, and the food web of the SFE. Many schematic conceptual model diagrams use boxes to depict ecosystem components and arrows to illustrate the relationships, flows, and interactions among them. The conceptual models developed by the IEP for its POD investigations (see below) include examples of schematic conceptual model depictions with few boxes and arrows, while some of the conceptual models developed for the “Delta Regional Ecosystem Restoration Implementation Plan” (DiGennaro et al. 2012, see below) and the “effects hierarchy” of factors affecting Delta Smelt abundance developed by Miller et al. (2012) provide examples of more complex schematics with a large number of boxes and arrows.

Conceptual models have become essential tools for summarizing, synthesizing, and communicating scientific understanding of ecosystem structure and functioning. They are also key to successful planning and implementation of ecological research and mathematical modeling as well as to adaptive management, restoration and recovery of ecosystems, and environmental science education (e.g., Thom 2000, Ogden et al. 2005, Fortuin et al. 2011). Conceptual models are also essential tools for identifying management and science priorities and for the selection of key ecological attributes to be used to evaluate the performance of management actions (i.e., performance measures) and assess the present relative to a desired state of an ecosystem (i.e., indicators) (Washington State Academy of Sciences 2012).
Conceptual models have clear limitations. For example, even the most complex conceptual models are highly simplified descriptions of a small part of an ecosystem – they can never tell the “whole” story. Just like our every-day mental models, they are also never final. To remain relevant, ecological conceptual models must evolve and change with the evolution of our knowledge about ecosystems. Furthermore, conceptual models identify key ecosystem components and relationships, but they do not quantify them and unless they are coupled with mathematical models, conceptual models cannot be used to make quantitative predictions.

Conceptual models can be used to make qualitative predictions about changes in ecosystem components and their relationships. These qualitative predictions can serve as testable hypotheses that help design scientific analyses and studies. The creation or revision of the conceptual models themselves usually forces the formulation of hypotheses and their testing with available data and information, as will be demonstrated in the later Chapters of this report. Qualitative predictions and testable hypotheses are also at the heart of active adaptive management. They are needed to design experimental adaptive management actions and the studies and monitoring needed to assess the outcomes from such actions. The fall outflow adaptive management plan (Reclamation 2011, 2012) provides an example of how a conceptual model was used to make qualitative predictions and design a comprehensive set of studies, the FLaSH studies. Finally, the formulation of conceptual models is usually the essential first step for constructing quantitative models. Mathematical models are sets of mathematical expressions that quantify the components and relationships in the conceptual models and can be used to make quantitative predictions about the state of ecosystem components and linkages under specific circumstances (Jackson et al. 2000). The (few) quantitative predictions in the fall outflow adaptive management plan (Reclamation 2011, 2012) are based on such mathematical models.

Ecological conceptual models generally link ecological “drivers” with ecological effects or “outcomes.” Drivers are physical, chemical, or biological factors of human or natural origin (for example, nutrients from natural soils and applied fertilizers). Outcomes can be physical, chemical or biological responses to the drivers (for example, phytoplankton growth and biomass), but can also be social and economic impacts on human components of the ecosystem (for example, harmful algal blooms that affect recreational use or costs of water treatment for drinking water supply). Drivers and outcomes are the components of the system under consideration. They are linked by mechanistic cause-effect relationships. Conceptual models can also be nested within each other, for example, to accommodate different temporal or spatial scales, or conceptual models can be coupled so that the outcome of one conceptual model becomes a driver in the next one. Drivers are often categorized in various ways, including their causal proximity to specific outcomes, whether they are natural or anthropogenic, and whether they can be altered by human management strategies and actions. Graphically, drivers are often arranged in hierarchical tiers that reflect these categories.

For example, Gentile et al (2001) describe a basic three-tiered approach that links environmental outcomes (tier 1) to proximal anthropogenic drivers termed “stressors” (tier 2) and the natural and anthropogenic drivers that act on these stressors (tier 3). Davis et al. (2010) show how different ecological regimes in Australian lakes (outcomes, tier 1) arise from the interplay of stressors (tier 2) and hydrological changes (tier 3) acting on the original ecological regime (tier 4). Carr et al. (2007) review a widely used five-tiered “Driver–Pressure–State–Impact–Response” (DPSIR) framework that focuses on identifying human-caused environmental problems and solutions. In this framework, the ultimate drivers (D) are social processes that result in specific human activities that manifest as proximal “pressures” (P) that change the “state” (S), or condition, of the environment. This can have “impacts” (I) on human well-being that are recognized as
problems. Some impacts are so severe that they require a human response (R), usually in the form of institutional solutions aimed at reducing high-priority impacts. The Puget Sound Partnership Science Panel (2012) recently used the DPSIR framework to develop a conceptual model that links management strategies (i.e., responses; e.g., reduce pollution) to anthropogenic drivers (e.g., human population growth) and pressures (e.g., pollution) that affect the state of ecosystem components (e.g., habitats and species) and impact the provisioning of ecosystem services (e.g., fishing). This model helped identify scientific knowledge gaps and decision-critical issues and questions that needed to be answered in response to management priorities.

Recent Conceptual Models for the San Francisco Estuary

Over the last decade, two integrated sets of conceptual models have been developed for portions of the SFE. The first conceptual model set was developed by the Ecosystem Restoration Program (http://www.dfg.ca.gov/ERP/) to evaluate restoration actions in the Delta under the “Delta Regional Ecosystem Restoration Implementation Plan” (DRERIP; DiGennaro et al. 2012). DRERIP conceptual models were developed for ecological processes, habitats, specific species, and stressors. The DRERIP conceptual models were built around environmental drivers, their expected effects termed “outcomes,” and cause-and-effect relationships between the two shown as one-way arrows termed “linkages.” In the graphical depiction of the DRERIP conceptual models, different arrow widths, colors, and styles denote the importance, degree of understanding, and predictability, respectively, of the driver-linkage-outcome relationships, while symbols next to the arrows denote the direction and nature of the effect (positive, negative, or non-linear) (DiGennaro 2012, Opperman 2012). The DRERIP species conceptual models include “transition matrix” diagrams depicting how environmental drivers affect the probability of one life stage successfully transitioning to the next.

The second set of conceptual models was developed by the IEP as a comprehensive conceptual framework intended to guide investigations of the POD and to synthesize and communicate results (Sommer et al. 2007, Baxter et al. 2010). This framework includes a “basic” POD conceptual model about key drivers of change affecting pelagic fish and their habitat (Fig. 6), more narrowly focused “species-specific” conceptual models about drivers affecting the different life stages of each of the four POD fish species (e.g., Fig. 7), and a broader “ecological regime shift” conceptual model that placed the POD decline in a longer-term historical context (not shown; see Baxter et al. 2010). The basic POD conceptual model placed the four fish species in the center of interacting drivers affecting the quantity and quality of their habitat (Fig. 6), while the species-specific models identified key seasonal drivers in red, with proximal causes and effects in yellow (Fig. 7).

The National Research Council Committee on Sustainable Water and Environmental Management in the California Bay-Delta (NRC Committee) (NRC 2012) called the POD conceptual model framework “an important example of supporting science. This framework identifies and links, in the context of both ecosystem structure and functioning, the key stressors that help to explain the decline of pelagic organisms.” The NRC Committee further noted that the “drivers of change” identified in the POD conceptual models “are quantifiable” and “suitable for model evaluation” and that the:
“types of stressors identified are integrative, reflecting co-occurring physical, chemical, and biotic changes. They also apply to multiple structural (food web structure, biodiversity) and functional (food transfer changes, biogeochemical cycling) changes taking place in the Delta. The framework and associated detail are both comprehensive and useful in terms of linking these drivers to changes taking place at multiple levels of the food web. This type of conceptual approach will also be useful for examining other drivers and impacts of ecological change, including observed changes in fish community structure and production; specifically, how these changes are affected and influenced by changes in physico-chemical factors (e.g., salinity, temperature, turbidity, nutrients/contaminants) and at lower trophic levels (phytoplankton, invertebrate grazers, and prey)” (NRC 2012, p. 34-35).

Since the release of the 2012 NRC report, the POD conceptual model framework has been used as the basis for additional conceptual models developed to aid planning and quantifying the ecological effects of active adaptive management of Delta outflow to improve fall low salinity habitat for Delta Smelt and to guide the associated fall low salinity habitat (FLaSH) studies (Reclamation 2011, 2012). A more complete summary of the POD and FLaSH conceptual models along with additional information about related conceptual and quantitative models in the SFE can be found in the initial FLaSH report (Brown et al. 2014, see also http://deltacouncil.ca.gov/science-program/fall-low-salinity-habitat-flash-studies-and-adaptive-management-plan-review-0).
One important new feature of the conceptual model developed for the fall outflow adaptive management plan and the FLaSH studies was the explicit consideration of interacting dynamic and relatively more stationary (geographically and temporally fixed) habitat components that was based on a conceptual model of environment-habitat-production linkages in tidal river estuaries developed by Peterson (2003). In the FLaSH conceptual model, the interactions among dynamic and stationary habitat components determine the characteristics of Delta Smelt habitat in the fall and lead to varying Delta Smelt outcomes. In essence, the dynamic flow and salinity regimes of the SFE move water, particles, and organisms across the estuary’s stationary topography, which has distinct physical features that modulate the dynamic habitat components. Together, these stationary and dynamic habitat components are hypothesized to control the survival, health, growth, fecundity, and, ultimately, the reproductive success of estuarine pelagic species, such as Delta Smelt. The interplay between stationary and dynamic habitat components also helps explain the distribution and movement of Delta Smelt across its range which cannot be understood – or managed – based on geography alone.

Numerous other conceptual and quantitative models have been developed for the SFE. Kimmerer (2004) summarized many of the earlier conceptual models. More recent conceptual model examples include those by Gilbert (2012) and Gilbert et al. (2011) as well as the five-tiered effects hierarchy by Miller et al. (2012). Recent examples of mathematical models of habitat use and population dynamics of Delta Smelt include models based on statistical approaches (e.g.,

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**Figure 7.** Species-specific conceptual model for Delta Smelt. This is one of four species-specific conceptual models developed as part of the conceptual framework for the pelagic organism decline (Baxter et al. 2010). The low salinity zone (LSZ) is defined as salinity 1-6. The Vernalis Adaptive Management Plan (VAMP) included reductions in spring exports with possible effects on Delta Smelt.
Chapter 3: Approach

This report is the result of a team effort by the IEP Management, Analysis, and Synthesis Team (MAST, often referred to as “we” in this report). Appendix A briefly describes the MAST and the report development process and schedule which included a public and independent expert peer review step that led to major revisions to the draft report.

General Approach

Our general approach in this report was to develop a new conceptual model framework for Delta Smelt and to use this framework to synthesize new scientific information and update and integrate existing conceptual models including the “basic” and “species-specific” POD conceptual models, the DRERIP “transition matrix” models, the tabular FLaSH conceptual model and the hierarchical conceptual model in Miller et al. (2012) described in Chapter 2.

The development of the new conceptual model framework was guided by the conceptual model literature (see Chapter 2) and by recommendations from the independent “FLaSH Panel” of national experts convened by the Delta Science Program. The FLaSH Panel recommended to:

“develop a schematic version of the [FLaSH] conceptual model that matches the revised, written version of the conceptual model in the draft 2012 FLaSH study report. The conceptual model in written and schematic form should continue to emphasize processes and their interactions over simple correlations, should ensure Delta Smelt vital rates remain central to thinking, and should be designed for routine use by scientists as an organizational tool and for testing hypotheses associated with the AMP [adaptive management plan]; it should be as complex as necessary to achieve these purposes. The conceptual model should also be able to encompass processes and interactions that extend before and after Fall Outflow Action periods, including areas both upstream and downstream of the LSZ” (FLaSH Panel 2012, page ii).

The conceptual modeling approach in this report is intended to provide a basis, not a substitute for the development or use of mathematical models. While mathematical models are outside of the scope of this report, we briefly discuss the promise and challenges of mathematical models for Delta Smelt, summarize some of the highlights of existing mathematical modeling efforts for Delta Smelt, and offer a brief description of two additional proposed mathematical modeling efforts — one qualitative and the other quantitative — we think are natural outgrowths of the information in this report (see Chapter 8). Development of a variety of flexible working tools to facilitate discussion of elements of the conceptual model is one intended outcome of the MAST effort. Even simple quantitative and qualitative models based on our revised conceptual model
In this Chapter, we introduce the new conceptual model framework for Delta Smelt. This framework consists of a series of nested and tiered conceptual models: a general life cycle conceptual model and more detailed life stage transition conceptual models. It was developed following recommendations by the FLaSH Panel (FLaSH Panel 2012) and extensive reviews of a draft version of this report (see http://www.water.ca.gov/iep/pod/mast.cfm and Appendix A). In Chapter 4 we review and synthesize existing information about drivers, habitat attributes, and Delta Smelt responses with a focus on new information since 2010. We use the drivers in the basic POD conceptual model as the basis for this synthesis. This information is then used to populate the nested conceptual models in the new conceptual model framework with key drivers and their linkages to Delta Smelt responses. The fully populated nested conceptual models are presented in Chapter 5. Chapter 6 focuses on Delta Smelt life history and population dynamics and trends. Chapters 4 and 6 include some new analyses of long-term monitoring data, but are largely based on a review and synthesis of the existing published literature. In Chapter 7, we compare data pertaining to ecosystem drivers (drivers), habitat attributes (drivers or outcomes) and Delta Smelt responses (outcomes) in four recent years with moderate to wet hydrology: the two most recent wet years (2006 and 2011) and the two drier years immediately before them (2005 and 2010). The intent is to assess the utility of the conceptual model for formulating and testing hypotheses that expand the comparative FLaSH approach (Brown et al. 2014) that focused on the fall to a more comprehensive year-round investigation of why Delta Smelt abundance increased in the wet year of 2011, but failed to respond to wet conditions in 2006. In each of the sections in Chapter 7 covering a specific life stage, the hypotheses inherent in the conceptual model are stated and the reasoning for including each hypothesis is explained. Although we attempted to develop independent hypotheses, this was not always possible because many drivers were related and important habitat attributes were influenced by multiple drivers and their interactions, as shown in the conceptual model diagrams and explored in Chapter 4.

Key insights from Chapters 4–7 are summarized in Chapter 8. In Chapter 8, we also discuss limitations of the analytical approaches in this report. In Chapter 9, we describe additional data and analyses needed to test hypotheses that could not be conclusively tested with the available data and our simple comparative analysis approach. We also present some ongoing or possible next steps for future years, including some recommendations for future synthesis and mathematical lifecycle modeling efforts aimed at Delta Smelt and other species and for future adaptive management, including the fall outflow adaptive management and FLaSH studies effort.

**Framework for the Delta Smelt Conceptual Model**

The updated Delta Smelt conceptual model framework in this report integrates and modifies features of the “basic” and “species specific” POD conceptual models (Baxter et al 2010), the FLaSH conceptual model (Brown et al. 2014), the DRERIP “transition matrix” conceptual models (DiGennaro et al. 2012), and the hierarchical conceptual model in Miller et al. (2012). It consists of two nested and linked conceptual models of increasing specificity:

1. A general life cycle conceptual model for the four Delta Smelt life stages (adults, eggs and larvae, juveniles, and subadults) that includes stationary landscape attributes and dynamic environmental drivers, habitat attributes, and Delta Smelt responses; and
2. More detailed life stage transition conceptual models for each of the four Delta Smelt life stages that describe relationships between environmental drivers, key habitat attributes, and the population-level probability of successfully transitioning from one life stage to the next. This probability is dependent on the effects of environmental drivers and habitat attributes on the growth, survival, reproduction, and movements of Delta Smelt but data are currently inadequate to provide causal links for most of these processes individually.

**General Life Cycle Conceptual Model**

The updated general life cycle conceptual model for Delta Smelt (Fig. 8) follows the FLaSH Panels (2012) recommendation to “ensure Delta Smelt vital rates remain central to thinking” and is structurally similar to the basic POD conceptual model (Fig. 6). The general life cycle conceptual model is divided vertically and horizontally into four sections representing four Delta Smelt life stages from eggs and larvae to adults occurring in four “life stage seasons” indicated in the center of the diagram (Fig. 8; tier 5 box, green shading). This is similar to the four seasonal compartments of the species-specific conceptual model diagram in Baxter et al. (2010). Importantly, these life stage seasons are not exactly the same as calendar-based seasons. Instead, they have somewhat variable duration and overlapping months. This is because life stage transitions from eggs to adults are gradual and different life stages of Delta Smelt often overlap for a period of one to three months. Delta Smelt responses (Fig. 8; tier 4 box with dark blue shading) to important habitat attributes throughout their usually annual life cycle are placed within a box representing habitat attributes important to their growth and survival, which conveys the idea that biotic and abiotic habitat elements drive Delta Smelt responses (Peterson 2003; Fig. 8; tier 3 box with light blue shading). For each life stage season, there are a set of natural and anthropogenic environmental drivers associated with the estuarine environment (Fig. 8; tier 2 box with purple shading) that generate the habitat attributes important to Delta Smelt growth and survival. Surrounding the environmental drivers box is a fourth, outer box that represents the stationary (geographically and temporally fixed) landscape attributes of the estuarine ecosystem associated with its physical geometry and the orientation and connections of its component waterbodies (Fig. 8; tier 1 box with grey shading). In contrast to this outer box, the components and processes described in the inner boxes of this conceptual model are dynamic in space and time. Note that the fixed landscape attributes are considered fixed in the context of Delta Smelt population biology in any particular year rather than across longer time scales. The different spatial and temporal scales for each tier of the conceptual model are shown in Figure 9.

The tiered components of the general life cycle conceptual model for Delta Smelt can vary over a wide range of spatial and temporal scales (Fig. 9). Landscape attributes of the San Francisco Estuary (tier 1) encompass local to estuarine-wide features and change slowly over decades or longer periods. Environmental drivers (tier 2) that affect Delta Smelt habitat attributes vary and manifest over the broadest range of spatial and temporal scales, from local variations over tidal or daily cycles to long-term changes at the watershed or even larger geographic scales. Similar to environmental drivers, habitat attributes of Delta Smelt (tier 3) can be highly dynamic at small spatial and temporal scales or change gradually over many years, but they don’t extend beyond the geographic range of the species, which in the case of Delta Smelt is the SFE. Delta Smelt responses (tier 4) vary in response to changing habitat attributes within subregions of the estuary. In this small fish species with its maximum age of two years and extremely small geographic range, population-level responses can range from rapid (e.g., in response to toxic spills) to more
slowly over the course of one or more years. Life stage seasons (tier 5) occur over the course of a year in seasonally occupied areas of the estuary.

Similar to the POD and DRERIP conceptual models, the updated Delta Smelt life cycle conceptual model includes only those landscape attributes and environmental drivers with plausible mechanistic linkages to outcomes, which in this case are changes in habitat attributes and resulting Delta Smelt responses in the four life stage seasons. These mechanistic linkages are depicted as arrows in a series of four new conceptual models for each life stage season (Fig. 10). These life stage season conceptual models are nested components of the general life cycle conceptual model as shown in Fig. 8. They will be described in detail in Chapter 5.

Data Sources

Our examination of environmental drivers in Chapter 4, Delta Smelt life history and population dynamics and trends in Chapter 6, and the evaluation of hypotheses about Delta Smelt responses to changing habitat attributes in Chapter 7 rely largely on results of previously published data and analyses, but in several cases we update these analyses with more recent data. We also include some additional analyses (described below). All these analyses depend largely on environmental monitoring data collected by IEP agencies during routine, long-term monitoring surveys.

Figure 8. A new conceptual model for Delta Smelt showing Delta Smelt responses (dark blue box) to habitat attributes (light blue box), which are influenced by environmental drivers (purple box) in four “life stage seasons” (green box). Environmental drivers are influenced by landscape attributes (grey box).
These surveys provide the long-term records and geographic coverage necessary and the data collected by these surveys are publicly available. Available data includes data on fish, invertebrates, phytoplankton, water quality variables, and flow. Use of these particular data sources does not reflect any preference for those data. Results from other ongoing research efforts were included as appropriate.

For the purposes of this report, we consider each stage, larvae through adults, of the Delta Smelt life cycle in the context of the monitoring programs that provide data on the Delta Smelt population. Delta Smelt eggs are not monitored and have in fact never been found in the wild. Monitoring surveys in the late winter and spring include the spring Kodiak trawl (SKT, Fig. 11), which samples maturing, spawning and post-spawning adults. The SKT is conducted monthly from January through May. Spring also includes the 20 mm survey (20 mm, Fig. 12), which samples larval and post-larval Delta Smelt and is conducted every two weeks from mid-March through mid-July. Summer includes the summer townet survey (TNS, Fig. 13); which samples juvenile fish and currently runs every two weeks from June through August. The Fall Midwater Trawl (FMWT, Fig. 14) survey samples subadult Delta Smelt monthly from September through mid-December. Each of these surveys samples fishes broadly within the upper SFE and generally covers the geographic habitat range used by Delta Smelt (Merz et al. 2011). Exceptions to complete coverage occur in some high outflow years when Delta Smelt can temporarily inhabit San Pablo Bay in association with decreased salinities caused by increased Delta outflows (Moyle 2002) and in other years when some adult fish move upstream of the geographic range of these surveys (probably to spawn) in the Yolo Bypass and Sacramento River (e.g., Feyrer et al. 2006, Merz et al. 2011). Also, FMWT and TNS sampling in the Cache Slough complex was instituted over several years starting in the 1990s for FMWT and 2000s for TNS. The current sampling locations have been in place since 2011. These exceptions to complete spatial coverage are believed to reflect small fractions of the population. Additional geographic coverage along
or outside of the margins of the other four monitoring surveys is provided by other IEP fish monitoring surveys such as the San Francisco Bay Study, trawling and seining conducted by the Delta Juvenile Fish Monitoring Program in the Sacramento River and the north Delta, as well as the fish salvage monitoring at the fish protection facilities associated with the SWP and CVP export pumps in the south Delta. All Delta Smelt life stages (larvae-adult) are also commonly collected from nearshore habitats and in shallow open water where trawls cannot be used effectively (e.g., Aasen 1999, Nobriga et al. 2005, Brown and May 2006); however, there are no data indicating these are preferred habitats, that these fish represent different populations (see Fisch et al. 2011), or that their abundance varies differently than data from the aforementioned trawl surveys would suggest.

Annual abundance indices for Delta Smelt life stages are calculated from the catch data provided by each of the four surveys (See Appendix B for details). Together, they provide a comprehensive account of long-term changes in the relative abundance of Delta Smelt (Fig. 3). The long series of abundance index records for the summer and fall have provided the basis for many data analyses and modeling studies (e.g., Jassby et al. 1995, Kimmerer 2002a,b, Bennett 2005, Manly and Chotkowski 2006, Thomson et al. 2010, MacNally et al. 2010, Maunder and Deriso 2011, Miller et al. 2012) and for regulatory actions (USFWS 2008). They have also been used to estimate absolute population abundance (Newman 2008). The Delta Smelt and other SFE fish abundance indices are generally considered useful indicators of the status and trends of the Delta Smelt population as well as of the status of other resident fishes in the SFE in general and serve as performance metrics for the success of management actions. All monitoring surveys have strengths and weaknesses, and the long-term fish monitoring programs in the SFE are no exception (Honey et al. 2004). In the case of Delta Smelt, strengths include reasonably good coverage of the geographic extent of Delta Smelt habitat and coverage of all life stages except

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**Figure 10.** Framework for the Delta Smelt life stage season conceptual models.
eggs (Gaines et al. 2006). They also include exceptionally long and consistent data records going back to 1959 in the case of the TNS, the oldest of the four surveys described here. There is a large amount of ancillary data (covariates), including data collected during the fish surveys, additional fish data from other monitoring surveys (Honey et al. 2004) as well as invertebrate, phytoplankton, water quality and hydrological data. Possible weaknesses include no measure of precision of abundance indices and imprecise estimates due to a high frequency of zero catches of Delta Smelt. These problems combine with survey design issues such as differences in Delta Smelt catchability with different nets and trawl regimes under changing environmental conditions, behavioral changes in distribution (Newman 2008) and the current low abundance of the species. For example, several studies have shown that Delta Smelt can exhibit lateral and vertical movements associated with tide and time of day (Bennett et al. 2002, Feyrer et al. 2013, Bennett and Burau 2014) but the overall frequency or effects of such local movements on abundance indices are unclear. Studies to further evaluate and address these issues are currently underway.

Two of the four fish monitoring surveys described here specifically target Delta Smelt; the other two do not. The SKT was designed and implemented specifically to improve detection of maturing adult Delta Smelt moving upstream in the winter and spring, particularly into the central and south Delta (Souza 2002). The 20 mm survey was designed and implemented specifically to capture late-stage larval Delta Smelt of about 20 mm in length; the SKT and 20 mm survey data help managers assess the risk of entrainment of these life stages by south Delta
The TNS was designed to target small juvenile Striped Bass of about 17-50 mm fork length (the distance from the snout to the indentation of the tail fin) (Stevens 1977, Turner and Chadwick 1972); however, Delta Smelt tend to be of appropriate size for capture by the TNS net during the survey period. This occurs because Delta Smelt (see below) and Striped Bass spawning overlaps in time and growth of both are linked to water temperature, such that peak larval abundance occurs in April or May in most years. The TNS traditionally started and ended based on mean length of Striped Bass; however, young Delta Smelt attain sizes vulnerable to the TNS net during the same time period Striped Bass are vulnerable (Miller 2000). The survey ends when young Striped Bass surpassed 38 mm fork length (Miller 2000). Thus, regardless of the particular number of sampling surveys in a year or the index calculation method, Delta Smelt juveniles are generally vulnerable to the TNS whenever it samples. Similarly, the FMWT survey was designed to capture young-of-the-year Striped Bass, but in the 60-140 mm fork length size range (Stevens 1977). Although the survey and gear is generally effective for small pelagic fishes, the cod-end mesh (1.3 mm stretch mesh) on the net is large enough to allow some smaller sub-adult Delta Smelt to escape during the first couple survey months (see Newman 2008 for an approach to correct this effect). Even though the gear is not completely effective at retaining all sub-adult Delta Smelt, FMWT provides a reasonable relative measure of sub-adult abundance through time (Kimmerer and Nobriga 2005), albeit with low precision at the current low catch levels and given additional variation related to changes in growth, and thus changes in retention in the net from year to year. With the aforementioned caveats, we believe these surveys provide useful and valid relative abundance measures to examine the various life stage transition
relationships described in this report as well as in many of the previously published studies cited in this report.

In addition to the annual abundance indices for Delta Smelt provided by the monitoring surveys described above, we also present annual indices of recruitment and survival. In this report, a survival index is simply the ratio of an abundance index for a particular life stage divided by the abundance index for a preceding life stage of the same Delta Smelt cohort. A recruitment index is the ratio of an abundance index for a particular life stage divided by an abundance index for a life stage of the preceding Delta Smelt year-class. These types of indices have been used in previous analyses (e.g. Miller et al. 2012), but it is important to note that they may compound the observation errors inherent in the annual abundance indices in complicated ways. This is likely more problematic for survival and recruitment indices that use the TNS and FMWT abundance indices because these surveys were not specifically designed to target Delta Smelt. It may be less problematic for the recruitment index calculated by dividing the 20 mm abundance index for larval and post-larval Delta Smelt by the preceding SKT abundance index for adult Delta Smelt because both surveys specifically target Delta Smelt. We use this recruitment index in some additional analyses included in this report. All other survival and recruitment indices are only used as a rough approximation and illustration of differences in recruitment and survival rates among different annual cohorts and life stages; they are not used for additional analyses.
Data Analysis

As noted previously, we review long-term trends in this report using published results, but in some cases include some additional analyses of long-term monitoring data (Chapters 4 and 7). These analyses are kept deliberately simple, for example, simple graphical explorations of time series, examinations of simple statistics such as medians and arithmetic means, and investigation of univariate relationships using simple correlation and least squares regression analyses. Such analyses are readily reproducible with the publicly available data described above. The purpose of presenting the results of these new analyses is to update previously published information with the most recent data. In many cases, the data presented in this report are summarized using boxplots. The center horizontal line in each box represents the median of the data. The upper and lower ends of the box represent the upper and lower quartiles of the data. These are also known as “hinges.” The “whiskers” are the lines extending above and below the box. The whiskers show the range of values falling within 1.5 times the inter-quartile distance from the nearest hinge. Values outside this range are shown as individual symbols. Asterisks denote values within 1.5 to 3.0 times the inter-quartile distance and circles denote values greater than 3.0 times the inter-quartile distance. Other types of plots are explicitly identified in the figure caption.

Some graphs and analyses refer specifically to the POD period. Analyses suggest the POD period started as early as 2002 or as late as 2004 (Thomson et al. 2010). We somewhat arbitrarily selected 2003-present as the POD period for this report. This period is not being recommended.
as the baseline for management agencies to use when considering recovery of Delta Smelt. The time period simply reflects the consistently low level of Delta Smelt abundance in recent years and a useful baseline for identifying years with improved Delta Smelt abundance indices, which would indicate improved environmental conditions for Delta Smelt. Similarly, we also consider the 1982-2001 period between the two major step declines in Delta Smelt abundance identified by Thomson et al. (2010) separately in some graphs and analyses. Finally, some graphs and analyses refer to calendar years while others refer to water years. In California, a water year starts on October 1 and ends on September 30 of the next calendar year. California water year classifications are based on calculations of annual unimpaired runoff, which represents the natural water production of a river basin, unaltered by upstream diversions, storage, and export of water to or import of water from other basins.

In Chapter 7, we explore a series of hypothesized driver-outcome linkages using a comparative approach. The purpose is to demonstrate the utility of our conceptual model framework for generating hypotheses about the factors that may have contributed to the 2011 increase in Delta Smelt abundance. Specifically, we compare Delta Smelt responses to habitat conditions in four recent years with moderate to wet hydrology: the two most recent wet years (2006 and 2011) and the two drier years immediately before them (2005 and 2010). This comparative approach and data sources (described in Chapter 4) are deliberately similar to the comparative approach used in the FLaSH investigation (Brown et al. 2014). This approach allows us to place the results of the FLaSH investigation in a year-round, life cycle context and to more comprehensively evaluate factors that may have been responsible for the strong Delta Smelt abundance and survival response in 2011, including any possible relevant antecedent conditions from 2010. We attempt to draw comparisons with a similar set of data collected during 2005 and 2006. Our working assumption is that different Delta Smelt abundances in 2006 and 2011 should be attributable to differing environmental conditions, in some cases attributable to management actions, and subsequent ecological processes affecting the Delta Smelt population.

In Chapter 9 we briefly describe three examples of additional mathematical modeling approaches that can be used to further explore some of the linkages and interactions in our conceptual models and complement previously published and other ongoing mathematical modeling efforts for Delta Smelt. Importantly, results from the three modeling examples in Chapter 9 are included for illustrative purposes only; peer-reviewed publications of these analyses need to be completed before they can be used to draw firm conclusions.

Chapter 4: Environmental Drivers and Habitat Attributes

The general approach of this Chapter is to focus on how environmental drivers and interactions among them create habitat attributes of importance to Delta Smelt. Specifically, we review and synthesize existing information about drivers and habitat attributes and Delta Smelt responses to habitat attributes with a focus on new information since Baxter et al. (2010). We use the drivers and habitat attributes depicted in the basic POD conceptual model (Fig. 6) as the basis for this synthesis. We consider habitat attributes important when there are published studies suggesting ecological responses by Delta Smelt. Each section focuses on a habitat attribute that can be the outcome of one or more environmental drivers. Physical habitat attributes are presented first,
followed by biological habitat attributes. The order of presentation does not imply any kind of ranking of relative importance. For simplicity, we consider all habitat attributes discussed here as equally important because, as noted in Chapter 2, habitat arises from the combination of all physical and biological attributes affecting a species. We fully acknowledge that as Delta Smelt research proceeds and the system continues to change, additional habitat attributes may need to be added to the conceptual model, while others may be deemphasized or even deleted.

Each section starts with the general importance of a specific habitat attribute for estuarine biota followed by a brief discussion of its linkages with environmental drivers and its dynamics in space and time. Each habitat attribute is then placed in the context of Delta Smelt biology.

**Water Temperature**

Water temperature is fundamental to aquatic ecosystem health and function. It directly influences biological, physical, and chemical properties such as metabolic rates and life histories of aquatic organisms, dissolved oxygen levels, primary productivity, and cycling of nutrients and other chemicals (Vannote and Sweeney 1980, Poole and Berman 2001, Null et al. 2013). Water temperature is an important variable for ectothermic (“cold-blooded”) animals, including all fishes and invertebrates in the SFE. In the most extreme case, when water temperature exceeds the thermal tolerance of an organism, it will die. Temperatures within the thermal tolerance of an organism control the rate and efficiency of many physiological processes, including activity, digestion, growth, reproductive development, and reproductive output. We return to these processes after giving an overview of water temperature variability and its drivers in the Delta.

Long term temperature records from selected sites in the SFE show substantial seasonal and daily fluctuations in water temperature (Kimmerer 2004). While daily variations are evident and likely important to organisms, seasonal variations are much greater (Wagner et al. 2011). Median water surface temperatures across all stations monitored by the IEP Environmental Monitoring Program (EMP) (Fig. 15) from 1975-2012 range from 9 °C in January (minimum: 6 °C) to 22 °C in July (maximum: 28 °C). There are also clear regional variations in water temperature (Fig. 16). In July and August, the hottest summer months, water temperatures are usually highest at monitoring stations in the south Delta (average 23-26 °C, maximum 28 °C), lower at stations in the northern and western Delta (average 21-23 °C, maximum 25 °C) and lowest at stations in Suisun and San Pablo Bays (average 19-21 °C, maximum 24 °C). In January, the coldest winter month, average water temperatures are uniformly below 10 °C in the entire Delta, but above 10 °C in San Pablo Bay.

There is currently little evidence for increasing water temperatures in the Delta, although with climate change such increases are expected over the course of the century (Cloern et al. 2011, Wagner et al. 2011, Brown et al. 2013). In Spring (March-June) water temperature at IEP EMP water quality monitoring stations in the Delta increased during 1996–2005 by about 0.2 °C per year, but a similar trend was not apparent for the longer-term data record from 1975-2005 or for stations in Suisun Bay (Jassby 2008). These findings are similar to the results of Nobriga et al. (2008) who found no long-term (1970-2004) trends in temperature data collected during summer fish monitoring surveys in the Delta. Nobriga et al. (2008) also noted that the long-term (1970-2004) mean July water temperature at TNS fish monitoring stations in the southern region of the Delta is 24 °C, with current mid-summer temperatures often exceeding 25 °C. This agrees with average monthly EMP data from 1975-2012 which shows July and August water temperatures at
a monitoring station located in Old River (station D28A) and in the San Joaquin River near the Port of Stockton (station P8) of more than 24 °C and 25 °C, respectively (Fig. 16).

In tidal systems, water temperature at a particular location is determined by the interaction between atmospheric forcing (e.g., air temperature and wind), tidal dispersion and riverine flows across the estuarine landscape (Monismith et al. 2009). In particular, estuarine water temperature is driven by heat exchange at the air–water interface and mediated by tidal and riverine flow dynamics and estuarine geomorphology (Enright et al. 2013). Wagner et al. (2011) found that regional weather patterns including air temperature and insolation (sunlight), are the primary drivers of water temperature variations in the SFE. Water flow and interaction with the stationary topography of the system also affects water temperature in the SFE, especially over shorter time scales and at smaller spatial scales. For example, Enright et al. (2013) showed that interaction
of tides with tidal marsh topography can have a mediating effect on water temperature in tidal sloughs and on thermal variability at smaller spatial scales. Wagner et al. (2011) showed that high winter and spring flows can temporarily lower water temperatures. Greenberg et al. (2012) found that the present riparian vegetation on Delta levees lowers insolation by about 9% compared to a hypothetical situation without vegetation and suggested that riparian vegetation thus contributes to locally cooler water temperatures. This suggests that at least to some degree, water temperature can be managed locally and for short periods. Over larger scales, however, these types of locally mediated effects are overwhelmed by the effects of air temperature and insolation.

Air temperature and insolation in the SFE are correlated with each other (Wagner 2012) and vary strongly with proximity to the Pacific Ocean because of the contrasting climate regimes prevailing in inland central California and the central California coast. While inland central California has a large annual air temperature range with hot, dry, sunny summers and cool, wet, and often foggy winters, the central California coast has a smaller annual air temperature range with cooler and often foggy summers and milder winters (Conomos et al. 1985). The SFE has a transitional climate with greater spatial and temporal variability in air temperature than either the coastal or the inland regions (Whipple et al. 2012). This is due to the interplay of the dynamic air masses from these regions across the stationary estuarine topography. In the summer, this interplay often results in strong afternoon winds from the ocean locally known as the “Delta breeze.” These onshore winds usually advance into the western and central Delta and, depending on the depth of the marine layer, often also into its marginal areas. In the Delta, these southwest to northeast winds can persist throughout the night and into the next morning and produce a marked decline in daily temperature. In the morning, this low is often followed by rapid warming once the winds subside and the high temperature inland air masses return to dominance (National Weather Service 2003). In the winter, ocean winds are weak and, during calm periods, cold air flows from the mountains into the estuary. This results in the formation of dense, overnight, near-surface fog locally known as “tule fog.” These calm and foggy periods are interrupted by winter storms. Many of these storms arrive from the south and southeast as “atmospheric rivers” that can often produce gale force winds and heavy rains lasting several days (Conomos et al. 1985, Dettinger and Ingram 2013).

The large variability in air temperature in the Delta is reflected by the larger annual variability in water temperature measured from 1998-2002 at continuous monitoring stations in the interior Delta compared to stations further upstream or downstream (Wagner et al. 2011). This high variability is also apparent in monthly water temperature data collected by the IEP Environmental
Monitoring Program since 1975 (Fig. 11). From 1975 to 2012, annual fluctuations in average monthly water temperature were greatest at stations in the south Delta (14-16 °C), smaller at stations in the northern and western Delta (12-13 °C), and lowest at stations in Suisun and San Pablo Bays (9-12 °C). Jassby (2008) reported that maximum daily air temperature could explain almost half the variability in maximum daily water temperature at the continuous monitoring station at Antioch during the summer months. The relationship between air and water temperature was also strong in all other months except January.

Wagner et al. (2011) and Wagner (2012) developed simple regression models for predicting water temperature at fixed temperature monitoring stations in the SFE using only air temperature and insolation on the day of interest and the water temperature from the previous day. Water temperature from the previous day accounts for both previous air temperature and the sources of water to the site, including advective flow from rivers or dispersive flow from more downstream reaches of the SFE. Each model had a different set of coefficients because of the differing influences of incoming river water or tidal exchange with San Francisco Bay. For stations with greater than 1 year of calibration data, model $R^2$ for daily average temperature exceeded 0.93, indicating that water temperature was highly predictable within the limits of the calibration data sets. High winter and spring flows were responsible for the largest divergences of the model outputs from measured temperatures.

The simple statistical models for water temperature developed by Wagner et al. (2011) and Wagner (2012) should be used with caution because they only predict temperature at the site of the recording instrument and do not explicitly account for mechanistic heat exchange. The analyses therefore do not incorporate the possible effect of site-specific features such as shading by riparian vegetation (Greenberg et al. 2012). Similarly, there are lateral and vertical variations in temperature on daily time scales (Wagner 2012) that could be important to organisms. For example, such variation might include substantial heterogeneity and formation of thermal refugia, which may be important to Delta Smelt.

In contrast to statistical modeling, which produces site-specific results, water temperature across regions is commonly modeled with computation-intensive deterministic simulation models. Such models use energy budgets to predict water temperature. Simple stochastic models are also possible. Like most statistical models, these stochastic models generally rely on the relationship between air and water temperature (Caissie 2006, Null et al. 2013). We are not aware that these types of models have been developed for the San Francisco Estuary.

Upper temperature limits for juvenile Delta Smelt survival are based on laboratory studies and corroborated by field data. Interpretation of the laboratory results is somewhat complicated as temperature tolerances can be affected by various factors including acclimation temperature, salinity, turbidity, and feeding status. Based on the critical thermal maximum, $CT_{\text{max}}$, juvenile Delta Smelt acclimated to 17 °C could not tolerate temperatures higher than 25.4 °C (Swanson et al. 2000). However, for juvenile Delta Smelt acclimated to 11.9, 15.7 and 19.7 °C, consistently higher $CT_{\text{max}}$ were estimated (27.1, 28.2 and 28.9 °C, respectively; Komoroske et al. 2014), which corresponded closely to the maximum water temperatures recorded in the TNS and FMWT surveys. Swanson et al. (2000) used wild-caught fish, while Komoroske et al. (2014) used hatchery-reared fish, which may have contributed to the differences in results. Based on the TNS (Nobriga et al. 2008) and the 20 mm Survey (Sommer and Mejia 2013), most juvenile Delta Smelt were predicted to occur in field samples when water temperature was below 25 °C. In a multivariate autoregressive modeling analysis with 16 independent variables, MacNally et al. (2010) found that high summer (June – September) water temperature had a negative effect
on Delta Smelt subadult abundance in the fall. Water temperature was also one of several factors affecting Delta Smelt life stage dynamics in the state-space model of Maunder and Deriso (2011) and in an individual-based Delta Smelt life-cycle model (Rose et al. 2013a,b).

In addition to lethal effects, water temperature also has direct effects on the bioenergetics (interaction of metabolism and prey density) of Delta Smelt (Bennett et al. 2008) and it may affect their tolerance to other habitat attributes, such as toxicity (Brooks et al. 2012) and predation risk. Responses of different life stages of Delta Smelt to various temperature, salinity, and turbidity conditions are currently being further assessed as part of a larger UC Davis laboratory study about the “fundamental niche” of Delta Smelt (Komoroske et al. 2014, R. Connon et al., U.C. Davis, unpublished data).

The topic of bioenergetics is an important consideration in much of the remainder of this report, so we address it in more detail here. In general, the total metabolic rate of a fish will increase with temperature to an optimum temperature at which, given unlimited food, there is the maximum ability to grow and develop reproductive products (eggs or sperm) in addition to maintaining the basal metabolic rate required for survival, which also increases with temperature (Houde 1989, Hartman and Brandt 1995). As temperature increases beyond the optimum, metabolic rate continues to increase but physiological processes become less and less efficient and more energy is required just to meet the basal metabolic rate of the organism. Eventually, the metabolic rate begins to decline as temperatures approach the physiological limits of the organism and the basal metabolic rate can no longer be maintained.

At temperatures beyond the optimum, the ability to grow and mature becomes increasingly impaired. Long-term exposure to such stressful temperatures can eventually be lethal. In addition, resistance to disease and contaminants can also be affected (Brooks et al. 2012). The responses to contaminants can vary depending on the type of contaminant. For example, low temperatures can decrease the toxicity of organophosphate insecticides, but increase the toxicity of pyrethroid and organochlorine insecticides (Harwood et al. 2009), a characteristic that has been used in toxicity identification and evaluation (Weston and Lydy 2010). The previous discussion assumes unlimited food, which is unlikely to be the case for Delta Smelt or any organism in nature. Even at the optimum temperature, growth and reproductive development will depend on the quantity and quality (energy and nutrient content) of the food consumed. If the fish is unable to ingest enough food to meet its nutrient and energetic requirements, including the energy expended to capture and digest prey, it will starve, after first depleting any available energy stores (fat or muscle). Given an array of food items, fish will generally choose larger prey items. This is because the energy required to detect, chase, and capture multiple smaller prey that are equivalent in nutritional value to a single large prey item will, in many cases, exceed the energy required to capture the single prey item. Note that these same ideas apply to predatory fish that might consume Delta Smelt.

Water temperature is also thought to affect the number of eggs produced by female Delta Smelt. Egg production (i.e., fecundity) of the population is influenced not only by individual female size and number (Bennett 2005, DFW unpublished), but also by the duration of a temperature “spawning window” (Bennett 2005, Mac Nally et al. 2010), variously defined as: 15-20 °C by Bennett (2005); 7-15 °C by Wang (1986); and 12-15 °C by Baskerville-Bridges et al. (2004b). Bennett (2005) further stated that during cool springs this spawning window persists longer, allowing more cohorts to recruit. Given a sufficiently long spawning window, individual females may also repeat-spay during the spawning season. This has been documented in culture (see Bennett 2005; J. Lindberg, U.C. Davis, personal communication 2013) and appears to occur
in the wild as well (L. Damon, CDFW, written communication 2012). Lindberg (U.C. Davis, personal communication 2013) observed that most females in culture spawned twice, some spawned three times and a very small number spawned four times. Each spawning was separated by a 4-5 week refractory period during February through June when water temperatures remained within the spawning window. Though laboratory conditions may not necessarily be representative of conditions in the wild, ripe females ready to release their second complete batch of eggs and developing a third batch have been detected in the wild during March and April (i.e., mid-season) suggesting that three spawns are possible (L. Damon, CDFW, written communication 2012). Thus, a longer spawning window would allow more females to repeat spawn adding both additional cohorts hatching under different conditions, and multiplying the fecundity of each repeat spawner (i.e., increasing the total fecundity of the individual), and thus, the total fecundity of the population. Moreover, in culture, individual females continued to grow through the spawning season and become more fecund with each batch of eggs (J. Lindberg, U.C. Davis, personal communication 2013). In the wild, the size of mature females generally increases month to month through the spawning season (Fig. 17), suggesting a potential increase in fecundity with each batch, but this has yet to be confirmed for wild fish. However, in culture, fish hatched later in the spawning season (mid-May to mid-June) grew up to be smaller-sized adults that started spawning later and had progeny with lower survival than the progeny of fish hatched earlier in the season (Lindberg et al. 2013). These observations are consistent with the reproductive patterns suggested for the wild Delta Smelt population (Bennett 2011). Overall, the effect of a prolonged spawning season on Delta Smelt population size and dynamics would seem to be positive; however, there is some uncertainty.

In the culture experiments reported by Bennett (2005), temperature strongly influenced hatching success of eggs. Specifically, Bennett (2005) reported that optimal hatching success and larval survival were estimated to occur at 15–17 °C based on studies conducted at 10, 15, and 20 °C. The data indicated that as incubation and early rearing temperatures increased, size at hatching and size at first feeding linearly decreased, possibly because basal metabolism of the developing embryo used more energy leaving less for growth. Fish that hatch relatively late in the season may experience high temperatures at a small size, which may reduce larval survival by several possible mechanisms. First, small size would limit the size of food items that the larvae could ingest because of smaller mouth size (see Nobriga 2002). Temperature may also affect food type and availability as discussed below. Second, small larvae are likely vulnerable to a larger range of predators for a longer period compared to larger larvae (e.g., “stage duration hypothesis;” Anderson 1988). Third, these fish could be potentially more vulnerable to transport toward the CVP and SWP export facilities, when Old and Middle River (OMR) flow restrictions are lifted. Restrictions are lifted when the 3-day mean water temperatures in Clifton Court Forebay (CCF) reach 25 °C or by the end of June.

As explained above, higher water temperatures increase energetic requirements and thus the food requirements of fish. To meet the increased need for food, it is possible that Delta Smelt spend more time foraging during the day. Since greater foraging time during the day increases visibility to predators, and those predators would also increase their foraging rates at higher temperatures, the encounter rate of predator and prey would likely increase at higher water temperatures. The net effect could be an increase in Delta Smelt predation risk (e.g., Walters and Juanes 1993). High temperatures can also decrease antipredator behavior, as described for Sacramento River Chinook Salmon (Oncorhynchus tshawytscha) (Marine and Cech 2004). In other words, the fish may make a behavioral choice to feed, grow, and become less vulnerable to predators as rapidly as possible, even though the short-term predation risk might increase. Water temperatures in the upper SFE are usually highest from June to September and decline rapidly between October and December.
The reported optimal culture temperatures for Delta Smelt larvae and late-larvae are 16.4 ± 0.25 °C (Komoroske et al 2014). Moreover, the chronic lethal thermal maximum for Delta Smelt varies by life stage (Komoroske et al. 2014). Juvenile and subadult Delta Smelt are observed in the field most commonly at temperature near or below 20 °C (Bennett et al. 2008, Nobriga et al. 2008), a temperature which is often exceeded beginning in May or June and continuing through September and more rarely in October (see Chapter 7). Thus, we suggest that the same tradeoffs between feeding and predation risk may persist through the warmer months and into early fall, but become less likely as the season progresses into late fall and winter. Note, however, that predation risk is also influenced by a complex suite of other factors such as turbidity, life stage, and proximity to predator habitat, so the level of risk to Delta Smelt can’t be determined.

Another possible indirect effect of higher water temperatures is that they may promote harmful algal blooms (HAB) (Lehman et al. 2005), which may degrade Delta Smelt habitat quality in the summer and early fall (Baxter et al. 2010). In the Delta, Lehman et al. (2013) found that blooms of the harmful cyanobacteria (blue-green algae) *Microcystis aeruginosa* required a water temperature of at least 19 °C for initiation. Other drivers of HABs and the possible effects of HABs are discussed more fully in a separate section of this Chapter. The combination of large seasonal and regional water temperature variability in the SFE and substantial direct and indirect effects of water temperature for all life stages of Delta Smelt means that this variable should be considered one of the most important habitat attributes for Delta Smelt. Differences in water

**Figure 17.** Individual female fork lengths by calendar day for mature female Delta Smelt collected in the Spring Kodiak Trawl Survey, January through May, 2005, 2006, 2010 and 2011. These data include both monthly distribution survey fish and directed survey fish. The directed survey (which targeted smelt spawning areas) was discontinued after January 2010.
temperature between regions or time periods may have important effects on the Delta Smelt population (Rose et al. 2013b).

**Salinity and the Size and Location of the Low Salinity Zone**

A dynamic salinity gradient from fresh water to salt water is one of the most characteristic features of an estuary (Kimmerer 2004). It originates from the mixing of fresh inland water with salty ocean water through tidal dispersion and gravitational circulation (Monismith et al. 2002). Many estuarine-dependent organisms occur in distinct salinity ranges (e.g., Kimmerer 2002a) and the extent and location of water with suitable salinities is thus an important habitat attribute for estuarine organisms. Over the time period of available monitoring data, there is no clear long-term trend in salinity levels and distributions in the estuary. Significant increases and decreases linked to changing flow patterns have been detected for various stations and months (e.g., Jassby et al. 1995, Enright and Culberson 2009, Shellenbarger and Schoellhamer 2011, Cloern and Jassby 2012).

The brackish (oligohaline) “low salinity zone” (LSZ) is an important region for retention of organisms and particles and for nutrient cycling. In the SFE, the LSZ provides important habitat for numerous organisms including Delta Smelt (Turner and Chadwick 1972, Kimmerer 2004, Bennett 2005). In this report we define the LSZ as salinity 1-6; however, other salinity ranges have been used by others, such 0.5-6 (Kimmerer et al. 2013) or 0.5-5 (Jassby 2008).

In the SFE, the position of the LSZ is commonly expressed in terms of X2, which is the distance from the Golden Gate in km along the axis of the estuary to the salinity 2 isohaline measured near the bottom of the water column (Jassby et al. 1995). X2 represents the approximate center of the LSZ (Kimmerer et al. 2013).

X2 is an index of the physical response of the estuary to freshwater outflow from the Delta; it decreases with increasing outflow because increasing freshwater outflow prevents seawater from moving landward. The X2 index was developed two decades ago as an easily-measured, policy-relevant “habitat indicator.” Its ecological significance for multiple species and processes was established through statistical analyses of biological responses to seasonally or annually averaged X2 values (Jassby et al. 1995) and has since been reaffirmed in additional studies (e.g., Kimmerer et al. 2002a,b, 2009, 2013, Thomson et al. 2010, Mac Nally et al. 2010). There is, however, still much uncertainty regarding the causal mechanisms for the observed biological responses of biota to X2. As with all statistically derived functional relationships, biological responses to X2 do not necessarily reflect direct causal relationships and it is generally recognized that some of the causal mechanisms may not be directly linked to the size and location of the LSZ.

Most of the scientific and management attention has focused on the LSZ and X2 from late winter to early summer (hereafter “spring X2”) depending on the species of interest, but in recent years the LSZ and X2 during the fall months (“fall X2”) has also received considerable scientific and policy attention. Annual abundance indices of several estuarine fish and invertebrate species have a negative relationship with spring X2, meaning that abundance indices increase when X2 and the LSZ are more westward and Delta outflow is higher in the late winter and spring months (Jassby et al. 1995, Kimmerer 2002a, Kimmerer et al. 2009). Delta Smelt summer abundance indices have a significant relationship with prior fall X2 and fall abundance (USFWS 2008, Mount et al.
2013). Changes in spring and fall X2 have also been linked to long-term fish declines in the SFE (Thomson et al. 2010, Mac Nally et al. 2010).

The size and location of the LSZ are considered key factors determining the quantity and quality of low salinity rearing habitat available to Delta Smelt and other estuarine species. LSZ size and location are determined by the interaction of dynamic tidal and river flows with the stationary topography of the region (Reclamation 2011, 2012, Kimmerer et al. 2013). In a recent study, Kimmerer et al. (2013) used the three-dimensional hydrodynamic “UnTRIM” model which has an unstructured grid (Casulli and Zanolli 2002, 2005) to produce detailed maps of the distribution of salinity in the SFE under different outflow conditions. These maps (figure 2 in Kimmerer et al. 2013 and Fig. 18 and 19 in this report) show that under low outflow conditions typical of summer and fall months (outflow = 140 m$^3$ s$^{-1}$, X2 = 85 km), the LSZ is in the western Delta confluence region, including the Sacramento and San Joaquin Rivers upstream of Chipps Island (Fig. 18), while under high outflow conditions typical of wet winter months (outflow = 1,440 m$^3$ s$^{-1}$, X2 = 51 km), the LSZ is much further west in San Pablo Bay. At intermediate outflows (intermediate X2 = 74 km), it is located east of Carquinez Strait and covers Suisun Bay and parts of Suisun Marsh (Fig. 19).

Kimmerer et al. (2013) also examined the relationships between X2 and the area, average depth, and volume of the LSZ. They found that these relationships were bimodal, with the largest volumes and areas and shallowest depths at X2 values below 50 km when the LSZ is located in the large San Pablo Bay, and secondary peaks at X2 values between 60 and 75 km when the LSZ overlays the smaller Suisun Bay (Fig. 20). Area and volume were smallest and depth greatest when the LSZ was constricted in Carquinez Strait (X2~50-60 km) and in the confluence region of the Sacramento and San Joaquin Rivers (X2~80-85 km).

Paleosalinity investigations going back several thousand years indicate that the Delta has historically been largely fresh, while the Suisun region has alternated between brackish (oligohaline) and fresh (Ingram and Malamud-Roam 2013, Drexler et al. 2014). The LSZ and X2 likely moved according to predictable annual and interannual rhythms. Interannually, X2 was most variable in the higher-flow winter and spring months and least variable in the low-flow fall months. Seasonally X2 moved from the west in winter and spring to the east in summer and fall. CDWR (CDWR 2007) computes monthly “unimpaired” outflows which remove the effects of dam operations and water diversions. Annual X2 dynamics based on these unimpaired flows may give a sense of these historical fluctuations (Fig. 21). It is important to note, however, that unimpaired flows are not the same as historical “natural” flows because they do not take into account upstream water losses (e.g., consumption and evaporation) or physical water body alterations such as channelization, groundwater depletion, draining of wetlands, and disconnection of floodplains. The historical wetlands, floodplains, and groundwater basins would have naturally retained and released water (Whipple et al. 2012) and likely affected flows and the LSZ in different ways than today’s man-made reservoirs. Work is currently underway at UC Davis, the San Francisco Estuary Institute, and elsewhere to explore these issues, but results have not yet been published (W. Fleenor, U.C. Davis, personal communication). At this time, considerable uncertainty remains regarding the natural ranges in the timing and volume of the historical seasonal and interannual freshwater flows and how they caused the LSZ to spread out and contract across the estuary’s historical landscape. There is, however, little doubt that interannual variations in precipitation and hence river flows caused a high degree of interannual variability in the size and location of the low-salinity zone (Dettinger 2011).
There is also no doubt that human water use and landscape alterations have changed flows into and out of the Delta and, consequently, salinity dynamics in the SFE, though changing precipitation patterns also play a role (Enright and Culberson 2009). Before the construction of today’s major reservoirs, upstream water diversions coupled with the isolation of floodplains and wetlands, which had naturally stored runoff, from river channels by levees exacerbated salinity intrusions into the Delta in dry years. This was especially evident during the severe drought from...
1929 to 1934 when salinities of 2 were observed at Paintersville Bridge which is located on the Sacramento River at a distance of about 136 km from the Golden Gate (Mathew 1931). Operation of the large CVP and SWP reservoirs that were constructed after this drought has prevented such severe salinity intrusions since then and X2 has remained west of Rio Vista located on the Sacramento River 100 km upstream of the Golden Gate. Beginning with the salinity requirements in SWRCB water right decision D-1275 of 1967, salinity and the position of the LSZ have also

**Figure 19.** Salinity distribution at intermediate outflow. The upper panel shows the area of the low-salinity zone (9,140 hectares) at X2 = 74 km (at Chipp Island). The lower panel shows the percentage of day that the low-salinity zone occupies different areas.
been increasingly regulated to protect “beneficial uses,” including habitat and fish protections (see Chapter 1).

CVP and SWP water exports from the Delta began in the early 1950s with the completion of the CVP C.W. Bill Jones Pumping Plant (formerly known as the Tracy Pumping Plant) in 1951 and then increased with the completion of SWP’s Harvey O. Banks Pumping Plant in 1968. Long-term variability in the trend of Delta outflow has been reduced seasonally for the period 1921-2006, in part due to water project operations (Enright and Culberson 2009), but also due to overriding climate changes. Analyzing data from 1956–2010, Cloern and Jassby (2012) found significant increases in water exports from the Delta in all months of the year except May, but in the first half of the year, these increases in exports did not significantly affect Delta outflow. We
show this by plotting the relationship between the Sacramento River Water Year Index, a measure of runoff, and average spring X2 (February-June) for two periods before (1956 to 1999) and after (2000-2013) the current flow and salinity requirements in SWRCB water right decision D-1641 became mandatory. The relationship appeared to remain essentially unchanged when the two time periods were compared (Fig. 22a). Cloern and Jassby (2012) further found that inflow to the Delta significantly increased in July and August, but these increases in inflow did not translate into significant increases in Delta outflow due to concurrent increases in exports during these months. Nevertheless, plots of recent data show that July and August outflows increased and the relationship between the Sacramento River Water Year Index and summer-time X2 (July-August) shifted downward in the years since the SWRCB water right decision 1641 went into effect in 2000 relative to previous years (Fig. 22b). The wet year 2006 did not fit this pattern because it had high summer X2 in spite of a high water year index. This means that with the exception of 2006, the LSZ has generally been located somewhat more westward in July and August since 2000 than from 1956 to 1999 under similar runoff conditions.
Figure 22. Plots of monthly X2 as a function of the Sacramento River Water Year Index (a measure of runoff) for the years 1956 to 1999 and 2000 to 2013 for: a, winter/spring; b, summer; and c, fall. The regression equation for each set of points is also shown. The index is calculated as: 0.4 * Current April to July Runoff Forecast (in millions of acre feet, maf) + 0.3 * Current October to March Runoff in (maf) + 0.3 * Previous Water Year's Index (if the Previous Water Year's Index exceeds 10.0, then 10.0 is used) (see http://cdec.water.ca.gov/cgi-progs/iodir/WSIHIST for further detail).
Cloern and Jassby (2012) also showed that significantly increasing exports combined with declining inflows led to significant declines in Delta outflow in each month from September to December. In plots of recent data, this led to a shallower slope of the relationship between the Sacramento River Water Year Index and fall X2 (September-December) and a more eastward LSZ location in the fall months of wetter years (below normal, above normal, and wet water year types) during 2000-2012 compared to 1956-1999, with the exception of two wet years at the end of the time series, 1997 and 1999, which fall on the 2000-2012 line (Fig. 16c, see also Feyrer et al. 2007, 2010). The areas with light blue shading in the three plots shown in Figure 16 show the range of X2 that places the LSZ over Suisun Bay and are associated with a high LSZ volume, area, and shallow LSZ depths (Kimmerer et al. 2013, Fig. 14). Fall X2 commonly fell into this range from 1956-1999 (in 18 of 44 years; Fig. 22c), but never after 2000. In 2011, the most recent wet year, fall X2 was lower than in the preceding wet years of 2006, 1997, and 1999, but still elevated relative to the majority of previous wet years. Overall, the changes in flows in the summer and fall months described by Cloern and Jassby (2012) have resulted in more muted seasonal and interannual variations in X2 and in the size and location of the LSZ in more recent years and possibly also relative to historical variability (Fig. 21).

Delta Smelt are found in the estuary at salinities up to 18 (Bennett 2005), but are most common in the in the LSZ (< 6) (Moyle et al. 1992, Sommer and Mejia 2013, Kimmerer et al. 2013). Sommer et al. (2011a) described Delta Smelt as a “diadromous species that is a seasonal reproductive migrant.” In the winter, adult Delta Smelt move upstream into fresh water for spawning. In the spring and summer, young Delta Smelt are transported or swim downstream into the LSZ (Dege and Brown 2004). Delta Smelt usually rear in low salinity habitat in the summer (Nobriga et al. 2008) and fall (Feyrer et al. 2007), although some Delta Smelt remain year-round in fresh water (Sommer et al. 2011a, Merz et al. 2011, Sommer and Mejia 2013).

The recruitment success of Longfin Smelt and age-0 Striped Bass increases linearly with more westward positions of the LSZ during spring (Jassby et al. 1995, Kimmerer 2002a). In contrast, the relationships of annual Delta Smelt indices with spring X2 are more complex because they have not been consistent over the period of record (Fig. 23). Jassby et al. (1995) found that from 1968-1991, the highest fall abundance indices for Delta Smelt coincided with intermediate values of average April-July X2 when the LSZ was positioned in Suisun Bay. Low fall abundances were, however, also observed at these intermediate X2 values. The analyses by Jassby et al. (1995) were later updated and augmented with an analysis of the relationship between Delta Smelt summer abundance and spring X2 (Kimmerer 2002a, Kimmerer et al. 2009).

We updated the analyses by Jassby et al. (1995) with more recent data and data from additional monitoring surveys to examine the hypothesis that during periods of relatively stable abundance (i.e. without step changes, Thomson et al. 2010), the abundance of different Delta Smelt life stages is related to spring outflow and the position of the LSZ as expressed by spring X2. To obtain spring X2, we first calculated mean monthly X2 values calculated from daily X2 values. We then averaged the mean monthly X2 values for February to June. This is different from the April-July period used by Jassby et al. (1995) for their Delta Smelt analyses, but similar to the spring X2 averaging period used by Kimmerer (2002a). Note that different averaging methods for calculating seasonal X2 values account for the small quantitative differences between results presented here and those of previously published analyses that used the same data, but this does not affect the overall patterns. We partitioned the data into the periods before, between, and after the 1981 and 2002 step declines in Delta Smelt abundance identified by Thomson et al. (2010). The 1981-1982 partition, but not the 2002-2003 partition, has been previously applied by Kimmerer (2002a) and Kimmerer et al. (2009).
Kimmerer (2002a) and Kimmerer et al. (2009) found that the relationship between spring X2 and Delta Smelt juvenile abundance indices was positive before the step decline in Delta Smelt abundance that started in 1981 (Thomson et al. 2010), suggesting that historically, Delta Smelt population recruitment may have benefitted from lower outflows and a more upstream LSZ in the late winter and spring. In our analysis, we found that the relationship was perhaps more unimodal than linear (Table 1, Fig. 23a) because a model that included a quadratic spring X2 term explained more of the variation in the data than a linear model that did not, although the statistical significance of the linear model was slightly higher than that of the quadratic model because of the loss of a degree of freedom due to the additional quadratic term included in
the quadratic model. Similar to Kimmerer (2002a) and Kimmerer et al. (2009), we found that in the period after the 1981 step change and also in the period after the 2002 step change, the relationship of log-transformed summer abundance with spring X2 shifted downward and became more clearly negative than unimodal (Fig. 23a). The relationship remained statistically significant at the $P < 0.05$ level in the period after the 1981 step decline, but is no longer statistically significant after 2001. Similarly, the relationship is also not significant across the entire 52-year time series (Table 1).

Kimmerer et al. (2009) found a non-significant and essentially flat relationship between spring X2 and the entire log-transformed sub-adult abundance time series for Delta Smelt; this remains the case when data from the five most recent years is included in the analysis (Table 1). Similar to Jassby et al. (1995), we found a weakly unimodal relationship between spring X2 and log-transformed Delta Smelt subadult abundance indices before the first step change, but this relationship was not statistically significant at the $P < 0.05$ level (Table 1, Fig. 23b). Similar to juvenile abundance, the relationship of log-transformed subadult abundance with spring X2 shifted downward in the periods after each of the two step changes and became more negative than unimodal (Fig. 23b), but again these relationships were not statistically significant at the $P < 0.05$ level (Table 1).

Taken together, these findings are generally consistent with previous conclusions that moderate hydrological conditions in the late winter and spring and a large LSZ located in the Suisun region can be beneficial to Delta Smelt population abundance (Jassby et al. 1995). Historically, this may have been the case for several life stages. At present, however, juvenile and subadult Delta Smelt seem to barely respond to spring X2. As Jassby et al. (1995) point out, this does not mean that there is no longer an effect of spring X2 on juveniles and subadults; the spring X2 effect may just be masked or weakened by changes in other habitat attributes. The relationships between these life stages and spring X2 clearly underwent downward shifts after each step decline. These persistent downward shifts indicate that occasional years with beneficial spring X2 conditions continue to have a positive effect on Delta Smelt, but they are by themselves not enough to overcome the depressed abundance levels and recover the population.

The downward shifts and changes in shape of the spring X2-Delta Smelt abundance index relationships (Fig. 23) also illustrate the difficulties of determining and understanding functional responses of biota to dynamic physical habitat attributes in changing ecosystems; the species of interest, other habitat attributes, and their interactions may all change as much or more than the habitat attribute under consideration. Further, these changes may not always be gradual, but can take the form of sudden step changes that may be associated with system-wide regime shifts (Davis et al. 2010, Baxter et al. 2010, Cloern and Jassby 2012). Moreover, prior conditions and prior abundance may also influence outcomes. In Chapter 9 of this report we give a relatively simple example of additional multivariate analyses aimed at exploring the effects of hydrology and prior abundance on the abundance and recruitment of Delta Smelt larvae. More sophisticated multivariate life cycle modeling that greatly exceeds the scope of this report is needed to account for these simultaneous changes and interactive effects on all life stages.

Changes in the size, location, and dynamics of the LSZ likely also interact in complex ways with other changes, such as changes in sediment and nutrient loadings and resulting turbidity and nutrient dynamics and their effects on Delta Smelt and the food web. For example, LSZ position affects recruitment of the invasive clam *Potamocorbula amurensis*, which may in turn affect phytoplankton and zooplankton biomass, size, and production (Thompson 2005, Winder and Jassby 2011), and has likely affected fish-X2 relationships (Kimmerer et al. 2002a).
Table 1. Summary of relationships between log-transformed annual abundance indices for four Delta Smelt life stages (response variable) and spring X2 (February-June, see text): Survey: see description of monitoring surveys in Chapter 3; Regression: least squares linear or quadratic regression: n, number of observations (years); P, statistical significance level for the model; R², coefficient of determination; adjusted R², R² adjusted for the number of predictor terms in the regression model. Bold font indicates statistically significant relationships.

<table>
<thead>
<tr>
<th>Life Stage</th>
<th>Season</th>
<th>Survey</th>
<th>Period</th>
<th>Regression</th>
<th>n</th>
<th>P</th>
<th>R²</th>
<th>Adjusted R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Juvenile</td>
<td>Summer</td>
<td>TNS</td>
<td>1959-2013</td>
<td>Linear</td>
<td>52</td>
<td>0.614</td>
<td>0.005</td>
<td></td>
</tr>
<tr>
<td>Juvenile</td>
<td>Summer</td>
<td>TNS</td>
<td>1959-1981</td>
<td>Linear</td>
<td>20</td>
<td>0.033</td>
<td>0.230</td>
<td>0.187</td>
</tr>
<tr>
<td>Juvenile</td>
<td>Summer</td>
<td>TNS</td>
<td>1959-1981</td>
<td>Quadratic</td>
<td>20</td>
<td>0.052</td>
<td>0.295</td>
<td>0.212</td>
</tr>
<tr>
<td>Juvenile</td>
<td>Summer</td>
<td>TNS</td>
<td>1982-2002</td>
<td>Linear</td>
<td>21</td>
<td>0.023</td>
<td>0.243</td>
<td>0.203</td>
</tr>
<tr>
<td>Juvenile</td>
<td>Summer</td>
<td>TNS</td>
<td>2002-2013</td>
<td>Linear</td>
<td>11</td>
<td>0.689</td>
<td>0.019</td>
<td></td>
</tr>
<tr>
<td>Subadult</td>
<td>Fall</td>
<td>FMWT</td>
<td>1968-2013</td>
<td>Linear</td>
<td>43</td>
<td>0.290</td>
<td>0.027</td>
<td>0.003</td>
</tr>
<tr>
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<td>Fall</td>
<td>FMWT</td>
<td>1968-1981</td>
<td>Linear</td>
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<td>0.699</td>
<td>0.017</td>
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<tr>
<td>Subadult</td>
<td>Fall</td>
<td>FMWT</td>
<td>1968-1981</td>
<td>Quadratic</td>
<td>11</td>
<td>0.295</td>
<td>0.263</td>
<td>0.079</td>
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<tr>
<td>Subadult</td>
<td>Fall</td>
<td>FMWT</td>
<td>1982-2002</td>
<td>Linear</td>
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<td>0.394</td>
<td>0.038</td>
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<tr>
<td>Subadult</td>
<td>Fall</td>
<td>FMWT</td>
<td>2002-2013</td>
<td>Linear</td>
<td>11</td>
<td>0.107</td>
<td>0.263</td>
<td>0.181</td>
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</tbody>
</table>

Ongoing studies coordinated by the IEP as part of the POD and FLaSH studies focus on the processes that link physics, chemistry, and biology in the LSZ and its habitat value for Delta Smelt and other native and non-native species. Similar to Monismith et al. (2002), preliminary results indicate that the strength of physical mixing (lateral dispersion) in the LSZ changes with the volume of freshwater outflow, underscoring the importance of variable hydrodynamics on not just the location of the LSZ, but how ecological services (nutrient mixing, organism dispersal) are influenced by variable estuarine outflow (Monismith, U.C. Berkeley, personal communication).

**Turbidity**

In this report, turbidity is considered an environmental driver that interacts with other environmental drivers, resulting in habitat attributes that directly affect Delta Smelt responses, rather than a stand-alone habitat attribute. Clearly, studies have shown that distribution of Delta Smelt is correlated with turbidity (e.g., Feyrer et al. 2007, Nobriga et al. 2008, Grimaldo et al. 2009, Sommer and Mejia 2013). In the conceptual model we chose to incorporate turbidity as a modifier of several important linkages between environmental drivers and habitat attributes that are important to Delta Smelt, primarily food visibility for small larvae and predation risk for all life stages. If we had incorporated turbidity as a habitat attribute and, for example, predation risk
was discussed separately from turbidity, there would have been a great deal of overlapping text between the two sections because turbidity interacts with the presence of predators to determine predation risk. Our approach is not ideal but should reduce redundant text and contribute to clarity of presentation. Nonetheless, we recognize that turbidity by itself could reasonably be considered as a habitat attribute. For example, it is possible that Delta Smelt experience stress in low turbidity habitat, which would in turn affect survival (likely through predation) but also in other direct ways such as lower growth and reduced egg production. However, we do not have evidence at this point to support that hypothesis.

In addition to salinity gradients, estuaries often have turbidity gradients. Turbidity is an optical property of water, which is the loss of transparency due to scattering of light by suspended particles. Typically, the upper reaches of estuaries have areas with high levels of suspended particles known as “estuarine turbidity maxima.” In many estuaries, these areas are located in or near the low salinity zone and are associated with higher numbers and enhanced growth for larvae of some species (Sirois and Dodson 2000a, b, Shoji et al. 2005). In the SFE, turbidity is largely determined by the amount of suspended inorganic sediment in the water (Cloern 1987, Ganju et al. 2007, Schoellhamer et al. 2012), although organic components can also play a role (USGS 2008). Sediment particles are constantly deposited, eroded, and resuspended, and are transported into, within, and out of the estuary. The amount of sediment that is suspended in the water column depends on the available hydrodynamic energy, which determines transport capacity, and on the supply of erodible sediment in the estuary and suspended sediments from the watershed.

In the upper SFE there are two main physical processes controlling turbidity. Suspended sediment is transported from the tributary watersheds into the system during high flows associated with winter and spring storm runoff (Schoellhamer et al. 2012). The first large storm of the rainy season often carries the highest concentrations of suspended sediment. Some portion of the transported sediment moves through the system to San Pablo and San Francisco Bay and the remainder is stored within the system as bottom sediment. During the remainder of the year, turbidity is primarily caused by interactions of this stored sediment with other environmental drivers (Schoellhamer et al. 2012). Water moving with the tides can resuspend fine sediments because of turbulence resulting from interactions between the bottom and water moving at high tidal velocities. At a larger scale, irregularities in the bottom topography may define geographic regions of greater turbulence and greater turbidity. In the upper estuary, such regions occur at a large bathymetric sill between Carquinez Strait and Suisun Bay and at another location within Suisun Bay (Schoellhamer 2001). Sediments may also be resuspended by turbulence related to wind waves. This process is mainly limited to areas with fine sediments on relatively shallow shoals where wind wave turbulence reaches the bottom. This process is most important in the shallows of Suisun, Grizzly, and Honker Bays and Liberty Island (Ruhl and Schoellhamer 2004, Warner et al. 2004, Morgan-King and Schoellhamer 2013). Thus, turbidity at any particular location is the result of several environmental drivers, including hydrology (transport from the watershed) and weather (wind and precipitation) interacting with the physical configuration of the upper SFE. Further, annual variation in these factors may have important effects. For example, during a drought there is little transport of suspended sediment and the same wind patterns during the summer may result in less turbidity than would occur after a wet year because less sediment was stored as benthic sediment during the winter. There is also evidence of longer term changes in turbidity (Schoellhamer et al. 2011, Hestir et al. 2013), along with regional differences.

In addition to the inorganic component of turbidity, organic matter (e.g., phytoplankton) also contributes to both suspended solids and the sediment load on the bed that is re-suspended with
wind and wave action (McGann et al. 2013). In the SFE, phytoplankton concentration varies spatially, seasonally, and on an inter-annual scale (Cloern et al. 1985, Jassby 2008, Cloern and Jassby 2012), and is controlled by multiple factors, including benthic grazing, climate, river inflows (Jassby et al. 2002), and nutrient dynamics (Glibert et al. 2011, Parker et al. 2012, Dugdale et al. 2013), which in turn are likely to affect the organic component of turbidity. Phytoplankton dynamics are discussed in detail in the ‘Food and Feeding’ section (below), but it is important to note here that plankton concentration comprises part of the SFE turbidity and is significant as it relates to productivity at higher trophic levels.

Among the geographic regions of the upper SFE, the Suisun region is one of the most turbid, when the system is not being influenced by storm flows. This results from strong turbulent hydrodynamics in the Suisun region caused by strongly interacting tidal and riverine flows, bathymetric complexity, and high wind speeds, which create waves that resuspend erodible benthic sediment in the large and open shallow bays of the Suisun region. The North Delta, especially the large open expanse of Liberty Island (flooded since 1998) and the adjacent Cache Slough region are also relatively turbid. Recent evidence suggests that Liberty Island acts as a sediment sink in the winter and a sediment source for the surrounding Cache Slough complex in the summer (Morgan-King and Schoellhamer 2013).

Turbidity is usually lower in the channels of the confluence of the Sacramento and San Joaquin Rivers compared to the Suisun region and North Delta region. Turbidity dynamics in the deep channels of the river confluence are driven more by riverine and tidal processes while high wind and associated sediment resuspension has little if any effect (Ruhl and Schoellhamer 2004). Turbidity is generally lowest in the south Delta (Nobriga et al. 2008). This may in part be due to sediment trapping by large, dense beds of *Egeria densa*, an invasive species of submerged aquatic vegetation (Hestir 2010). In winter/spring during the comparison years the highest Secchi disc depths (lowest turbidity) were found in the freshwater regions of the estuary (< 1 salinity), except for the Cache Slough region in the north Delta which was as turbid as the saltier regions of the estuary (Fig. 24).

There is strong evidence for an initial increase followed by a more recent long-term decline in sediment transport into the upper estuary, likely due to anthropogenic activities during the last century and a half (Schoellhamer et al. 2013, Wright and Schoellhamer 2004). Schoellhamer et al. (2013) presented a conceptual model of the effects of human activities on the sediment supplies in the SFE with four successive regimes:

1. The natural state.
2. Increasing sediment supplies due to mining, deforestation, agricultural expansion, etc.
3. Decreasing sediment supply due to sediment flushing during high flow events and sediment trapping behind dams and dikes.
4. A new altered state of low sediment supplies. The pulse of increased sediment inputs during and after the California gold rush and the more recent decline in these inputs is apparent in isotopic data from sediment cores taken in the estuary (Drexler et al. 2014).

The recent declines in sediment supplies have led to a long-term increase in water clarity in the upper Estuary (Jassby et al. 2002, Feyrer et al. 2007, Jassby 2008). Jassby et al. (2002) documented a 50% decrease in total suspended-solids concentration (TSS, a laboratory measurement of total suspended solids), approximated by suspended sediment concentration
(SSC, an optical measurement done in the field for these data) in the Delta from 1975-1995. Jassby (2008) found that the downward trend continued in the decade after 1995, although at a slower pace than over the entire 1975-2005. From 1975-2005, there were significant declines in SSC of up to 6% per year at 8 of 10 Delta stations (Jassby 2008). Jassby et al. (2005) showed that TSS concentrations in the north Delta dropped sharply toward the end of the 1982–1983 El Niño-Southern Oscillation (ENSO) event, which was associated with extremely high outflows, and did not recover afterward. This step decrease after 1983 has been corroborated by further trend analyses of TSS (Hestir 2013). Following the El Niño event of 1997–1998, there was a 36% step decrease in SSC in San Francisco Bay as the threshold from transport to supply regulation was crossed as an anthropogenic erodible sediment pool was depleted (Schoellhamer 2011). Sediment trapping by dense beds of *Egeria densa* may be further reducing available sediment in the Delta (Hestir 2010). While other anthropogenic factors may have also contributed to long-term changes in turbidity (e.g., export operations; Arthur et al. 1996), quantitative analyses of the effects of these factors have not been conducted.
Before the step decline in SSC and the onset of the pelagic organism decline in the late 1990s and early 2000s (i.e. the “pre-POD” period), water transparency (roughly the opposite of turbidity) measured with a Secchi disc at all IEP EMP stations was usually highest in November and lowest in June (Fig. 25). From 2003-2012 (i.e. the “POD” period), average water transparency was not only higher (by an average of 16 cm Secchi depth) than in the previous period, but the annual dynamics also shifted forward by a month, to greatest transparency (i.e. lowest turbidity) in October and lowest transparency in May. The greatest differences in average water transparency between the pre-POD and POD periods occurred in September and October (28 and 30 cm difference between monthly averages, respectively) and the smallest differences in January-May (10 cm). While the EMP has collected turbidity data (nephelometric turbidity (NTU) measurements) since 1975, long-term fish monitoring surveys have traditionally collected Secchi disc data and only in recent years have incorporated turbidity. Therefore, Secchi disc data are presented in the majority of this report when relating Delta Smelt abundance to water clarity conditions.

Multiple field and modeling studies have established the association between elevated turbidity and the occurrence and abundance of Delta Smelt. The abundance of larval/postlarval Delta Smelt larvae was well explained by salinity and Secchi depth, a proxy for turbidity (Kimmerer et al. 2009). Sommer and Mejia (2013) and Nobriga et al. (2008) found that late-larval and juvenile Delta Smelt are strongly associated with turbid water, a pattern that continues through fall (Feyrer et al. 2007). Long term declines in turbidity may also be a key reason that juvenile Delta Smelt now rarely occur in the south Delta during summer (Nobriga et al. 2008). Thomson et al. (2010) found that turbidity (water clarity) was the only significant predictor variable that was shared by three of the four POD species; all other significant predictor variables were unique to each species. Grimaldo et al. (2009) found that the occurrence of adult Delta Smelt at the fish salvage facilities was linked, in part, with high turbidity associated with winter “first flush” events. Turbidity may also serve as a behavioral cue for small-scale (lateral and vertical movements in the water column) and larger-scale (migratory) Delta Smelt movements (Bennett and Burau 2014).

Delta Smelt are visual feeders, and feed primarily between dawn and dusk (Hobbs et al. 2006, Slater and Baxter 2014). As for all visual feeders, visual range and prey density determine feeding success of Delta Smelt. Visual range depends on size, contrast and mobility of the prey, retinal sensitivity and eye size of the visual feeder, and on the optical habitat attributes such as light scattering, absorption, and intensity (Aksnes and Giske 1993). Optical habitat attributes are affected by turbidity from suspended organic particles, such as algae and detritus, and inorganic particles, such as sand and silt. Somewhat counterintuitively, some level of turbidity appears important to the feeding success of larval Delta Smelt. Baskerville-Bridges et al. (2004a) conducted laboratory experiments in which alga densities (0, 0.5 x 10^6 cell/mL, and 2 x 10^6 cell/mL or 1, 3, and 11 NTU) and light levels (range tested: 0.01 μmoles/s x m^2, 0.3 μmoles/s x m^2, 1.9 μmoles/s x m^2) were manipulated and first-feeding success of larval Delta Smelt was quantified. They found that maximum feeding response occurred at the highest alga concentrations and light levels tested. In a subsequent experiment, when alga were removed entirely, the feeding response was very low. The addition of algae or some other form of suspended particle is standard practice for successfully rearing Delta Smelt larvae in culture facilities (Mager et al. 2004, Baskerville-Bridges et al. 2005, Werner et al. 2010b, Lindberg et al. 2013). Presumably the suspended particles provide a background of stationary particles that helps the larvae detect moving prey. Sufficient turbidity also appears to be important to reduce overall environmental stress and increase survival of larval Delta Smelt (Lindberg et al. 2013). Thus, it seems likely that turbidity is important to the feeding success and survival of larval Delta Smelt.
Smelt in the wild. Recent research on juvenile Delta Smelt, however, suggests that influence of turbidity on feeding success may vary across life stages and field conditions. Hasenbein et al. (2013) exposed juveniles to varying turbidities (5-250 NTU) and observed a negative relationship between turbidity and feeding rates, with a marked decline in feeding at 250 NTU. However, feeding rates were highest at 12 NTU and stable in the 12-120 NTU turbidity range, which is likely within the range experienced by juvenile Delta Smelt in typical summer conditions in the Delta. Turbidity values of 250 NTU are generally not observed during the summer; therefore, the typical summer turbidity range in the Delta likely does not limit juvenile feeding success.

In addition to its effects on feeding, turbidity may also reduce predation risk. Based on the general recognition that fish assemblages are often partitioned between turbid-water and clear-water assemblages (Rodríguez and Lewis 1997, Whitfield 1999, Quist et al. 2004), and that turbidity can influence the predation rate on turbid-adapted fishes (Rodríguez and Lewis 1997, Gregory and Levings 1998, Quist et al. 2004), it has generally been assumed that juvenile and adult Delta Smelt are closely associated with turbidity in order to minimize their risk of predation in their generally open-water habitat. There may also be complex interactions between feeding and predation risk that are mediated by turbidity. Recent laboratory work has shown that in light (as opposed to dark) conditions, the vertical distribution of larval Delta Smelt shifts upward in the water column when turbidity is increased from clear (< 2 NTU) to 24 NTU (L. Sullivan, San Francisco State University, unpublished data), suggesting that larval Delta Smelt may use turbidity to safely forage in surface waters that may be more food-rich. Interestingly, when a predator cue (water, after containing juvenile Striped Bass for 1 hr) is added to clear water, the distribution of larval Delta Smelt becomes bimodal, with increased densities near the surface and

![Figure 25. Average and median Secchi depth in cm from monthly sampling at IEP Environmental Monitoring Program stations. Data are shown for the time period up to the pelagic organism decline (1975-2002) and after the decline (2003-2012).](chart.png)
closer to the bottom (L. Sullivan, San Francisco State University, unpublished data). Thus, while laboratory studies have demonstrated that larvae have improved feeding success at higher (but not too high, see above) turbidities, in natural settings, turbidity and predation risk may interact (e.g., Miner and Stein 1996) to affect Delta Smelt habitat choice and feeding success.

Turbidity may also be a migration cue for Delta Smelt. A recent field study investigated behavioral responses of Delta Smelt to winter “first flush” events in the Sacramento and San Joaquin Rivers near their confluence (W. Bennett, U.C. Davis, unpublished data). A first flush is defined as an increase in flow and turbidity associated with the onset of winter rain. This study found lateral turbidity gradients that changed with the tides and before and after first flush events and coincided with lateral Delta Smelt movements toward the channel during flood tides and toward the shoreline during ebb tides. The researchers concluded that this behavior likely facilitates maintaining channel position or moving upriver and cross-channel gradients in water turbidity may act as a behavioral cue. Feyrer et al. (2013) also found small-scale lateral and vertical gradients in turbidity in the lower Sacramento River just prior to a winter-time first flush event. In their study, turbidity and salinity were highest in the lower half of the water column and during flood tides and lowest during ebb tides in the center of the channel in the upper half of the water column. This coincided with observations of Delta Smelt which were more frequently caught throughout the water column during flood tides than during ebb tides when they were observed only in the lower half of the water column and sides of the channel. Feyrer et al. (2013) concluded that Delta Smelt may actively move in the water column by keying in on turbidity and salinity gradients or because of the physics underlying them.

**Entrainment and Transport**

The egg, larval, and juvenile stages of estuarine fishes and invertebrates along with small and weakly swimming adult stages are subject to involuntary transport (advection) by riverine and tidal flows. Entrainment is a specific case of involuntary transport. It refers to situations when altered flows misdirect and transport fish and other organisms in directions in which they would not normally travel or where they will encounter unfavorable conditions and increased risk of mortality. In this report, we use the term entrainment to specifically refer to the incidental removal of fishes and other organisms in water diverted from the estuary, primarily by CVP and SWP export pumping (Arthur et al. 1996, Grimaldo et al. 2009, Castillo et al. 2012).

Ultimately, watershed hydrology determines how much water can flow into and through the Delta; however, water flows into, within, and out of the Delta are manipulated in many ways. Water is: routed through and around artificial channels, gates, and barriers; stored in and released from reservoirs; discharged from agricultural and urban drains; and diverted with large and small pumps. Perhaps the greatest flow alterations in the Delta have taken place in Old and Middle Rivers (collectively referred to as “OMR”) in the central Delta (Fig. 2). Historically, these river channels were part of the tidal distributary channel network of the San Joaquin River (Whipple et al. 2012). Today, they are a central component of the CVP and SWP water conveyance system through the Delta. Water from the Sacramento River in the north now flows through the northern Delta (down Georgiana Slough, through Three-Mile Slough and around Sherman Island) and eastern Delta (via the artificial “Delta cross-channel” and down the forks of the Mokelumne River) to OMR in the central Delta, then to the SWP and CVP. The SWP and CVP pumps are capable of pumping water at rates sufficient to cause the loss of ebb tide flows and to cause negative net flows (the advective component of flow after removal of the diffusive tidal flow component) through OMR toward the pumps (see Grimaldo et al. 2009), thus greatly altering regional hydrodynamics and water quality (Monsen et al. 2007). Under these conditions, fish
and other aquatic species in the Delta may be transported toward the pumps (Arthur et al. 1996, Brown et al. 1996, Moyle et al. 2010), may swim toward the pumps if they are behaviorally inclined to follow net flow (Grimaldo et al. 2009), or may move toward the pumps if they are employing tidal surfing behavior (Sommer et al. 2011).

The SWP and CVP have large fish salvage facilities intended to reduce fish loss from the system due to entrainment - the State Skinner Fish Protective Facility (SFPF) and the federal Tracy Fish Collection Facility (TFCF). The SFPF and TFCF are located at the intakes to the State and federal export pumps on Old River in the southwestern Delta (Fig. 2). Both facilities have fish directing louvers and collecting screens that are used to capture and collect fish before they reach the pumps. The “salvaged” fish are then trucked to and released back into the western Delta. A variable fraction of these fish survive the capture, handling, trucking and release process (Miranda et al. 2010a,b, Aasen 2013, Afentoulis et al. 2013, Morinaka 2013a). The number of salvaged fish is monitored and reported as an index of SWP and CVP salvage and entrainment losses (Morinaka 2013b, more information and data available at http://www.dfg.ca.gov/delta/apps/salvage/Default.aspx). The SWP differs from the CVP in having a regulating reservoir, Clifton Court Forebay that temporarily stores water from Old River to improve operations of the SWP pumps. A change in the location of SWP water diversion from Italian Slough to Old River through CCF in 1969 may have led to a substantial increase in pre-screen losses at the SWP (Heubach ca. 1973, Kano 1990).

Fish have been salvaged since 1958 at the TFCF and since 1968 at SFPF, and the quality of the historical salvage data has improved over time. Delta Smelt salvage data is available since May 1979 for both the TFCF and SFPF (ftp://ftp.delta.dfg.ca.gov/salvage/). Juveniles less than 30 mm fork length are less efficiently captured in the salvage facilities (Kimmerer 2008, Morinaka 2013a) and Delta Smelt larvae less than 20 mm fork length have not been reported in the salvage data, although entrainment losses of Delta Smelt larvae have been calculated to be substantial under some circumstances (Kimmerer 2008). Development of a quantitative monitoring methodology for entrained Delta Smelt larvae at the CVP and SWP was recognized as necessary to refine triggers for protective actions (USFWS 2008). The current methodology for monitoring larval Delta Smelt at the TFCF and SFPF has provided presence-absence data since 2008 (Morinaka 2013b). Improved methods for sampling fish larvae have been reported at the TFCF (Reyes et al. 2012).

Despite these caveats salvage of Delta Smelt has been used as a rough index of entrainment losses. Delta Smelt salvage data since 1993 is considered more reliable than salvage data from earlier years. The difference in reliability is due to a change in count frequency from twice a day (0100 and 1300) from July 1978 to July 1992 to every two hours thereafter and an increased focus on proper identification of Delta Smelt following its State and federal listings as threatened (Morinaka 2013b).

Similar to the TNS and FMWT results for Delta Smelt, Delta Smelt salvage has declined dramatically since the beginning of this time series (Fig. 26). This is similar to trends for Chinook Salmon and Striped Bass salvage (not shown), but opposite to trends for Largemouth Bass and Bluegill (Lepomis macrochirus) salvage (Fig. 27), two species that may be benefiting from conditions resulting from an apparent ecological regime shift (Baxter et al. 2010). The ratio of Delta Smelt salvage divided by the previous year’s FMWT index has been used as a simple indicator of relative interannual entrainment losses. For adult (December-March) salvage, this ratio has been variable over time, but particularly high in the first three years of this time series (1980-1982, with 1982 being a wet year) and again during the beginning of a series of drought years in 1989 and in the fairly dry “POD” years 2003-2005 (Fig. 26). Current management
provisions to protect Delta Smelt (USFWS 2008) are aimed at keeping this ratio at no more than the average during the period of 2006-2008.

Delta Smelt were salvaged nearly year-round in the beginning of this time series. Delta Smelt salvage since 2005 has occurred mostly from January through June, with substantial decline of May-June juvenile salvage since the mid 2000s (Fig. 28) and virtual disappearance of older juveniles from July-August salvage since the year 2000 (Fig. 29) and subadults since the early 1990s (Fig. 30). These patterns coincide with the near disappearance of Delta Smelt from the central and southern Delta in the summer (Nobriga et al 2008) and in the south Delta in the fall (Feyrer et al. 2007). Historically, adult and larval-juvenile (> 20 mm FL) Delta Smelt salvaged were not separately recorded and reported, but based on length measurements of a subset of salvaged fish, adults were predominantly salvaged between December and March or April.
and most Delta Smelt larvae and juveniles were historically salvaged from April through July (Kimmerer 2008, Grimaldo et al. 2009).

Salvage data are routinely used to track and manage incidental take at the SWP and CVP and have been used to explore factors affecting entrainment and to estimate the effects of the SWP and CVP on Delta fishes. For example, Grimaldo et al. (2009) found that OMR flows and
It is important to remember, however, that salvage is only a very rough indicator of Delta Smelt entrainment. Based on mark-recapture experiments using cultured Delta Smelt, salvage was a very small fraction of total entrainment losses because of major pre-screen losses and low fish facility efficiency (Castillo et al. 2012). Experimental studies with cultured Chinook Salmon, Steelhead (*Oncorhynchus mykiss*), and Striped Bass have consistently shown that a large fraction (63% to 100%) of the entrained fish are not salvaged due to pre-screen losses and capture inefficiencies at the SWP fish facility (Brown et al. 1996, Gingras 1997, Clark et al. 2009). In addition, a mark–recapture test using field collected juvenile Chinook Salmon in CCF resulted in only 0.32% of the fish being salvaged (see Castillo et al. 2012). Pre-screen losses are generally turbidity account for much of the intra-annual variability in the salvage for juvenile and adult Delta Smelt.

**Figure 28.** A: Total reported May-June salvage for juvenile Delta Smelt and the corresponding mean salvage density based on the total monthly salvage and water volume exported by CVP and SWP. B: Both salvage and salvage density standardized by the Fall Midwater Trawl (FMWT) index for the previous year.
attributed to increased predation and other unfavorable habitat conditions near the SWP and CVP pumps (e.g. Kano 1990, Brown et al. 1996, Gringas and McGee 1997, Clark et al. 2009, Castillo et al. 2012). For juvenile and adult Delta Smelt, Castillo et al. (2012) found that 94.3% to 100% of marked fish groups released into the SWP CCF were never salvaged and that salvage of marked fish decreased as the distance from the release site to SFPF increased and as residence time in CCF increased.

Large pre-screen losses of Delta Smelt in CCF are likely due to increased predation, especially when Delta Smelt spend a relatively long time in the reservoir in the presence of predators. MacWilliams and Gross (2013) used a particle tracking model to estimate residence time of passive particles, which can be considered surrogates for weakly swimming Delta Smelt. In 21-

**Figure 29.** A: Total reported July-August salvage for juvenile Delta Smelt and the corresponding mean salvage density based on the total monthly salvage and water volume exported by CVP and SWP. B: Both salvage and salvage density standardized by the Fall Midwater Trawl (FMWT) index for the previous year.
day simulations with the three-dimensional (3D) hydrodynamic model UnTRIM, MacWilliams and Gross (2013) found that the time particles spend in CCF varies greatly with wind and SWP operating conditions. They estimated transit times for passive particles (e.g., larval Delta Smelt) from the radial gates to the SFPF of 4.3 days under moderate export conditions (average daily SWP export rate of 2,351 cfs) and 9.1 days under low export conditions (689 cfs). The CVP does not have a regulating reservoir in the Delta and CVP pre-screen losses in the river channels leading to the TFCF are likely different from SWP pre-screen losses, but there are no studies quantifying these differences.

In general, Delta Smelt salvage increases with increasing net OMR flow reversal (i.e., more negative net OMR flows) and when turbidity exceeds 10-12 NTU (USFWS 2008, Grimaldo et al. 2009). Based on field and salvage data, Kimmerer (2008) calculated that from near 0% to 25% of larval-juvenile and 0% to 50% of the adult Delta Smelt population can be entrained at

**Figure 30.** A: Total reported July-August salvage for sub-adult Delta Smelt and the corresponding mean salvage density based on the total monthly salvage and water volume exported by CVP and SWP. B: Both salvage and salvage density standardized by the Fall Midwater Trawl (FMWT) index for the same year.
the CVP and SWP annually, in years with periods of high exports. Although methods to calculate proportional loss estimates have since been debated (Kimmerer 2011, Miller 2011), a number of modeling efforts suggest that entrainment losses can adversely affect the Delta Smelt population (Kimmerer 2011, Maunder and Deriso 2011, Rose et al. 2013a, b).

High winter entrainment of Delta Smelt has been suspected as a contributing cause of both the early 1980s (Moyle et al. 1992) and the POD-era declines of Delta Smelt (Baxter et al. 2010). In addition to entraining Delta Smelt, water exports may likely also have indirect effects on Delta Smelt by contributing to adverse alterations of their habitat, for example, by changing Delta outflow and the size and location of the LSZ (see above) or by entraining food organisms (Jassby et al. 2002). The magnitude of these indirect effects of water exports on the Delta Smelt population has, however, not yet been quantified.

Delta Smelt are most vulnerable to entrainment when, as adults, they move from brackish water into fresh water, or as larvae, when they move from freshwater in the southern and central Delta into the brackish water of Suisun Bay. While some Delta Smelt live year-round in fresh water far from the CVP and SWP, most rear in the low-salinity regions of the estuary, also at a relatively safe distance from the SWP and CVP pumps. The timing, direction and geographic extent of the spawning movements of adult Delta Smelt affect their entrainment risk (Sweetnam 1999, Sommer et al. 2011a). Unlike the years prior to the 1990s, when high salvage of adult and juvenile Delta Smelt occurred at high, intermediate or low export levels, the risk of entrainment for fish that move into the central and south Delta is currently highest when net Delta outflow is at intermediate levels (~20,000 to 75,000 cfs) and OMR flow is more negative than -5000 cfs (USFWS 2008). In contrast, when adult Delta Smelt move upstream to the Sacramento River and into the Cache Slough region or do not move upstream at all, entrainment risk is appreciably lower. As explained later in this report, adult Delta Smelt may not move very far upstream during extreme wet years because the region of low salinity habitat becomes fresh and suitable for spawning (e.g., Suisun Bay or Napa River).

Transport mechanisms are most relevant to larval fishes, which have comparatively little ability to swim or otherwise affect their location. Dispersal from hatching areas to favorable nursery areas with sufficient food and low predation is generally considered one of the most important factors affecting the mortality of fish larvae (Hjort 1914, Hunter 1980, Anderson 1988, Leggett and Deblois 1994). Larvae of various smelt species exhibit diverse behaviors to reach and maintain favorable position within estuaries (Laprise and Dodson 1989, Bennett et al. 2002). Such nursery areas provide increased feeding success, growth rates and survival (Laprise and Dodson 1989, Sirois and Dodson 2000a, b, Peterson 2003, Hobbs et al. 2006). Until recently it was thought that larval Delta Smelt were transported from upstream hatching areas to downstream rearing areas, particularly the shallow productive waters of Suisun Bay (Moyle et al. 1992). Spring distributions of post-larval and small juvenile Delta Smelt support this view (Dege and Brown 2004). The distributions of these life stages were centered upstream of X2, but approached X2 as fish aged. These distributions could be displaced, and shifted up or down estuary with outflow and the shifting position of X2 (Dege and Brown 2004). More recent evidence suggests, however, that the timing and extent of downstream movement by young Delta Smelt is more variable than previously thought and that some may remain in upstream areas throughout the year (Sommer et al. 2011a, Contreras et al. 2011, Merz et al. 2011, Sommer and Mejia 2013).

Adult spawning site selection affects the potential importance of transport and entrainment to larvae. The risk of larval entrainment appears to increase with proximity to the south Delta export pumps (Kimmerer and Nobriga 2008). Larvae hatching in the San Joaquin River channel from
Big Break upstream to the city of Stockton and tidal channels south of these locations, can be affected by several interacting processes. Flows from the San Joaquin, Calaveras, Mokelumne and Cosumnes rivers act to cause net downstream flow, whereas export levels at the south Delta pumps act to reverse net flows in the lower San Joaquin River. High export rates can create negative flows past Jersey Point on the lower San Joaquin River (“Qwest,” see Dayflow documentation: http://www.water.ca.gov/dayflow/output/Output.cfm) and negative OMR flows (Fig. 31). Since the onset of the POD in 2002, positive average monthly OMR flows have only occurred in 9 months (6%) during the wettest years and average monthly Qwest flows were negative in just under half (49%) of all months (Fig. 31). Tidal conditions can also act in favor of downstream transport or entrainment depending upon whether the Delta is filling or draining in response to the fortnightly spring-neap cycle (Arthur et al. 1996). The combination of high export and low inflow can create very asymmetrical tides in OMR that covary with net negative flow resulting in stronger floods compared to ebbs, which may also contribute to fish entrainment.

**Predation Risk**

Small planktivorous fishes, including osmerids, serve as prey for larger fishes, birds and mammals. As prey, they have the critically important trophic function of transferring energy to higher trophic levels. Consequently, they are often subjected to intense predation pressure (Gleason and Bengsten 1996, Jung and Houde 2004, Halffredsson and Pedersen 2009). Prey fish populations compensate for high mortality through high reproductive rates, including strategies such as repeat spawning by individuals and rapid maturation (Winemiller and Rose 1992, Rose et al. 2001). Predation can be a dominant source of mortality for fish larvae, along with starvation and dispersion to inhospitable habitats (Hjort 1914, Hunter 1980, Anderson 1988, Leggett and Deblois 1994).

Since predation is a natural part of functional aquatic ecosystems, predators are likely not responsible for long-term declines in populations of prey fishes, such as Delta Smelt, without some additional sources of stress that disrupt the predator-prey relationship (Nobriga et al. 2013). Predation may become an issue when established predator-prey relationships are disrupted by habitat change or species invasions (Kitchell et al. 1994). As described in Chapter 1, the SFE has been extensively modified (Nichols et al. 1986, Cohen and Carlton 1998, Whipple et al. 2012, Cloern and Jassby 2012) so disrupted relationships between predators and prey are certainly plausible. For example, prey may be more susceptible to predation if they are weakened by disease, contaminants, poor water quality, or starvation. Similarly, the creation of more “ambush habitat” (e.g. structures, weed beds), declines in turbidity levels, or the introduction of a novel piscivore also may dramatically shift the existing predator-prey relationships (Ferrari et al. 2014).

All of these changes have in fact taken place in the estuary, especially in the central and south Delta (Feyrer and Healey 2003, Nobriga et al. 2005, Brown and Michniuk 2007).

Virtually all fishes of appropriate size will feed on fish larvae when available and predation is theoretically maximal when larve lengths are 10% of the length of the predator (Paradis et al. 1996). Presently, Mississippi Silverside (*Menidia audens*) is thought to be the most substantial predator of Delta Smelt larvae (Bennett and Moyle 1996, Bennett 2005, Baerwald et al. 2012). Juvenile and adult Delta Smelt have also been reported from the stomach contents of Striped Bass (Stevens 1963, Stevens 1966, Thomas 1967), White Catfish (*Ictalurus catus*) and Black Crappie (*Pomoxis nigromaculatus*) (Turner 1966a,b). Stevens (1963) reported “freshwater smelt” to be a very common component of Striped Bass stomach contents (nearly 100% frequency of occurrence in fifteen stomachs with food) on the Sacramento River near Paintersville Bridge.
during March-April 1963. During 1963-1964, Stevens (1966) also evaluated seasonal variation in the diets of juvenile Striped Bass throughout the Delta; only age 2 and age 3 Striped Bass contained more than trace amounts of Delta Smelt. The highest reported predation on Delta Smelt was 8% of the age 2 Striped Bass diet by volume during the summer. Thomas (1967) reported on spatial variation in Striped Bass diet composition based on collections throughout the SFE and the Sacramento River above tidal influence. The field collections occurred from 1957-1961; data were collected on age 1 and older Striped Bass but data were only summarized as all ages combined. Delta Smelt accounted for 8% of the spring diet composition and about 16% of the summer diet composition in the Delta.

Several authors tested hypotheses about inverse correlations between estimates of adult and juvenile Striped Bass abundance and indices of Delta Smelt relative abundance or survival (MacNally et al. 2010, Thomson et al. 2010, Maunder and Deriso 2011, Miller et al. 2012, Nobriga et al. 2013). None of these statistical analyses has found evidence for the expected inverse correlation. Modeling studies indicate that Striped Bass predation rates on prey are affected by temperature and predator abundance (mostly the latter; Loboschefsky et al. 2012). However, the links between prey abundance and predator abundance vary from strong to non-existent, depending on the strength of their interaction in the food web (Essington and Hansson 2004). It is not currently known if changes in juvenile Striped Bass abundance correspond with changes in population-level or per capita Striped Bass predation rate on Delta Smelt (Nobriga et al. 2013).

Recent modeling efforts show that Delta Smelt declines are negatively associated with metrics assumed to reflect the abundance of predators in the estuary (Maunder and Deriso 2011, Miller

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**Figure 31.** Flows in cubic feet per second for Qwest (positive values are seaward), Old and Middle River (OMR) (positive values are seaward), and total exports for years since the beginning of the pelagic organism decline (POD). Maximum monthly average Qwest values in 2006 and 2011 omitted to improve graph display. Values are 50,086 cfs in April 2006, 35,477 in May 2006, and 32,884 cfs in April 2011 (Qwest and Export data are from 2013 Dayflow, OMR data are from USGS).
et al. 2012). These metrics are composites of the relative abundance of Mississippi Silverside, Largemouth Bass and other centrarchids; species that are potential predators of concern because of their increasing abundance (Fig. 27; Bennett and Moyle 1996, Brown and Michniuk 2007, Thomson et al. 2010), and because of inverse correlations between Largemouth Bass abundance and Delta Smelt abundance (Nobriga and Feyrer 2007, Thomson et al. 2010, Maunder and Deriso 2011). These correlations could represent predation on Delta Smelt by Largemouth Bass, or alternatively, the very different responses of the two species to changing habitat within the Delta (Moyle and Bennett 2008). Current data suggest that Largemouth Bass populations have expanded as the SAV *Egeria densa* has expanded and have come to dominate parts of the Delta (Brown and Michniuk 2007). *E. densa* and Largemouth Bass are particularly prevalent in the central and southern Delta (Brown and Michniuk 2007) and Largemouth Bass may contribute to the pre-screen losses of Delta Smelt entrained into the SWP and CVP export pumps (see above). Largemouth Bass will readily eat Delta Smelt when the opportunity exists (Ferrari et al. 2014). However, there is little evidence that Largemouth Bass are major consumers of Delta Smelt due to low spatial co-occurrence (Nobriga et al. 2005, Baxter et al. 2010; L. Conrad, California Department of Water Resources, unpublished data). Thus, the inverse correlations between these species may not be mechanistic. Rather, they may reflect adaptation to, and selection for, different environmental conditions.

As noted above, predation on fish larvae can also be an important source of mortality. Juvenile and small adult fishes of many species will consume fish larvae when they are available. Major predators of the eggs and larvae of nearshore coastal and pelagic estuarine forage fishes can include invertebrates (DeBlois and Leggett 1993) and numerous small fishes not typically thought of as “piscivorous” (Johnson and Dropkin 1992), including adults of their own species (Takasuka et al. 2003). Bennett and Moyle (1996) and Bennett (2005) noted this and specifically identified Mississippi Silversides (hereafter, Silversides) as potential predators on Delta Smelt larvae. These authors also documented increases in the Silverside population from the mid-1970s through 2002. Consumption of Delta Smelt larvae by Silversides in the Delta was recently verified using DNA techniques (Baerwald et al. 2012). Larval predation is discussed in more detail in the next Chapter.

**Contaminants**

Fish are particularly sensitive to alterations in the chemical composition of the natural aquatic environment, as these changes can have significant impacts on their behavioral and physiological systems (Radhaiah et al. 1987). Chemical alterations can be the result of natural processes, for example the changes in local water quality associated with tidal water movements or natural biogeochemical processes, or they can be caused by pollution from watershed- or land-based sources of nutrients, such as nitrogen compounds, and contaminants, such as pesticides, metals, and contaminants of emerging concerns (CECs). The movement of contaminants through aquatic ecosystems is complex and dynamic, and many contaminants are difficult to detect and expensive to monitor (Scholz et al. 2012).

Portions of the SFE are listed as “impaired” on California’s 303(d) list of Impaired Water Bodies due to metals, pesticides, legacy pollutants, and nutrients that exceed established water quality objectives (SWRCB 2010). In particular, the entire SFE has been listed as impaired due to pollution with metals, such as mercury and selenium, and pesticides such as chlorpyrifos, DDT (Dichlorodiphenyltrichloroethane), and diazinon. The entire Delta, but not the bays of the SFE, is also listed for observed toxicity to aquatic organisms. In addition, the Stockton Ship Channel
in the southeastern Delta is listed for enrichment with nutrients, organic compounds, and low dissolved oxygen levels; Old River in the south-central Delta is listed for elevated salinity (electrical conductivity; EC) and total dissolved solids (TDS). Delta Smelt are likely exposed to a variety of these contaminants throughout their life cycle; however, the frequency and magnitude of the effects of contaminants on Delta Smelt health and reproduction are not very well understood in the SFE (Johnson et al. 2010, Brooks et al. 2012). The following sections describe the potential effects of key contaminants on Delta Smelt.

**Pesticides**

Pesticides produce many physiological and biochemical changes in freshwater organisms through their influence on the activities of several enzymes (Khan and Law 2005). Specifically, pesticides can have an adverse effect on hormones or other chemical messengers important to the health of an individual. Previous work has shown that chronic exposure to low levels of pesticides may even have a more adverse effect on fish than a single acute exposure to high levels. Chronic exposures were associated with changes in behavior and physiology that could influence survival and reproduction of wild fish (Ewing 1999). Biochemical and physiological stresses induced by exposure to pesticides can result in metabolic disturbances, retardation of growth, as well as reduction in longevity and fecundity (Murty 1986).

Pesticides are among the key contaminants believed to have contributed to the Delta Smelt decline (Johnson et al. 2010, Brooks et al. 2012, NRC 2012). Because pesticide concentrations in surface water are typically highest during the winter and spring, pesticides are most likely to affect the adult and larval life stages; however, effects may occur during any life stage as pesticides are seasonally and geographically widespread (Kuivila and Hladik 2008). Kuivila and Moon (2004) found that peak densities of larval and juvenile Delta Smelt sometimes coincided in time and space with elevated concentrations of dissolved pesticides in the spring. These periods of co-occurrence lasted for up to 2–3 weeks. While concentrations of individual pesticides were lower than would be expected to cause acute mortality, little is known of the sublethal effects of pesticides on Delta Smelt. Although little evidence exists for acute effects of pesticides on fish or invertebrates, several studies have documented sublethal effects on fish health (Werner et al. 2008, Werner et al. 2010a, Werner et al. 2010b).

Herbicides and fungicides were among the most commonly detected classes of pesticides observed in water and sediment in the Delta and are also found in fish tissue (Orlando et al. 2013, Smalling et al. 2013). Herbicides are known to affect primary producers, while insecticides can affect invertebrate prey species (e.g., Brander et al. 2009, Weston et al. 2012), which could lead to contaminant-mediated food limitation for Delta Smelt. Fungicides have been found to cause endocrine disruption in fish, including reduced fecundity (Ankley et al. 2005). Recent work has shown that the insecticide esfenvalerate affects swimming behavior of exposed larval Delta Smelt (Connon et al. 2009). It was also found to alter the expression of genes involved in neuromuscular activity and immune response, detoxification, and growth and development (Connon et al. 2009). Additionally, insecticides are known to affect predator-prey relationships for fish, as well as lead to endocrine disruptions (Scholz et al. 2000, Junges et al. 2010, Relyea and Edwards 2010, Riar et al 2013, Forsgren et al. 2013). Contamination of aquatic systems by pyrethroid insecticides was recently found to lead to genetic point mutations in the nontarget, aquatic amphipod *Hyalina azteca*, resulting in differences in pyrethroid sensitivity. Wild populations of *H. azteca* collected from areas with high sediment concentrations of pyrethroids exhibited remarkable resistance to pyrethroids compared to laboratory cultures and the observed
resistance was highly coupled to the presence of a genetic mutation. The LC50s (concentration that is lethal to 50% of the exposed population) of previously-exposed wild populations were up to two orders of magnitude greater than LC50s of laboratory cultures. Moreover, the presence of a genetic mutation was detected in 100% of *H. azteca* that survived exposure to high pyrethroid concentrations. The development of such resistance can result in costs to genetic and biological diversity, including reduced fitness, and may lead to impacts to the food web (Weston et al. 2013). The presence of such resistance and genetic mutations in Delta Smelt as a result of pyrethroids or other pesticide exposure has not been investigated.

It is also important to note that environmental factors such as temperature and salinity affect pesticide toxicity in fish (Coats et al. 1989, Lavado et al. 2009). For that reason, seasonal variation in environmental factors may result in greater risk to certain life stages. The results above are for dissolved pesticides; pesticides may also be bound to sediments, representing another possible mechanism of exposure. Pesticides, such as pyrethroids and organochlorines, that strongly bind to sediment may be particularly important to the adult and larval life stage of Delta Smelt as these life stages occur during the winter and spring, when rain events (including the “first flush”) transport sediment and associated contaminants into the Delta; however, as the mechanisms that influence the desorption rates of pesticides are complex (e.g., temperature, contact time, pesticide) (e.g., Xu et al. 2008, Cornelissen et al. 1998), exposure rates for Delta Smelt lifestages are likely multifaceted and difficult to predict.

**Ammonia and Ammonium**

Agricultural operations, wastewater treatment plant effluent, and other sources contribute to the accumulation of nutrients in the Delta. Nutrients, such as ammonium (a cation) and ammonia (its toxic, unionized form) are of particular concern in the Delta, as they can have significant negative effects on Delta Smelt and their habitat. Ammonium is increasingly converted into ammonia as pH rises. Delta Smelt spawning and larval nursery areas in the northern Delta are at particular risk to exposure to ammonia/um, mainly due to discharge by the Sacramento Regional Wastewater Treatment Plant (SRWTP) into the lower Sacramento River (Connon et al. 2011a). However, effects of nutrients such as ammonia/um are likely at all Delta Smelt life stages, as nutrients are discharged throughout the Delta year-round.

Recent work demonstrated that Delta Smelt exposed to ammonia exhibited membrane destabilization, which may lead to increased membrane permeability as well as increased susceptibility to synergistic effects of multi-contaminant exposures (Connon et al. 2011a, Hasenbein et al. 2013b); however, the concentrations of ammonia used in these studies were higher than the concentrations typically experienced by Delta Smelt in the wild. In other fish species, sublethal concentrations of ammonia/um have also led to histological effects such as gill lamellae fusions and deformities (Benli et al. 2008). Other work has also shown that neurological and muscular impacts of ammonia/um resulted in slowed escape response and subsequent mortality (McKenzie et al. 2008).

**Metals and Other Elements of Concern**

Historic mining sites, industrial and domestic wastewater discharges, and agricultural runoff are largely responsible for the presence of metals and other elements of concern in the Delta. Metals of particular importance in the Delta include copper and mercury; selenium is a trace element
of concern. Delta Smelt exposed to copper exhibited reduced swimming velocities and suffered
digestive and neurological effects (Connon et al. 2011b). Other sublethal effects on fish caused
by exposure to these elements include reduced fertility and growth, impaired neurological and
endocrine functions, and skeletal deformities that affect swimming performance (Boening 2000,
Chapman et al. 2010). These elements are often associated with sediment and may be particularly
important to the adult and larval life stages, since sediment is transported with significant rain
events, including the “first flush.”

**Contaminants of Emerging Concern**

Contaminants of emerging concern (CECs) such as pharmaceuticals, hormones, personal care
products, and industrial chemicals are of increasing concern because they are widespread
in the aquatic environment, biologically active, and are relatively unregulated (Kolpin et
al. 2002, Pal et al. 2010). The California State Water Resources Control Board is currently
investigating CECs in the Delta (http://www.sccwrp.org/ResearchAreas/Contaminants/
ContaminantsOfEmergingConcern/ EcosystemsAdvisoryPanel.aspx). CECs originate from many
sources including industrial and domestic wastewater. They are responsible for a myriad of
sublethal effects in fish including endocrine disruption, changes in gene transcription and protein
expression, and morphological and behavioral changes (Brander 2013). Though the effects of
CECs have been well studied in other fish species, the extent to which they influence Delta Smelt
remains unclear.

**Polycyclic Aromatic Hydrocarbons (PAHs)
and Polychlorinated Biphenyls (PCBs)**

The PAHs and PCBs found in the Delta are largely from urban and industrial sources. PAHs are
formed during the incomplete burning of coal, oil, gas, garbage, and other organic substances.
PCBs are synthetic organic chemicals that were used in many industrial and commercial
applications. PCBs were banned in 1979, but continue to persist in the environment. PAHs and
PCBs bind strongly to sediment and therefore are likely to be associated with the “first flush”
and may be particularly important to the adult and larval life stages of Delta Smelt. Almost all
sediments sampled in the Delta in 2006 contained PAHs (mean concentration of 0.3 parts per
million in Suisun Bay) and PCBs (mean concentration of 0.8 parts per million in Suisun Bay)
(SFEI 2007). Studies have found PAHs and PCBs in surface water, with concentrations in excess
of established water quality objectives (Thomson et al. 2000, Oros et al. 2006). Both PCBs and
PAHs can cause endocrine disruption in fish (Brar et al. 2010, Nicolas, 1999); however, specific
impacts on Delta Smelt have not been documented.

**Contaminant Mixtures**

While the individual effects of the aforementioned contaminants can be severe, recent work has
demonstrated that the interaction of the contaminants within mixtures can have both synergistic
and antagonistic effects, exacerbating potential impacts on fish physiology (e.g., Jordan et
al. 2012). There is increasing evidence that compounds in mixtures show adverse effects at
concentrations at which no effects were observed for single toxicants (e.g., Baas et al. 2009,
Silva et al. 2002, Walter et al. 2002). For example, recent work on Mississippi Silversides
has demonstrated that contaminant mixtures resulted in endocrine disruptions such as varied
expression of mRNA levels for estrogen-responsive genes, reduced mean gonadal somatic indices (GSI), testicular necrosis, and biased sex ratios (Brander et al. 2013). Studies have also shown that mixtures can affect predator-prey interactions (Relyea and Edwards 2010) and cause liver abnormalities (Sacramento Splittail, *Pogonichthys macrolepidotus*; Greenfield et al. 2008). Other work on Striped Bass has demonstrated that contaminant mixtures can be maternally-transferred to fish eggs, resulting in larvae with impaired growth and abnormal brain and liver development (Ostrach et al. 2008).

Due to the unpredictability of their effects on organisms, the synergistic effects of contaminant mixtures have received a great deal of attention both within pharmacology and environmental sciences (Arnold et al. 1996, Ashby et al. 1997, Berenbaum 1989, Greco et al. 1995, Liang and Lichtenstein 1974). Currently, one of the greatest challenges in chemical mixture research is how to deal with the infinite number of combinations of chemicals and other stressors, as well as their interactive effects, on organisms (Baas et al. 2010). Additional challenges also exist trying to relate lab-based findings to wild populations for studies examining the effects of individual contaminants and contaminant mixtures on organisms using exposure concentrations that are environmentally representative. Therefore, while the potential for exposure to contaminant mixtures in all Delta Smelt life stages is highly probable, any specific effects of such interactions on Delta Smelt remain unknown.

**Food and Feeding**

The presence of food is, obviously, a critical habitat attribute for any organism; however, the factors determining the quantity and quality of available food can be quite complex. In this section, we begin with a brief review of information about trophic processes in the upper SFE. We then discuss the available data on prey consumed by Delta Smelt. Finally, we provide a review of information on factors possibly affecting abundance and quality of food organisms.

Estuaries are commonly characterized as highly productive nursery areas for a suite of organisms. Productivity of estuarine ecosystems is often fueled by detritus-based food webs. In the SFE, much of the community metabolism in pelagic waters does result from microbial consumption of organic detritus. However, evidence suggests that metazoan production in pelagic waters is primarily driven by phytoplankton production (Sobczak et al. 2002, 2005, Mueller-Solger et al. 2002, 2006, Kimmerer et al. 2005). Protists (flagellates and ciliates) consume both microbial and phytoplankton prey (Murrell and Hollibaugh 1998, York et al. 2010) and are an additional important food source for many copepod species in the estuary (Rollwagen-Bollens and Penry 2003, Bouley and Kimmerer 2006, Gifford et al. 2007, McManus et al. 2008). However, the conversion of dissolved and particulate organic matter to microbial biomass and then to zooplankton is a relatively slow and inefficient process. Shifts in phytoplankton and microbial food resources for zooplankton might favor different zooplankton species. Moreover, phytoplankton production and biomass in the SFE is low compared to many other estuaries (e.g., Jassby et al. 2002, Kimmerer et al. 2005, Wilkerson et al. 2006, Cloern and Jassby 2012). The recognition that phytoplankton production might impose limits on pelagic fishes, such as Delta Smelt, through food availability has led to intense interest in factors affecting phytoplankton production and species composition and in management actions aimed at enhancing high-quality phytoplankton production. In addition, there is a major need to understand other trophic pathways given the observation that larger Delta Smelt periodically can take advantage of epibenthic prey (see below).
Phytoplankton biomass (measured as chlorophyll-\(a\)) has been routinely monitored in the estuary since the 1970s. The 1975-2012 median chlorophyll-\(a\) concentration across all IEP EMP stations is 2.8 \(\mu\)g/L (n = 13482, interquartile range (IQR) = 5 \(\mu\)g/L). Seasonally, the highest chlorophyll-\(a\) concentrations tend to be observed in May and June and the lowest concentrations in December and January (Fig. 32). Regionally, monitoring stations in the South Delta/San Joaquin River usually have the highest chlorophyll-\(a\) concentrations. There has been a well-documented long-term decline in phytoplankton biomass (chlorophyll-\(a\)) and primary productivity (estimated from measurements of chlorophyll-\(a\) and of water column light utilization efficiency) to very low levels in the Suisun Bay region and the lower Delta (Jassby et al. 2002). Jassby et al. (2002) detected a 47% decline in June–November chlorophyll-\(a\) and a 36% decline in June–November primary production between the periods 1975–1985 and 1986–1995. Jassby (2008) updated the phytoplankton analysis to include the more recent data (1996–2005) from the Delta and Suisun Bay. Jassby (2008) confirmed a long-term decline in chlorophyll-\(a\) from 1975 to 2005 but also found that March–September chlorophyll-\(a\) had an increasing trend in the Delta from 1996 to 2005. Suisun Bay did not exhibit any trend during 1996–2005. A similar pattern was noted for primary production in the Delta. These chlorophyll-\(a\) patterns continued to hold through 2008 according to a more recent study by Winder and Jassby (2011). In the most recent decade (2003-2012), the median chlorophyll-\(a\) concentration across all IEP EMP stations was 2 \(\mu\)g/L (n = 2620, IQR = 2 \(\mu\)g/L), compared to the 1975-2002 median chlorophyll-\(a\) concentration of 3 \(\mu\)g/L (n = 10862, IQR = 6 \(\mu\)g/L) (Fig. 32). Most of the decrease was due to declines during May-October and especially the near-elimination of the formerly common “spring bloom” of phytoplankton in May (Fig. 32). In summary, phytoplankton biomass and production in the Delta and Suisun Bay seem to have reached a low point by the end of the 1987–1994 drought. While they recovered somewhat in the Delta, chlorophyll-\(a\) stayed consistently low in Suisun Bay through the POD years.

**Figure 32.** Interquartile ranges (boxes) and medians (lines) for chlorophyll-\(a\) measured monthly at all IEP EMP stations from 1975-2002 (blue) and 2003-2012 (red). Data from [http://www.water.ca.gov/BDMA/](http://www.water.ca.gov/BDMA/).
A major reason for the long-term phytoplankton reduction in the upper SFE after 1985 is benthic grazing by the invasive overbite clam (*Potamocorbula amurensis* also known as *Corbula amurensis*) (Alpine and Cloern 1992), which became abundant by the late 1980s (Kimmerer 2002). The overbite clam was first reported from San Francisco Estuary in 1986 and it was well established by 1987 (Carlton et al. 1990). Prior to the overbite clam invasion, the invasive Asiatic freshwater clam (*Corbicula fluminea*) (introduced in the 1940s) colonized Suisun Bay during high flow periods and the estuarine clam *Mya arenaria* (also known as *Macoma balthica*, an earlier introduction) colonized Suisun Bay during prolonged (> 14 month) low flow periods (Nichols et al. 1990). Thus, there were periods of relatively low clam grazing rates while one species was dying back and the other was colonizing, resulting in neither reaching high abundances. The *P. amurensis* invasion changed this formerly dynamic clam assemblage because *P. amurensis*, which is tolerant of a wide range of salinity, can maintain large, permanent populations in the brackish water regions of the estuary. *P. amurensis* biomass and grazing usually increase from spring to fall which contributes to the reduction in phytoplankton biomass from May to October relative to historical levels. In addition, the grazing influence of *P. amurensis* extends into the freshwater Delta beyond the clam’s typical brackish salinity range, presumably due to tidal dispersion of phytoplankton-depleted water between regions of brackish water and fresh water (Kimmerer and Orsi 1996, Jassby et al. 2002).

Phytoplankton production in the SFE has been considered primarily light-limited because nutrient concentrations commonly exceed concentrations limiting primary production. According to some recent work, shifts in nutrient concentrations and ratios may, however, also contribute to the phytoplankton reduction and changes in algal species composition in the SFE. Nutrients may also play a larger role in regulating phytoplankton dynamics in the estuary as the estuary clears and light availability increases (see turbidity section above).

While phosphorus (total phosphorous and soluble reactive phosphorous) concentrations declined in the Delta and Suisun Bay region over the last few decades, nitrogen (total nitrogen and ammonium) concentrations increased. These changes have been attributed to the operation of the Sacramento Regional Wastewater Treatment Plant (SRWTP), a large secondary treatment facility that was completed in 1984 (VanNieuwenhuysen 2007, Jassby 2008). As stated previously, ammonia has two forms, un-ionized ammonia (NH$_3$) which is toxic to aquatic organisms and the ammonium ion (NH$_4^+$) which is considerably less toxic to animals and an important nutrient for plants and algae (Thurston et al. 1981). Ammonia exists in equilibrium between the two forms dependent primarily on the pH of the water, but also temperature, with increases in pH and temperature favoring the un-ionized form (Thurston et al. 1981). Dugdale et al. (2007) and Wilkerson et al. (2006) found that high ammonium concentrations prevented the formation of diatom blooms but stimulated flagellate blooms in the lower estuary. They propose that this occurs because diatoms preferentially utilize ammonium in their physiological processes even though it is used less efficiently and at high concentrations ammonium can prevent uptake of nitrate (Dugdale et al. 2007). Thus, diatom populations must consume available ammonium before nitrate, which supports higher growth rates, can be utilized or concentrations of ammonium need to be diluted. A recent independent review panel (Reed et al. 2014) found that there is good evidence for preferential uptake of ammonium and sequential uptake of first ammonium and then nitrate, but that a large amount of uncertainty remains regarding the growth rates on ammonium relative to nitrate and the role of ammonium in suppressing spring blooms.

Glibert (2012) analyzed long-term data (from 1975 or 1979 to 2006 depending on the variable considered) from the Delta and Suisun Bay and related changing forms and ratios of nutrients, particularly changes in ammonium, to declines in diatoms and increases in flagellates and
cyanobacteria. Similar shifts in species composition were noted by Brown (2009), with loss of
diatom species, such as *Thalassiosira sp.*, an important food for calanoid copepods, including
*Eurytemora affinis* and *Sinocalanus doerrii* (Orsi 1995). More recently, Parker et al. (2012)
found that the region where blooms are suppressed extends upstream into the Sacramento River
to the SRWTP, the source of the majority of the ammonium in the river (Jassby 2008). Parker
et al. (2012) found that at high ambient ammonium concentrations, river phytoplankton cannot
efficiently take up any form of nitrogen including ammonium, leading to often extremely low
biomass in the river. A study using multiple stable isotope tracers (Lehman et al. 2014) found
that the cyanobacteria *M. aeruginosa* utilized ammonium, not nitrate, as the primary source
of nitrogen in the central and western Delta. In 2009, the ammonia concentration in effluent
from SRWTP was reduced by approximately 10%, due to changes in operation (K. Ohlinger,
Sacramento Regional County Sanitation District, personal communication). In spring 2010
unusually strong spring diatom blooms were observed in Suisun Bay that co-occurred with low
ammonia concentrations (Dugdale et al. 2013).

Jassby (2008) suggested the following comprehensive explanation for his observations.
Phytoplankton production in the lower Delta is associated with flow and residence time; however,
other factors introduce a substantial degree of interannual variability. Benthic grazing by *C.
fluminea* is likely a major factor as grazing can exceed rates of primary production (Lucas et
al. 2002, Lopez et al. 2006) and are abundant year round at some locations in the Delta (Fuller
2012). Current data are inadequate to estimate the overall magnitude of the grazing effect of
*C. fluminea*. In Suisun Bay, benthic grazing by *P. amurensis* is a controlling factor that keeps
phytoplankton at low levels. Thus, metazoan populations in Suisun Bay are dependent on
importation of phytoplankton production from the upstream portions of the Delta. Upstream
Delta phytoplankton can be lost via exports and within-Delta depletion; Cloern and Jassby
(2012) reported phytoplankton losses equivalent to 30% of the primary production in the
Delta. Ammonium concentrations and water clarity have increased; however, these two factors
should have opposing effects on phytoplankton production. These factors likely also contribute
to variability in the interannual pattern but the relative importance of each is unknown. The
interactions among primary production, grazing, and transport time can be complex (Lucas et

The changes in phytoplankton production and invasion and establishment of the overbite clam *P.
amurensis* were also accompanied by a series of major changes in consumers (Winder and Jassby
2011). Many of these changes likely negatively influenced pelagic fish production, including
Delta Smelt. The quantity of food available to Delta Smelt is a function of several factors,
including but not limited to seasonal trends in prey abundance and prey species specific salinity
tolerances, which influence distribution (Kimmerer and Orsi 1996, Hennessy and Enderlein
2013). Seasonal peaks in abundance vary among calanoid copepods consumed as prey by Delta
Smelt, *E. affinis* in April-May (Fig. 33), *P. forbesi* in July (Fig. 34), and *A. sinensis* in Sep-Oct
(Fig. 35). Upstream, the calanoid copepod *S. doerrii* is most abundant May-June (Fig. 36). The
seasonal trend in cladocerans (Fig. 37) and mysid (Fig. 38) prey are similar, being most abundant
in summer.

From March through June, larval Delta Smelt rely heavily on first juvenile, then adult stages of
the calanoid copepods *Eurytemora affinis* and *Pseudodiaptomus forbesi*, as well as cladocerans
(Nobriga 2002, Hobbs et al. 2006, Slater and Baxter 2014), and *Sinocalanus doerrii* (Fig. 39).
Nobriga (2002) found that Delta Smelt larvae expressed positive selection for *E. affinis* and *P.
forbesi*, consuming these prey species in greater proportion than available in the environment.
Such selection was not noted for other zooplankton prey. Regional differences in food use occur,
with *E. affinis* and *P. forbesi* being major prey items downstream in the LSZ with a transition to *S. doerrii* and cyclopoid copepods as major prey items upstream into the Cache Slough-Sacramento River Deepwater Ship Channel (CS-SRDWSC) (Fig. 39).

Juvenile Delta Smelt (June-September) rely extensively on calanoid copepods such as *E. affinis* and *P. forbesi*, especially in freshwater (salinity < 1) and CS-SRDWSC but there is great variability among regions (figs. 40-43). Larger fish are also able to take advantage of mysids,
cladocerans, and amphipods (Moyle et al. 1992, Lott 1998, Feyrer et al. 2003, Steven Slater, California Department of Fish and Wildlife, unpublished data) (Figs. 34-37). The presence of several epibenthic species in diets therefore indicates that food sources for this species are not confined to pelagic pathways. Such food sources may be especially important in regions of the estuary where there is extensive shoal habitat such as Liberty Island (Steven Slater, California Department of Fish and Wildlife, unpublished data).

Figure 34. Density (number/m³) of adult *Pseudodiaptomus forbesi* (*P. forbesi*) by month for three salinity ranges. Each month 16 stations were sampled across all salinity ranges. Horizontal lines represent single samples within a salinity range and boxes without whiskers indicate 2 samples within a salinity range. Data from the IEP Zooplankton Study index stations. See Chapter 3: Data Analyses for explanation of boxplots.
Subadult Delta Smelt (September through December) prey items are very similar to those of juvenile Delta Smelt but with increased variability in diet composition (Moyle et al. 1992, Lott 1998, Steven Slater, California Department of Fish and Wildlife, unpublished data) (Figs. 40-43) coinciding with the seasonal decline in pelagic zooplankton, such as *P. forbesi* (Fig. 34) and mysids (Fig. 38). Food habits of adult Delta Smelt during the winter and spring (January-May) have been less well documented (Moyle et al. 1992). In 2012, diet of adults in the LSZ and
< 1 ppt were found to include cyclopoid copepods, other than *Limnoithona* spp., with a mix of larger prey types, amphipods, cladocerans, cumaceans, and larval fish and in CS-SRDWSC the calanoid copepod *S. doerrii* continued to be a large portion of the diet (Steven Slater, California Department of Fish and Wildlife, unpublished data) (Fig. 44). Larval fish found in stomachs of Delta Smelt in the higher salinity areas were primarily Pacific Herring (*Clupea pallasii*), with

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**Figure 37.** Density (number/m³) of all cladoceran taxa by month. Each month 16 stations were sampled across all salinity ranges. Horizontal lines represent single samples within a salinity range and boxes without whiskers indicate 2 samples within a salinity range. Data from the IEP Zooplankton Study index stations. See Chapter 3: Data Analyses for explanation of boxplots.

**Figure 38.** Density (number/m³) of all mysid shrimp taxa by month. Each month 16 stations were sampled across all salinity ranges. Horizontal lines represent single samples within a salinity range and boxes without whiskers indicate 2 samples within a salinity range. Data from the IEP Zooplankton Study index stations. See Chapter 3: Data Analyses for explanation of boxplots.
some Longfin Smelt, and Prickly Sculpin (Cottus asper) in the Sacramento River and CS-SRDWSC region; no Delta Smelt larvae were found in the stomachs of adults (Steven Slater, California Department of Fish and Wildlife, unpublished data).

The large proportion of benthic amphipods, cumaceans, and some cladocerans (Camptocercus spp.) in the diet is a notable change from Delta Smelt diet in the 1970s. Delta Smelt diets historically did include amphipods, notably Corophium spp. (Moyle et al. 1992), yet it was a small fraction of a mostly pelagic based diet. The considerable use of benthic invertebrates for food in recent years is believed to be in large part due to food limitation associated with the long-term decline and changes in composition of the pelagic food web (Slater and Baxter 2014). The quality of benthic invertebrates as food is not currently understood, but amphipods are lower in energy (calories per gram) than copepods (Cummins and Wuychek 1971, Davis 1993) and mysids (Davis 1993).

As noted previously, the changes in phytoplankton production and phytoplankton species abundances observed and the invasion of P. amurensis may have had important consequences for consumer species preyed upon by Delta Smelt. For example, there has been a decrease in mean zooplankton size (Winder and Jassby 2011) and a long-term decline in calanoid copepods, including a major step-decline in the abundance of the copepod E. affinis. These changes are possibly due to predation by the overbite clam (Kimmerer et al. 1994) or indirect effects of clam grazing on copepod food supply. Predation by P. amurensis may also have been important for other zooplankton species (Kimmerer 2008). Northern Anchovy Engraulis mordax abandoned the low salinity zone coincident with the P. amurensis invasion, presumably because the clam reduced planktonic food abundance to the point that occupation of the low-salinity waters was no longer energetically efficient for this marine fish (Kimmerer 2006). Similarly, Longfin Smelt Spirinchus thaleichthys shifted its distribution toward higher salinity in the early 1990s, also presumably because of reduced pelagic food in the upper estuary (Fish et al. 2009). There was also a decline in mysid shrimp (Winder and Jassby 2011), including a major step-decline in 1987–1988, likely due to competition with the overbite clam for phytoplankton (Orsi and Mecum 1996). Mysid shrimp had been an extremely important food item for larger fishes like Longfin Smelt and juvenile Striped Bass (Orsi and Mecum 1996), and may be consumed by larger Delta Smelt (Moyle et al. 1992). The decline in mysids was associated with substantial changes in the diet composition of these and other fishes, including Delta Smelt (Feyrer et al. 2003, Bryant and Arnold 2007). The population responses of Longfin Smelt and juvenile Striped Bass to winter–spring outflows changed after the P. amurensis invasion. Longfin Smelt relative abundance was lower per unit outflow after the overbite clam became established (Kimmerer 2002b). Age-0 Striped Bass relative abundance stopped responding to outflow altogether (Sommer et al. 2007). One hypothesis to explain these changes in fish population dynamics is that lower prey abundance reduced the system carrying capacity (Kimmerer et al. 2000, Sommer et al. 2007).

In addition to a long-term decline in calanoid copepods and mysids in the upper Estuary, there have been numerous copepod species introductions (Winder and Jassby 2011). P. forbesi, a calanoid copepod that was first observed in the estuary in the late 1980s, has replaced E. affinis as the most common Delta Smelt prey during the summer. It may have a competitive advantage over E. affinis due to its more selective feeding ability. Selective feeding may allow P. forbesi to utilize the remaining high-quality algae in the system while avoiding increasingly more prevalent low-quality and potentially toxic food items such as M. aeruginosa (Mueller-Solger et al. 2006, Ger et al. 2010a). After an initial rapid increase in abundance, P. forbesi declined somewhat in abundance from the early 1990s in the Suisun Bay and Suisun Marsh regions but maintained its abundance, with some variability, in the central and southern Delta (Winder and Jassby 2011).
Figure 39. Percentage by weight of prey types found in the digestive tracts of larval and young juvenile Delta Smelt (≤ 20 mm fork length) collected from 1-6 ppt, < 1 ppt, and Cache Slough-Sacramento River Deepwater Ship Channel (CS-SRDWSC) in A) 2005, B) 2006, C) 2010, and D) 2011. Number of digestive tracts examined are shown above the columns. Mean fork length (mm) of Delta Smelt is also shown.
Figure 40. Percentage by weight of prey types found in stomachs of age-0 Delta Smelt collected from > 6 ppt during April through December in A) 2005, B) 2006, C) 2010, and D) 2011. Number of stomachs examined are shown above the columns. One fish examined in August 2006 had an empty stomach. Mean fork length (mm) of Delta Smelt is also shown.
Figure 41. Percentage by weight of prey types found in stomachs of age-0 Delta Smelt collected from 1-6 ppt during April through December in A) 2005, B) 2006, C) 2010, and D) 2011. Number of stomachs examined are shown above the columns. Mean fork length (mm) of Delta Smelt is also shown.
Figure 42. Percentage by weight of prey types found in stomachs of age-0 Delta Smelt collected from < 1 ppt during April through December in A) 2005, B) 2006, C) 2010, and D) 2011. Number of stomachs examined are shown above the columns. Mean fork length (mm) of Delta Smelt is also shown.
Figure 43. Percentage by weight of prey types found in stomachs of age-0 Delta Smelt collected from Cache Slough-Sacramento River Deepwater Ship Channel (CS-SRDWSC) during April through December in A) 2005, B) 2006, C) 2010, and D) 2011. Number of stomachs examined are shown above the columns. Mean fork length (mm) of Delta Smelt is also shown.
Figure 44. Percentage by weight of prey types found in stomachs of adult Delta Smelt collected in 2012 during January through May from A) > 6 ppt, B) 1-6 ppt, C) < 1 ppt, and D) Cache Slough-Sacramento River Deepwater Ship Channel (CS-SRDWSC). Number of stomachs examined are shown above the columns. One fish examined from 1-6 ppt in May had an empty stomach. Mean fork length (mm) of Delta Smelt is also shown.
Although substantial uncertainties about mechanisms remain, the decline of *P. forbesi* in the Suisun region may be related to increasing recruitment failure and mortality in this region due to competition and predation by *P. amurensis*, contaminant exposures, and entrainment of source populations in the Delta (Mueller-Solger et al. 2006, Winder and Jassby 2011, Durand 2010).

The abundance of a more recent invader, the cyclopoid copepod *Limnoithona tetraspina*, significantly increased in the Suisun Bay region beginning in the mid-1990s. It is now the most abundant copepod species in the Suisun Bay and confluence region of the estuary (Bouley and Kimmerer 2006, Winder and Jassby 2011). Gould and Kimmerer (2010) found that it grows slowly and has low fecundity. Based on these findings they concluded that the population success of *L. tetraspina* must be due to low mortality and that this small copepod may be able to avoid visual predation to which larger copepods are more susceptible. It has been hypothesized that *L. tetraspina* is an inferior food for pelagic fishes including Delta Smelt because of its small size, generally sedentary behavior, and ability to detect and avoid predators (Bouley and Kimmerer 2006, Gould and Kimmerer 2010). Nevertheless, this copepod has been found in the guts of Delta Smelt when *Limnoithona* spp. occurs at extremely high densities relative to other zooplankton (Slater and Baxter 2014). Recent experimental studies addressing this issue suggest that larval Delta Smelt will consume and grow on *L. tetraspina*, but growth is slower than with *P. forbesi* (Kimmerer et al. 2011). It remains unclear if consuming this small prey is energetically beneficial for Delta Smelt at all sizes or if there is a breakpoint above which larger Delta Smelt receive little benefit from such prey.

*L. tetraspina*, also reached considerable densities in Suisun Bay and the western Delta over the last decade (Hennessy 2010), although its suitability as food for pelagic fish species remains unclear.

Preliminary information from studies on pelagic fish growth, condition, and histology provide additional evidence for food limitation in pelagic fishes in the estuary (IEP 2005). In 1999 and 2004, Delta Smelt growth was low from the Sacramento-San Joaquin confluence through Suisun Bay relative to other parts of the system. Delta Smelt collected in 2005 from the Sacramento-San Joaquin confluence and Suisun Bay also had high incidence of liver glycogen depletion, a possible indicator of food limitation (Bennett et al. 2008). As previously noted, warm water temperatures during the summer period may have exacerbated lack of food by raising the metabolic rate of Delta Smelt. Based on data for histopathology, date of birth from otoliths, and growth rates from otoliths of Delta Smelt in 2005, Bennett et al. (2008) proposed a novel strategy for Delta Smelt survival in 2005. Natural selection appeared to favor individuals with a specific set of characters, including relatively slow larval development, but faster than average juvenile growth in July. Water temperatures in July typically include the annual maximum (Fig. 16). The salinity field can also change rapidly as freshwater flow out of the Delta changes. Many of these fish surviving into the pre-adult stage had also hatched earlier in the spawning season (i.e., before May).

For many fishes, success at first feeding is believed to be critical to larval survival and a major cause of year-class variability (e.g., “critical period hypothesis,” Hjort 1914, Leggett and DeBlois 1994). In Rainbow Smelt *Osmerus mordax* a related smelt species, calculated larva mortality rates were related to feeding conditions at first feeding that varied on a predictable cycle of 15 days associated with tide and photoperiod (Sirois and Dodson 2000b). In feeding experiments, copepod evasion behavior affected capture by larval Striped Bass, and *E. affinis* was among the more easily captured species (Meng and Orsi 1991). There has been a long-term decline in calanoid copepods in the upper estuary, particularly in the Suisun Region (Winder and Jassby 2011), potentially reducing feeding success, growth and thereby survival. Currently, *E. affinis*
abundance peaks in spring (Hennessy 2010, 2011) coincident with hatching of Delta Smelt. *E. affinis* abundance has been negatively related to X2 since the overbite clam invasion (Kimmerer 2002b). When X2 is “high” outflow is low and *E. affinis* densities are low. These lines of evidence suggest that the first feeding conditions may improve in springs with higher outflow.

Changes in the quality and quantity of available prey may have contributed to the observed reduction in the mean size of Delta Smelt in fall since the early 1990s (Sweetnam 1999, Bennett 2005); however, mean size subsequently increased. The importance of food resources as a driver is supported by Kimmerer (2008), who showed that Delta Smelt survival from summer to fall is correlated with biomass of copepods in the low salinity zone, the central 50% of the summer Delta Smelt distribution. Other variations of this correlation were shown by Maunder and Deriso (2011) and Miller et al. (2012). Miller et al. (2012) have tested for an explicit influence of prey density during the fall. Miller et al. (2012) found a stronger correlation between Delta Smelt abundance during the fall and prey density during the fall than for prey density during the summer.

**Harmful algal blooms**

Periodic blooms of the toxic blue-green alga *Microcystis aeruginosa* during late summer, most commonly August and September are an emerging concern for Delta Smelt (Lehman et al. 2005, Lehman et al. 2013). Although this harmful algal bloom (HAB) typically occurs in the San Joaquin River away from the core summer distribution of Delta Smelt, some overlap is apparent during blooms and as cells and toxins are dispersed downstream after blooms (Baxter et al. 2010). Density rankings of *Microcystis* at TNS stations were highest in the south Delta, east Delta and lower San Joaquin River regions; yet *Microcystis* distribution may be expanding north over time (Morris 2013). Moreover, studies by Lehman et al. (2010) suggest that Delta Smelt likely are exposed to microcystins, which may degrade their habitat and perhaps affect the distribution of Delta Smelt (Baxter et al. 2010). For example, these HABs are known to be toxic to another native fish of the region, Sacramento Splittail (Acuña et al. 2012a) and the alien Threadfin Shad (Acuña et al. 2012b). Histopathology evidence from Lehman et al. (2010) suggested the health of two common fish in the estuary, Striped Bass, and Mississippi Silversides, was worse at locations where microcystin concentrations were elevated.

Indirect effects are also likely as *Microcystis* blooms are toxic to copepods that serve as the primary food resources of Delta Smelt (Ger et al. 2009, 2010a,b). Ger et al. (2009) determined toxicity of one form of microcystin (LR) to two species of calanoid copepods, *E. affinis* and *P. forbesi*, which are important as food to Delta Smelt. They found that, although the copepods tested were relatively sensitive to microcystin-LR compared to other types of zooplankton, ambient concentrations in the Delta were unlikely to be acutely toxic. However, chronic effects were not determined and Lehman et al. (2010) found that *Microcystis* may indeed contribute to changes in phytoplankton, zooplankton and fish populations in the Delta.

Factors that are thought to cause more intensive *Microcystis* blooms include warmer temperatures, lower flows, high nitrogen levels, and relatively clear water (Lehman et al. 2005, Baxter et al. 2010, Lehman et al. 2013, Morris 2013). These conditions occur during dry years in the SFE. Both *Microcystis* abundance and microcystin concentrations have been greater in recent years with dry year conditions (Lehman et al. 2013). These factors can also interact. For example, low flows can provide less dilution of ammonium from wastewater treatment plants (Jassby and Van Nieuwenhuyse 2005, Dugdale et al. 2012, Dugdale et al. 2013) and *Microcystis* can
readily utilize ammonium as a primary nitrogen source during blooms (Lehman et al. 2013). The intensity and duration of *Microcystis* blooms are expected to increase over the long-term, along with any negative impact on aquatic organisms, due to increased frequency of drought conditions associated with climate change (Lehman et al. 2013).

Chapter 5: Updated Conceptual Models for Delta Smelt

In this Chapter we transfer the information on drivers and Delta Smelt responses reviewed and presented in Chapter 4 into the conceptual model framework established in Chapter 3. The Delta Smelt general life cycle conceptual model recognizes the pervasive, year-round importance of the tier 1 landscape attributes and the seasonal importance of the various tier 2 environmental drivers and tier 3 habitat attributes to the tier 4 life stage transitions of Delta Smelt in the four tier 5 “transition seasons” (Fig. 45). Some habitat attributes—food, toxicity, and predation—affect life stage transitions in all seasons, while other habitat attributes—temperature, entrainment and transport, size and location of the low salinity zone, and harmful algal blooms—affect some life stage transition more than others. Clearly, adequate food must be available at all life stages for Delta Smelt to survive. Toxicity is included during all seasons because we know that contaminants of various types are present throughout the year; however, little is known about the direct or indirect effects of contaminants at ambient concentrations on individual Delta Smelt or the population as a whole. Predation is included in all seasons because we recognize that predation is likely the ultimate cause of mortality for most individual fish; however, responses of Delta Smelt to other habitat attributes and environmental drivers such as food availability and turbidity can modify predation risk.

The mechanistic linkages between landscape attributes, environmental drivers, habitat attributes and Delta Smelt responses in the four life stage seasons are depicted as one-way arrows in four new “life stage transition” conceptual models (Figs. 46-49). As mentioned in Chapter 3, the life stage transition conceptual models are nested components of the general life cycle conceptual model (Fig. 8). Each life stage transition conceptual model (Figs. 46-49) includes the habitat attributes hypothesized to affect the transition of Delta Smelt from one life-stage to the next. Hypotheses selected for detailed consideration in Chapter 7 are indicated by “H” in the diagrams. The models also show the landscape attributes and environmental drivers. While the models include many linkages among individual landscape attributes, environmental drivers, and habitat attributes, they do not include linkages between individual habitat attributes and the specific biological processes (growth, survival, reproduction) underlying the life stage transitions. The primary reason for this simplification is that the available data are generally inadequate to fully describe and differentiate among specific functional relationships and mathematical modeling that could help estimate them is beyond the scope of this report. Instead, the combined effects of all habitat attributes on the life stage transition probability are depicted by one upward arrow in each life stage transition conceptual model. This does not imply, however, that all habitat attributes have an equal role in determining life stage transition probability and population success or that the role of each habitat attribute remains constant from year to year.

In the remainder of this Chapter we briefly describe the linkages and associated hypotheses depicted in each of the life stage transition conceptual model diagrams (figs. 46-49). These
An Updated Conceptual Model of Delta Smelt Biology

**Figure 45.** Delta Smelt general life cycle conceptual model.

![Delta Smelt Life Cycle Conceptual Model](image1)

**Figure 46.** Conceptual model of drivers affecting the transition from Delta Smelt adults to larvae. Hypotheses addressed in Chapter 7 are indicated by the “H-number” combinations.

![Transition Model](image2)
Figure 47. Conceptual model of drivers affecting the transition from Delta Smelt larvae to juveniles. Hypotheses addressed in Chapter 7 are indicated by the “H-number” combinations.

Figure 48. Conceptual model of drivers affecting the transition from Delta Smelt juveniles to subadults. Hypotheses addressed in Chapter 7 are indicated by the “H-number” combinations.
hypotheses are stated and addressed in more detail in Chapter 7. All hypotheses focus on the life stage that is transitioning to (i.e. occurs prior to) the next life stage, for example, adults but not eggs and larvae, larvae and post-larvae but not juveniles, and so on. That said, it is important to remember that all life stages overlap and all transitions except for the transitions from adults to eggs and from eggs to freshly hatched larvae are gradual, not abrupt, and delineations of life stages are somewhat arbitrary (see Chapter 3).

The life stage conceptual model for the transition of adult Delta Smelt to eggs and larvae (Fig. 46) includes 5 habitat attributes. Because of the lack of information about specific contaminant effects on Delta Smelt noted above, there are no specific hypotheses regarding the effects of contaminants and possible direct or indirect toxicity on Delta Smelt, but based on the information discussed in Chapter 4, the model does recognize that effects on Delta Smelt or its food supply may be occurring. Food availability and visibility are hypothesized to be important with respect to providing nutrition that allows Delta Smelt to grow into healthy, large adults that can produce a large numbers of high quality eggs as well as multiple clutches of eggs over the spawning season. The availability of food is considered dependent on both food production and the availability of such food to the fish. There are two hypotheses related to predation risk. The first is that turbidity, created by the interaction of high winter and spring flows with the erodible sediment supply in the watershed and within the Delta, influences the vulnerability of Delta Smelt to predators that co-occur with them. The second is that Delta Smelt behaviors that bring Delta Smelt close to channel edges may increase their vulnerability to Largemouth Bass, which generally occupy nearshore and vegetated habitats such as SAV beds. Entrainment risk in this life stage transition conceptual model is focused on adults. Entrainment of adults would reduce the reproductive
potential of the population. Entrainment risk depends on the distribution of the adult Delta Smelt in relation to water diversions, and the magnitudes of water diversions and flows. Delta water temperature determines the beginning and duration of the spawning season (hereafter “spawning window”).

The life stage conceptual model for the transition of Delta Smelt eggs and larvae to juveniles includes 4 habitat attributes (Fig. 47). Food production and availability is important for the survival of larvae to juveniles. Food quantity is dependent on multiple interacting factors. Turbidity is important for early feeding by delta smelt larvae. Predation risk focuses on predation of Mississippi Silversides on Delta Smelt larvae because of recent evidence that such predation occurs. Predation risk is hypothesized to depend on co-occurrence of the two species, with Mississippi Silverside generally being associated with shallower waters, turbidity, which decreases the effectiveness of predators, and water temperature, which affects energy requirements of predators (hunger level). In addition to its effect on predator bioenergetics, water temperature is hypothesized to affect the length of the spawning season (spawning window). If food availability is sufficient, then a longer spawning window may allow the adult population to produce multiple clutches of eggs, resulting in more young. This hypothesis could arguably be included in the previous life stage transition conceptual model, but considering it here allows for consideration of predation on larvae in the context of the time period over which larvae are being produced. Larvae are also at risk of entrainment or transport to unfavorable areas. The magnitude of this risk is hypothesized to depend on an interaction of spring hydrology and water exports. As indicated by numerous arrows, winter and spring hydrology affect Delta Smelt spawning and larval rearing habitat in many ways. We thus also include a more general hypothesis about the hydrological effects on Delta Smelt larval abundance and recruitment.

The life stage conceptual model for the transition of Delta Smelt juveniles to subadults includes 4 habitat attributes (Fig. 48). In addition, there is a stand-alone hypothesis dealing with population dynamics. Juvenile growth and survival is hypothesized to depend on availability and quantity of food. Food production during this summer period is hypothesized to involve complex interactions of clam grazing, nutrients, hydrology and harmful algal blooms. The probability of observing a harmful algal bloom is hypothesized to be a function of the same factors but with temperature playing an important role. Harmful algal blooms may also affect Delta Smelt directly through production of toxic microcystins. Summer water temperatures are hypothesized to have a very direct effect on juvenile Delta Smelt with water temperatures hypothesized to reach stressful levels, affecting their bioenergetics and the area of suitable habitat. The transition probability hypothesis is that at the currently small population sizes, survival from juvenile to subadult is density independent, meaning independent of the number of individuals present (see Chapter 6 for details).

The life stage conceptual model for the transition of Delta Smelt subadults to adults includes 6 habitat attributes (Fig. 49). As for the previous conceptual model, there is a stand-alone hypothesis dealing with population dynamics. As in the previous conceptual model, growth and survival are hypothesized to depend on food availability and food production and availability depends on interactions of a variety of landscape attributes and environmental drivers. Toxicity is recognized as potentially important but no specific hypotheses have been tested. Harmful algal blooms may still be present with hypothesized direct effects on Delta Smelt subadults and indirect effects on their food. Predation risk on subadult Delta Smelt is hypothesized to depend on co-occurrence of Delta Smelt with the two most likely predators, Largemouth Bass and Striped Bass. Largemouth Bass occurrence is linked with that of SAV and the vulnerability of prey to both predators is affected by turbidity and bioenergetics. Water temperature is mainly
hypothesized to have an effect through bioenergetics because water temperature becomes less stressful than in the summer. In this conceptual model the size and location of the LSZ is considered both a landscape attribute and a habitat attribute. In the earlier conceptual models, the LSZ was mainly viewed as a landscape attribute that interacted with other landscape attributes and environmental drivers to create habitat attributes. In this conceptual model the size and position of the LSZ is hypothesized to have certain characteristics that directly determine habitat quantity and quality for Delta Smelt. The transition probability hypothesis is that at the currently small population sizes, survival from subadult to adult is density independent, meaning independent of the number of individuals present (see Chapter 6 for details).

Chapter 6: Delta Smelt Population Biology

This Chapter consists of two main parts. In the first part, we introduce general concepts in population biology that are utilized in the following sections of this Chapter and to generally describe Delta Smelt population dynamics. Explaining these concepts and population trends now is intended to reduce repetitive text in the remaining sections and to reduce possible confusion for readers unfamiliar with the concepts. The concepts are discussed specifically in the context of Delta Smelt.

In the second part of this Chapter, we review information about the life history and population trends of each Delta Smelt life stage represented in our conceptual models, starting with adults. While we describe trends over the entire available time series for each life stage, we pay particular attention to differences in Delta Smelt abundance and life stage transitions between the two most recent wet years, 2006 and 2011. Our working assumption is that these differences should be attributable to differing habitat conditions and, in some cases, management actions. Differences in habitat conditions between these two years will be further explored in Chapter 7.

Population Biology

Recruitment is the addition of new individuals to a population through reproduction or immigration. In fisheries science, the term recruitment was first used by Ricker (1954) to describe the addition of fish of a new generation to a fish population, in other words, the number of young surviving to a particular age or life stage. We use the term recruitment to refer to production of larvae, juveniles, subadults, or adults by adults of the previous generation. Relationships between numbers of spawning fish or other measures of potential spawning stock (e.g., numbers of subadult or mature prespawning fish) and the numbers of fish of a given age or life stage in the subsequent generation are known as stock-recruitment relationships.

Stock-recruitment relationships have been described for many species and are a central part of the management of commercially and recreationally fished species (Myers et al. 1995, Touzeau and Gouze 1998). Different forms of stock-recruitment relationships are possible, including density-independent, density-dependent, and density-vague types. The density-independent type occurs when the current size of the population has little or no effect on the number of recruits (except possibly when stock size is extremely low). This type of population growth is rare in fish
populations and occurs when environmental factors largely determine the survival and number of recruits (e.g., the Longfin Smelt outflow abundance relationship; see Myers 1998). Density dependence occurs when the current population size affects survival and abundance of recruits and thus population growth. In such populations, within the lower range of stock size, the number of recruits is strongly and positively related to stock size. At some point as stock size increases, competition for food (or some other limiting factor) between the adult population and recruits affects survival and abundance of recruits; cannibalism is another means by which recruitment can be affected by stock size. Thus, the growth and survival of the recruit population strongly depends on the density of the stock population. In reality it’s difficult to determine which type of response is occurring (e.g., Myers and Barrowman 1996). Moreover, a predominantly annual fish, such as Delta Smelt, is predicted to conform poorly to models that assume density-dependent recruitment (Winemiller 2005), which appears to be the case (e.g., Rose et al. 2013).

The idea of density dependence is related to the idea of carrying capacity. The carrying capacity of an ecosystem is the number of individuals of all species that can be supported by the available resources. In reality it can be very difficult to apply this idea to a single species in an ecosystem because of the complex relationships among species and the seasonal, annual, and other changes in resource availability. The density vague type of population growth refers to situations where there is not a statistically demonstrable stock-recruitment relationship observable in available data.

In density-dependent stock-recruitment relationships, the factors causing the density dependence can operate at various points in the life cycle of the new generation. For some species, the concept of density dependence is separated into two concepts. In this formulation, density-dependent stock recruitment is limited to the direct effects of the adult stock on recruitment of the next generation, as described above. For example, if a large spawning stock has a limited spawning area, as in the case of salmonids, then successive waves of female spawners are known to re-excavate previous nests while building their own, substantially increasing mortality of the eggs. Density dependence could also occur at the larval or juvenile stage if adults are predatory and feed on young, or if adults are in direct competition for food or space with young. The second concept of density-dependent survival is often inextricably linked to density-dependent stock-recruit relationships because the mechanisms causing declines in recruits at high stock levels are unknown. In density-dependent survival, the abundance of young affects their own survival.

In the case of Delta Smelt, density dependent survival could occur if many of the larvae starved because of insufficient food supplies due to competition with other Delta Smelt larvae, or other species. Because many Delta Smelt die after their first spawning, density-dependent survival is certainly the dominant mechanism for the species and for the remainder of this report the direct effects of adults on survival of eggs and larvae are assumed to be minimal. If resources were sufficient for larvae and juvenile fish to survive in large numbers, the surviving subadults might overwhelm food sources (i.e., surpass carrying capacity), resulting in low survival and poor reproductive output. Thus, it is important to understand species ecology and survival between life stages to understand how density dependence is affecting a population. This is particularly important for fishes in estuaries where environmental factors can create large variation in habitat size and food web productivity from season to season and year to year, thus affecting carrying capacity and the potential for density-dependent survival.

Density-independence is more straightforward. In this case, the population is controlled by factors unrelated to the density of the population. For example, high water temperatures will affect individual fish, whether the population is large or small. In reality, populations can be
affected by both density-dependent and density-independent factors at different times. This interaction is the basis for the idea of compensatory density dependence. In this formulation, a population is governed by density independent factors when population size is small. As the population increases and approaches the carrying capacity, density-dependent factors become important and the population growth rate declines. Fluctuations in carrying capacity, as noted above, are an added complication. Again, it is essential to understand the ecology of the species and survival between life stages to understand the relative importance of density dependent and density independent factors.

Unfortunately, Delta Smelt were never of sufficient interest as a commercial or recreational species to warrant development of stock-recruitment models until they were listed. Data now used to develop stock-recruitment models for Delta Smelt started becoming available after the initiation of fisheries studies and monitoring surveys in the late 1950s (TNS initiated 1959; FMWT initiated 1967) in association with the planning and operation of the CVP and SWP. These IEP fish monitoring surveys were designed to produce relative abundance indices or catch-per-unit-effort (CPUE, e.g., number per trawl) that could be used to monitor trends in abundance over time. More recently, annual abundance indices based on these surveys have also been incorporated into stock-recruitment relationships (e.g., Moyle et al. 1992, Sweetnam and Stevens 1993, Miller 2000, Bennett 2005, Maunder and Deriso 2011). Neither of these early IEP fish monitoring surveys (TNS, FMWT) were specifically designed to monitor Delta Smelt, but instead targeted primarily the commercially and recreationally more important Striped Bass. As researchers began using TNS and FMWT indices for Delta Smelt analyses, they began investigating how the indices performed and means to improve them (see Wadsworth and Sommer 1996, Miller 2000, Newman 2008). This work is ongoing and also includes similar investigations for the newer SKT (initiated in 2002) and 20 mm survey (initiated in 1995) monitoring surveys.

The two stock-recruitment relations based on the longest data records include the relationship of the FMWT abundance index with the FMWT abundance index in the previous year and the relationship of the TNS abundance index with the FMWT abundance index in the previous year (Fig. 50). Because of the large changes that have occurred in the Delta ecosystem, including the invasion by \( P. \ amurensis \) and the POD, these plots can be difficult to interpret because carrying capacity is assumed to have changed (Bennett 2005, Kimmerer et al. 2000, Sommer et al. 2007). It does appear that there is much more variability associated with the FMWT relationship compared to the TNS relationship. This might indicate variable survival between the juvenile and subadult life stage.

In any form of a stock-recruitment model, there is a point at which low adult stock will result in low juvenile abundance and subsequent low recruitment to future adult stocks. This can occur even under favorable environmental conditions while the stock “rebuilds” itself. From a stock-recruitment perspective, the recent low abundance of Delta Smelt is of particular concern. Since about 2002, the current population is smaller than at any time previously in the record, with the exception of the 2011 year class. This strong year class suggests that Delta Smelt have yet to reach low levels where the stock will need years to rebuild, at least to pre-POD levels (Fig. 3).

In addition to their use in exploring stock-recruitment relationships, ratios of annual Delta Smelt abundance indices can also be used to obtain rough estimates of relative annual recruitment and survival rates (figs. 51 and 52). As for the stock-recruitment relationships these recruitment and survival indices should be interpreted with caution given the large changes that have taken place in the Delta and the absence of estimates of variability for the indices. The main utility of these
indices is identifying years with relatively high or low survival for a specific life stage transition or life stage transitions with differences in annual variability.

Here, we use the ratios of abundance indices for different life stages of the same generation as indices of survival (survival indices, Fig. 51) and the ratios of current to preceding year abundance indices as indices of recruitment (recruitment indices, Fig. 52). For the density-independent case, recruitment rate is independent of the size of the adult population. The number
of recruits produced is the product of recruitment rate and the size of the adult population. For this report, we assume that the estimates have sufficiently low and comparable uncertainty to provide worthwhile interpretations, as long as caution is exercised. It is also important to remember that abundance, survival, and recruitment index values are only meaningful in a relative, not in an absolute sense.

The annual stage to stage survival indices from larvae to juveniles, subadults, and adults are shown in Figure 51. The relative recruitment rates from adults and subadults in one year to larvae, juveniles, and subadults the next year are shown in Figure 52. We recognize that a life cycle model with environmental covariates is needed to fully assess the combined effects of stock-recruitment and stage-to-stage survival indices on Delta Smelt population dynamics. Nevertheless, examination of the recruitment and survival index data sets reveal several interesting patterns for the POD period (2003-2013).
Figure 52. Delta Smelt recruitment indices based on the annual adult, larval, juvenile, and subadult abundance indices provided by the Spring Kodiak Trawl (SKT, adults), 20 mm Survey (20 mm, larvae), Summer Townet Survey (TNS, juveniles), and Fall Midwater Trawl (FMWT, subadults).
First, interannual variability in these stock and survival indices declines from larval recruitment (coefficient of variation (CV): 92%), to subsequent larvae to juvenile survival (CV: 67%), juvenile to subadult survival (CV: 43%), to subadult to adult survival (CV: 38%). This result is consistent with expected highly dynamic patterns of recruitment and survival for an annual opportunistic species such as Delta Smelt. The pattern of reduced variability in survival for larger fish suggests that older fish may no longer be vulnerable to some forms of mortality affecting earlier life stages either because a factor is no longer important when larger fish are present (e.g., effect of summer high water temperatures on juveniles) or that larger fish escape some forms of mortality (e.g., larger fish are no longer eaten by the large variety of predators able to consume larvae).

Second, the patterns of adult and larval abundance (Fig. 3) and adult to larvae recruitment (Fig. 52a) suggest: (1) even a small adult Delta Smelt stock can produce a large number of larvae under the right habitat conditions; but (2) larval recruitment is not a good predictor of juvenile survival and subsequent adult stock size. In other words, good larval recruitment sets the stage for population recovery, but good survival through subsequent life stage transitions is needed to realize its potential.

Third, there are clear contrasts in Delta Smelt responses between the two wet years 2006 and 2011 (the years of particular interest in this report) (Figs. 51 and 52). Since the initiation of the SKT survey for adult Delta Smelt in 2002 (indices calculated beginning in 2003), the recruitment of larvae from adults was greatest in the two wet years 2006 and 2011 (Fig. 52a) compared to the other, drier years in the time series, but in 2006 very strong adult to larvae recruitment was followed by very poor larvae to juvenile survival in the summer (Fig. 51a) and only average survival in the fall (Fig. 51b) and winter (Fig. 51c). This led to low abundance of the subsequent life stages of the 2006 cohort. Survival from larvae to juveniles and subadults was much better in 2011 and, along with good recruitment, led to the highest juvenile and adult abundance indices since the onset of the POD (Fig. 3). In other words, good recruitment set the stage for population recovery, but a substantial abundance increase was realized only in 2011. Unfortunately the 2011 abundance increase was short-lived; it was immediately followed by poor recruitment and survival in 2012 and abundance indices for the 2012 and 2013 cohorts were once again at the low levels typical for the POD period (Fig. 3). Several consecutive years of good recruitment and survival are likely needed for a more sustained increase of the Delta Smelt population abundance to pre-POD abundance levels. Population declines such as the decline experienced by Delta Smelt do not only reduce the number of individuals, but can also reduce the genetic diversity present in the population. While the 2011-2012 data suggest that recovery of Delta Smelt abundance can still be fairly rapid via high larval recruitment followed by good survival (Figs. 51 and 52) recovery of genetic diversity is a much slower process which is an important conservation concern (Fisch et al. 2011).

Small Delta Smelt population size affects the effective population size \((N_e)\), a measure of the genetic properties of a population and the abundance at which significant genetic diversity is lost due to inbreeding (Falconer and Mackay 1996, Schwartz et al. 2007, Antao et al. 2010). In many species \(N_e\) may be orders of magnitude smaller than the census population size \((N)\) and low \(N_e/N\) ratios indicate the population may be in danger of losing genetic variability, potentially resulting in reduced adaptability, population persistence, and productivity (Hauser et al. 2002). For Delta Smelt, Fisch et al. (2011) detected a genetic bottleneck in each of four sampling years (2003, 2005, 2007 and 2009) and observed a significant decline in effective population size between sampling years 2003 and 2007 (Fisch et al. 2011). The genetic signal of the decline in \(N_e\) is corroborated by the observed abundance index declines and support the hypothesis that decreases
in Ne and allelic richness have likely occurred over the last few decades (Fisch et al. 2011). Genetic changes within the Delta Smelt population deserve continued evaluation with respect to changes in population size.

In addition, Delta Smelt recruitment and the fecundity of adult Delta Smelt likely vary substantially from year to year (Rose et al. 2013b). Delta Smelt fecundity is a function of female size (Bennett 2005, Lindberg et al. 2013). The mean size of adult Delta Smelt declined in the early 1990s (Sweetnam 1999), possibly due to changes in the food web (see Chapter 4), but substantially recovered in the late 2000s. Another possible reason is that in some recent years, there may have been selection for smaller, late-spawned larvae as a result of export pumping schedules (Bennett 2011). For example, Bennett (2011) proposed that high export pumping in late winter may have resulted in high entrainment mortality of offspring from larger, fitter, early spawning females, which produced larger, fitter offspring (Bennett 2011). Further, Bennett et al. (2008) and Bennett (2011) posited that curtailment of export pumping in mid-April related to the Vernalis Adaptive Management Program (VAMP), allowed for greater survival of later-spawned, smaller larvae. The major concern is that these smaller later-spawned larvae have less opportunity to grow to large adult size, especially when food is scarce. If correct, the combined effects of export pumping and food supply on Delta Smelt growth and size could have a nonlinear impact on overall fecundity and population success. This is corroborated by the results from individual-based modeling which showed that growth in fall-winter and the subsequent number of eggs produced per adult were the most important factor determining the success of the next generation (Rose et al. 2013b). Moreover, repeated losses of early-spawned larvae could potentially have a negative effect on expression of this important phenotype and result in eventual loss of genetic variability in the population, and contribute to the genetic bottlenecks reported by Fisch et al. (2011).

Given the unprecedented low abundance of Delta Smelt since 2002 (Fig. 3, summer and fall), serious consideration should be given to evaluation of Allee effects. Allee effects occur when reproductive output per fish declines at low population levels (Berec et al. 2006). In other words, below a certain threshold the individuals in a population can no longer reproduce rapidly enough to replace themselves and the population, exhibiting inverse density dependence, spirals to extinction. For Delta Smelt, possible mechanisms for Allee effects include processes directly related to reproduction and genetic fitness such as difficulty finding mates, genetic drift, and inbreeding (Gascoigne et al. 2009), although none of these effects have been documented yet in Delta Smelt (Fisch et al. 2011). Other mechanisms related to survival such as increased vulnerability to predation (Gascoigne and Lipcius 2004) are also possible. While theoretical work suggests that Allee effects might be common in nature, empirical evidence for Allee effects in natural populations of fishes remains relatively sparse (Myers et al. 1995, Liermann and Hillborn 1997), possibly because they are often masked by measurement errors (Gregory et al. 2010). Recent meta-analytical work by Keith and Hutchings (2012) suggests that Allee effects in marine fish species might be more common than previously thought. But even in the absence of “true” Allee mechanisms, small population size (Hutchings 2013) can produce an emergent Allee effect and prevent recovery of collapsed fish populations even when threats are reduced (Kuparinen et al. 2012). This may be one of the reasons why recovery of many collapsed fish populations remains slow despite large reductions in fishing (Pauly et al. 1998, Hutchings et al. 2010). This finding challenges the traditional fisheries management view that depleted populations will grow and recover rapidly when fishing pressure is relaxed (Hilborn and Walters 1992). In addition, the interactive effects of multiple Allee effects may have important implications for species conservation, but have not yet been well explored in ecology (Berec et al. 2006).
Compensatory density dependence predicts that a fish’s population growth or survival rates can increase when abundance is low and decrease if abundance increases beyond a carrying capacity (Rose et al. 2001). If compensatory density dependence occurred in 2011, Delta Smelt survival would be expected to increase as long as the carrying capacity of the environment was not exceeded. Therefore, the sudden increase in subadult abundance in 2011 is consistent with the higher survival predicted by compensatory density dependence at low population abundance coupled with widespread availability of good habitat conditions throughout the year. Among the remaining comparison years, both 2005 and 2006 show evidence of compensatory recruitment to larvae (Fig. 52a). Adult abundance was moderately high in 2005, but low in 2006 and 2010 (Fig. 3). As predicted by compensatory density dependence processes, the recruitment index to larvae was higher in 2006 than in 2005. However, low adult abundance in 2010 did not give way to a similarly high recruitment index (Fig. 52a). In addition, the relatively high recruitment index in 2006 did not result in a higher larval abundance index compared to 2005 (Fig. 3). These inconsistencies, combined with a small number of comparison years, prevent any firm conclusion regarding compensatory recruitment or survival.

Similarly, if compensatory density-dependent survival was important we might expect larva to juvenile survival to be lower when larva production per adult was higher assuming similar adult populations. This was not the case for 2006, 2010, and 2011, which had relatively similar values for the SKT abundance index (figs. 3). In 2006, larval survival was low with high larval production per adult, and 2010 and 2011 had very similar larval survivals with similar adult abundances. Finally, in 2011, the highest population of juveniles led to the highest population of subadults and adults (2012 SKT), which argues against compensatory density-dependent survival. These comparisons argue against strict compensatory density dependence operating within the POD years. It seems more likely that population dynamics are driven by density independent relationships with factors such as summer water temperatures and resource availability (fluctuations in carrying capacity); however, the evidence is not conclusive. In particular, we do not understand how carrying capacity fluctuates over seasons and years or how other factors, such as predation, affect carrying capacity (Walters and Juanes 1993; Walters and Korman 1999).

**Adults**

**Life History**

The Delta Smelt is generally considered a diadromous seasonal reproductive migrant, and in the winter, many adult Delta Smelt move upstream into fresh water for spawning (Moyle et al. 1992, Bennett 2005, Sommer et al. 2011). These movements may be a specific change in behavior in response to one or more environmental cues, for example, to the rapid and often dramatic environmental changes during winter first flush periods (Sommer et al. 2011, Bennett and Burau 2014). Focused, fixed-station sampling in the winters of 2009-10 and 2010-11 revealed higher catch of Delta Smelt at higher turbidity levels, as well as an asymmetry in probability of catch with respect to tidal phase; catch was highest in the channels during flood tide, but highest near the shoreline during ebb tides (Bennett and Burau 2014). This change in horizontal channel position with respect to tidal direction has recently been confirmed by a second study in the fall of 2012 that used the “SmeltCam,” an underwater video camera attached to the cod-end of the FMWT net to detect Delta Smelt (Feyrer et al. 2013). This study demonstrated that during flood tides, Delta Smelt were relatively abundant throughout the water column, but less abundant during ebb tides, and found only in the lower portion of the water column and closer
to shorelines. This asymmetry in catch supports the idea of a “tidal surfing” behavior during migration that may minimize energetic costs of upstream movement and allow Delta Smelt to follow favorable conditions with respect to turbidity and salinity (Feyrer et al. 2013). Variations of this behavior would allow fish to maintain position in the channel (stay on the edge during flood or ebb tide) or move downstream (move into the channel on ebb tide).

It is also possible that Delta Smelt movements do not represent a change in behavior; rather, fish are simply expanding their foraging or refuge distribution to habitat upstream when it becomes turbid or otherwise more suitable during and after the first flush period (Murphy and Hamilton 2013). The specific mechanism for the seasonal change in distribution, however, may be more a matter of terminology than of ecological relevance for a fish with as small a home range as Delta Smelt. Here, we acknowledge the existence of both possibilities, but will use the term “spawning migration” to simply refer to a directed movement upstream or downstream occurring prior to and during the spawning season. Using this definition, this seasonal change counts as a migration since it represents a relatively predictable and substantial change in distribution that has adaptive value including potential spawning, foraging and refuge functions (Lucas and Bara 2001).

The Delta Smelt spawning migration from their low-salinity rearing habitat into freshwater usually occurs between late December and late February, typically during first flush periods when inflow and turbidity increase on the Sacramento and San Joaquin Rivers (Grimaldo et al. 2009, Sommer et al. 2011a). Increased catches of Delta Smelt in the Delta Juvenile Fish Monitoring Program’s Chipps Island Trawl Survey and at the south Delta salvage facilities are unimodal in most years and occur within a couple of weeks of first flush events, suggesting that adult Delta Smelt are responding to environmental changes and migrating rapidly upstream once the first flush occurs (Grimaldo et al. 2009, Sommer et al. 2011a). However, spawning migrations are not always upstream. During occasional periods of very high river flows that spread freshwater habitat throughout much of the estuary, some Delta Smelt “migrate downstream” from rearing habitats in Suisun Bay and the Delta to freshwater spawning habitats as far west as the Napa River (Hobbs et al. 2007). Also under high flow conditions, it is possible that some Delta Smelt may not migrate in any direction; if their brackish-water rearing habitat becomes fresh, they can presumably spawn in suitable areas nearby. In addition, there is a small subset of the population that appears to remain in the Cache Slough complex year around; these fish presumably stay in the region for spawning (Sommer et al. 2011).

Osmerids generally spawn in shallow waters (Moulton 1974, Murawski et al. 1980, Hirose and Kawaguchi 1998, Martin and Swiderski 2001, Bennett 2005). It is believed that Delta Smelt spawn over sandy substrates in shallow areas based on the observation that first hatch larvae are collected in high concentrations in areas near expansive sandy shoals (Bennett 2005, L. Grimaldo, U.S. Bureau of Reclamation, unpublished data); confirmation of this hypothesis has not been verified through egg collections or observations of spawning adults, except in mesocosm studies (J. Lindberg, U.C. Davis, unpublished data). Pilot studies to identify egg deposition areas have been conducted by the IEP but these efforts were unsuccessful; it is unknown whether it was due to the method used, locations selected, or because of the low probability of detecting eggs from a relatively rare species.

The Delta Smelt is an opportunistic strategist (Nobriga et al. 2005). Opportunistic strategists are characterized by their short life spans, but high intrinsic rates of population increase driven by rapid maturation and repeat spawns over a protracted spawning season (Winemiller and Rose 1992). The importance of per capita fecundity to the success of the Delta Smelt population was recently highlighted in an individual-based modeling study (Rose et al. 2013a,b). In culture,
Delta Smelt can spawn up to four times per year depending on water temperature (J. Lindberg, U.C. Davis, unpublished data). Recent evidence indicates that Delta Smelt can spawn multiple times in the wild if water temperatures stay cool in the later winter and early spring (Wang 2007, L. Damon, CDFW, written comm. 2013). The ability of Delta Smelt to spawn multiple times in the wild could substantially increase per capita fecundity over previous estimates for individuals of a specific size. It could also be a contributing factor to the large interannual variability in adult to larvae recruitment (Fig. 52a).

**Population Trends**

Adult Delta Smelt are monitored by the Spring Kodiak Trawl (SKT) survey which was initiated by CDFW (then CDFG) in 2002 and runs from January to May each year (Honey et al 2004). An indexing method was recently developed by CDFW for the SKT survey, allowing for year to year comparisons as well as comparisons with the abundance indices for other life stages (Fig. 3). The SKT index time series used in this report comprises 11 annual indices, from 2003 to 2013; no index is available for 2002. Each index represents the abundance of adult fish hatched in the previous calendar year that survive to spawn at the beginning of the next calendar year.

The highest SKT index on record occurred in 2012 (147), as a result of the high 2011 abundance of younger fish, and the lowest in 2006 (18). Of the four comparison years, 2005 had the highest SKT index (51), followed by 2010 (27) and 2011 (20) and then 2006 (18). While the SKT index was thus lower in the two wet years than in the two drier years, the SKT index increased substantially in each of the years following the two wet years; however it increased only 2-fold from 2006 to 2007 while it increased 7-fold from 2011 to 2012 (Fig. 3). It is also possible that the SKT is less effective during very high flow events. Delta outflow at times exceeded 200,000 cfs in winter 2011 and 300,000 cfs in winter 2006. These high flow events might have contributed to the low SKT indices in these two wet years, if Delta Smelt remained near shore to avoid displacement or moved into San Pablo Bay with the LSZ. In both cases they would be outside of SKT sampling range. Further evaluations are needed, however, to investigate and quantify this hypothesized effect.

The annual adult Delta Smelt abundance indices track the annual abundance indices of subadults calculated from the previous years’ FMWT survey closely (Fig. 53; see also Kimmerer 2008). The relationship is particularly strong at higher fall abundance indices (FMWT index > 50), with more variability at lower abundance indices. Before the POD decline in 2002, all Delta Smelt FMWT indices were greater than 50 (Fig. 3). Thus, the FMWT might provide a useful surrogate for estimating long-term abundance trends in the adult Delta Smelt population prior to the initiation of the SKT survey in 2002, but great caution is warranted with the approach because this hindcasting would rest on only four data points with high leverage (2003-2005, 2012) and assume stable subadult to adult survival relationships and habitat conditions, neither of which is likely true. Moreover, the Kodiak trawl more efficiently captures Delta Smelt than the FMWT net. The SKT survey was set up to target Delta Smelt, while the FMWT survey was designed to monitor young Striped Bass, which tend to be larger than Delta Smelt during fall; however, there is no reason to expect the difference in capture efficiency to affect the relationship, unless such differences were a function of population size (i.e., efficiency was different above and below FMWT = 50). The utility of the FMWT as a descriptor of long-term adult population trends in the absence of long-term data from the SKT will benefit from ongoing IEP efforts to quantitatively estimate the efficiency of the FMWT and to compare efficiencies of different trawling gear and protocols. While survival from subadults in the fall (FMWT) to adults in the winter and spring (SKT) (Fig. 53) has been more stable than adult to larvae recruitment and survival between other
life stages (Figs. 51 and 52), it nevertheless shows some variability, especially when abundance is low. These data suggest that at least in the POD decade, adult numbers appear largely driven by juvenile abundance and the influence of changes in winter-time habitat attributes is less important and relatively stable from year to year.

The number of adult spawners affects population dynamics through production of eggs. Potential reproductive output is proportional to the number of adult female spawners, the clutch size for females of a specific size, and the number of egg clutches produced by each female. Although egg production in the wild has not yet been documented, we can evaluate the relationship of the SKT adult population index to the 20 mm Survey abundance index (Fig. 54). This relationship does not appear to be strong during the POD period (linear regression, P > 0.05). This suggests that egg production or subsequent hatching of eggs and survival of larvae and thus overall recruitment of larvae from the previous generation’s adults is affected by other factors than adult population size. Hypotheses about the effects of habitat attributes in our conceptual model on adult growth and fecundity and recruitment of young are explored in Chapter 7.

Clutch sizes of fish collected in the SKT were not measured, but annual fork lengths of Delta Smelt collected in the SKT did not vary greatly (Fig. 55). It does not appear that clutch size should have varied much in the POD years, including the four comparison years 2005-6 and 2010-11, with 2003 as the exception where the median length was greater than 70 mm standard length (Fig. 55). For Delta Smelt, which are now considered seasonal indeterminant spawners (i.e., they spawn multiple times), total reproductive output of an individual female should vary with: 1) size at the onset of the spawning window because batch fecundity is a function of size (Bennett 2005, CDFW unpublished data), 2) length of the spawning window, which is the number of days with suitable water temperatures for spawning (see larval section below) and determines the number of batches possible; and 3) growth during the spawning window, which can potentially improve batch fecundity over time (see larval section below). Obviously, reproductive output will be higher in years when adult females are larger, abundances are higher, and the spawning window is prolonged such that multiple clutches are produced. Note that maximum reproductive output of the adult population at the beginning of spawning is not often realized due to mortality arising from density-dependent (e.g., food limitation or predation) or density-independent (e.g., entrainment, contaminants) mechanisms. According to Bennett (2011), larvae from bigger, early-spawning females may be disproportionally lost to CVP and SWP entrainment. In this report, we consider years when there are bigger females and/or a higher spawning stock size to be better in terms of reproductive potential than years when adult female size and spawning stock are smaller.

**Larvae**

**Life History**

Adult Delta Smelt, through their selection of spawning sites and spawn timing, largely determine the early rearing habitat and environmental conditions encountered by larvae. Given the Delta Smelt’s annual life cycle, small size at maturity, relatively low fecundity, and small egg size compared to other fishes, life history theory suggests that parental care, here limited to selection of spawning sites and spawn timing, should be an important factor in reproductive success (Winemiller and Rose 1992). Since eggs have not been detected routinely in the wild, spawning and early rearing habitat locations are inferred from collection of ripe adults and early stage
larvae, which occur from the Delta margins through eastern Suisun Bay (see: http://www.dfg.ca.gov/delta/projects.asp?ProjectID=SKT; Wang 1986, 1991, 2007). In culture, Delta Smelt begin spawning as water temperatures increase to 10-12 °C, at which time individual females accompanied by several males select appropriate water velocities and release gametes close to the substrate from dusk to dawn (Baskerville-Bridges et al. 2004b). In lab experiments, females deposited significantly more eggs on sand and gravel substrates as compared to other substrates offered for egg deposition (J. Lindberg, U.C. Davis, unpublished data). Based on periodicity in egg deposition in culture, Bennett (2005) proposed that spawning likely coincides with peak tidal currents (i.e., spring tides), which would result in hatching near neap tides. Such a strategy would limit the initial tidal dispersal of larvae.

In culture, larvae hatch after an 11-13 day incubation period at 14.8-16.0 °C and begin a short period of buoyancy (or positive phototaxis; Baskerville-Bridges et al. 2004b) prior to slowly settling to the bottom (Mager et al. 2004). After this buoyant period, Mager et al. (2004) found that larvae were demersal unless actively swimming to feed, which occurred only during daylight hours. Exogenous feeding begins at 5-6 days post-hatch as the last of the yolk sac is absorbed; the lipid globule is absorbed at 10 days (Mager et al. 2004) providing some nutritional reserve if feeding conditions are poor. Larvae probably remain somewhat bottom oriented until swim...
bladder and fin development are complete at about 65 days of age and about 20 mm TL (Mager et al. 2004, Baskerville-Bridges et al. 2004b), at which time they can fully control their buoyancy and efficiently use tidal and river currents to migrate. The center of distribution for Delta Smelt larvae and young juveniles is generally downstream of the spawning habitat, but upstream of and varying in association with X2 during spring (Dege and Brown 2004).

Early larval stages of Delta Smelt (4-15 mm) tended to be poorly collected by gear previously used in historical SFE egg and larval surveys (Striped Bass Egg and Larva Survey; sled-mounted 500 micron mesh net with 0.38 m$^2$ mouth area), but with growth and development greater proportions of the population become vulnerable. This observation led to a sampling gear change in the mid-1990s from the historical egg and larval gear to new gear targeting more vulnerable post-larvae and early juvenile Delta Smelt (i.e., 20 mm Survey). The improved catch and distribution information resulting from this change has since proven valuable to the management of Delta Smelt, and the 20 mm Survey results are now considered essential information (USFWS 2008). In the mid-2000s, an abundance index was developed from 20 mm data (Gleason and Adib-Samii 2007) that has since been used to index abundance trends of larvae in spring (e.g., Hieb et al. 2005, Contreras et al. 2011). We use 20 mm Survey abundance indices as one Delta Smelt end-point to evaluate the support for our hypotheses concerning the environmental drivers and habitat attributes responsible for abundance and survival of larvae.
Population Trends

The highest larval abundance indices on record occurred in the late 1990s, shortly after the initiation of the 20 mm survey in 1995. The lowest larval abundances were observed in 2007-2010 (Fig. 3). In 2011, larval abundance improved substantially from the recent minimum in 2007, and achieved levels comparable to those earlier in the 2000s (Fig. 3). Although 2011 larval abundance compared favorably to that of 2010, it remained below levels of 2005 and 2006. Thus, the modest larva abundance in 2011 did not appear sufficient to explain the high FMWT index observed in 2011 (Fig. 3). As explained above, larval abundance does not track the abundance of the parent generation very well (Fig. 54). In contrast, subsequent life stages of the same cohort track larval abundance and abundance relationships of larvae (log 20 mm index) with juveniles (log TNS index) and subadults (log FMWT index) in the same year are statistically significant (Fig. 56). However, the linear regression based on the FMWT explains less variance than the linear regression based on the TNS suggesting more variability in the abundance of the older life stages. This suggests that factors affecting juvenile mortality rates also play an important role in eventual recruitment.
Juveniles

Life History

During summer, juvenile Delta Smelt primarily rear in the west Delta, Suisun Bay, and Cache Slough complex (Moyle 2002, Bennett 2005, Merz et al. 2011, Sommer and Mejia 2013). As in late spring and fall, the center of distribution of the fish occurs in the low salinity zone, with the exception of the Cache Slough complex. The degree to which the fish use particular geographic areas depends on salinity, temperature, and turbidity (Nobriga et al. 2008); other factors that may affect their summer distribution include Microcystis distribution, and possibly prey density, bathymetric features, or other water quality constituents. As noted previously, Delta Smelt used to be common in the central and south Delta during the summer months, but this is no longer the case (Nobriga et al. 2008).

Population Trends

Relative abundance of juvenile Delta Smelt is presently indexed by the Summer Townet Survey (TNS). The survey was not designed specifically to measure Delta Smelt abundance and catches are low (Honey et al. 2004). Nonetheless, patterns in the annual abundance index provide a useful basic measure of population trends.

The TNS index rebounded substantially in 2011, but declined to a value consistent with low recent year indices in 2012 (Fig. 3). This pattern of persistently low abundance is consistent with the POD, which began over a decade ago (Sommer et al. 2007, Thomson et al. 2010). During the last decade, TNS abundance indices were especially low from 2005-2009 (Fig. 3). The onset of the 2005-2009 period of low juvenile abundance was characterized by extremely low larvae to juvenile survival in 2005 and 2006 (Fig. 51). Larval survival to juveniles recovered somewhat in the following years, but TNS indices stayed low (Fig. 3). Historically (e.g., early 1970s), high levels of Delta Smelt abundance during summer apparently allowed density dependent effects to occur between summer and fall in some years; this conclusion was still supported after the species declined in the early 1980s, but the apparent carrying capacity was lower (Bennett 2005). The available trawl data suggest that this trend of declining carrying capacity has continued as suggested by the very low Fall Midwater Trawl indices produced by a range of juvenile TNS abundance levels, during the POD years (Fig. 57).

Subadults

Life History

During fall, subadult Delta Smelt primarily rear in the western Delta, Suisun Bay, and Cache Slough complex (Moyle 2002, Bennett 2005, Sommer and Mejia 2013). The center of distribution is in the low-salinity zone (Sommer et al. 2011), with the exception of the Cache Slough complex. The degree to which the fish use particular geographic areas depends on salinity and turbidity (Feyrer et al. 2007). Other factors that may affect their distribution during the fall include Microcystis distribution and water temperature in the early fall (September-October), and possibly prey density.
**Population Trends**

Population trends for subadult Delta Smelt are presently indexed by the FMWT. Like the TNS, the FMWT was not designed specifically to measure Delta Smelt relative abundance and catches are low (Honey et al. 2004, Newman 2008). The data are nonetheless a useful basic measure of population trends, except perhaps at very low abundance (i.e., FMWT index values less than about 50; Fig. 53). However, the general agreement between the FMWT and subsequent Spring Kodiak Trawl (SKT) sampling (Fig. 53), suggests that FMWT results are a reasonable indicator.

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**Figure 56.** Relationship of annual index of Delta Smelt abundance from the 20 mm survey (20 mm) with the annual indices from the summer townet survey (TNS) and fall midwater trawl survey (FMWT). Year labels correspond to the comparison years of interest. The linear regressions with all index values log-transformed to address non-normal distributions in the raw data are: Log 20 mm index = 0.57 + 0.87(Log TNS index), n = 19, p < 0.05, $R^2 = 0.44$ and Log 20 mm index = 1.30 + 0.81(Log FMWT index), n = 19, p < 0.05, $R^2 = 0.27$. 

![Graph showing relationship between Delta Smelt abundance indices from different surveys.](image-url)
of general trends in abundance of adult Delta Smelt.

The FMWT index rebounded substantially in 2011, but declined to a value consistent with low recent-year indices in 2012 (Fig. 3). During the last decade, FMWT indices were especially low from 2005-2010 (Fig. 3). After the rebound in 2011, the index went back to a lower level similar to the 2005-2010 period. Since 2003, the juvenile to subadult survival index was lowest in 2004. During the four comparison years, the juvenile to subadult survival index was lowest in 2010, but relatively high in the other three years and highest in 2011 (Fig. 51).

Historically, high levels of Delta Smelt abundance during summer apparently resulted in density-dependent mortality between summer and fall in some years (Bennett 2005). This conclusion was still supported after the species declined in the early 1980s, but the apparent carrying capacity, meaning the magnitude of the FMWT index relative to the TNS index, was lower (Fig. 57). The available FMWT data suggest that these trends of density-dependent mortality during the summer-fall and declining carrying capacity have continued (Fig. 57). The close correlation of the FMWT and SKT (Fig. 53) indicates that the factors likely affecting survival of Delta Smelt to the adult spawning population operate earlier in the life cycle (i.e., between the egg and subadult life stages). Additional mortality certainly occurs between the FMWT and SKT but the lack of variability around the regression line suggests there is not a lot of variability in the rate of that mortality. Thus, the relative annual spawning stock appears to be largely determined by fall of the birth year.

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**Figure 57.** Plots of fall midwater trawl (FMWT) abundance index as a function of summer towent survey (TNS) abundance index for 1982-2013 and 2003-2013. Note the very different scales for both axes. Lines are LOWESS smooths.
Chapter 7: Using the Conceptual Model—Why did Delta Smelt abundance increase in 2011?

In this Chapter, we further explore Delta Smelt responses and habitat attributes as depicted in the driver and life stage transition conceptual model diagrams presented in Chapter 5. The purpose is to demonstrate the utility of our conceptual model framework for generating hypotheses about the factors that may have contributed to the 2011 increase in Delta Smelt abundance. For each life stage transition, we explore a series of hypothesized linkages among ecosystem drivers, habitat attributes, and Delta Smelt responses. We evaluate these hypotheses by comparing habitat conditions and Delta Smelt responses in the wet year 2011 to those in the prior wet year 2006 and in the drier years 2005 and 2010.

In this Chapter we briefly describe the comparative approach and the hydrological conditions during the four years that are the focus of our comparisons. We then state and explore each hypothesis for the adult, larval, juvenile, and subadult life stages of Delta Smelt using data sources described in Chapter 3. Key points from these evaluations, as well as previous report Chapters, along with benefits and limitations of the comparative approach are summarized and discussed in Chapter 8. In several cases, we lacked suitable data or other necessary information to evaluate our hypotheses; these data and information gaps are described in Chapter 9. Chapter 9 also includes a brief review of some of the more complex mathematical analyses used in recent peer-reviewed publications, such approaches currently being used by others, and three examples of additional mathematical modeling approaches that can be used to further explore some of the linkages and interactions in our conceptual model and complement previously published and other ongoing mathematical modeling efforts for Delta Smelt.

Comparative Approach

The comparative approach used for evaluating the hypotheses stated in this Chapter is similar to the approach taken in the FLaSH investigation (Brown et al. 2014, see also http://deltacouncil.ca.gov/science-program/fall-low-salinity-habitat-flash-studies-and-adaptive-management-plan-review-0). This allowed us to place the results of the FLaSH investigation in a year-round, life cycle context as recommended by the FLaSH Panel (FLaSH Panel 2012). Specifically, we compared data from the two most recent wet years, 2006 and 2011, and the two years that immediately preceded them, 2005 and 2010. To conduct our comparisons, we determined how Delta Smelt responses or habitat attributes would be expected to respond in the different years and then compared the expected response to the observed response. If the expected and observed responses were similar, the hypothesis was considered to be supported.

Moderate to wet hydrological conditions tend to benefit many estuarine organisms, including Delta Smelt (Sommer et al. 2007). But low recruitment or low survival at any point in the predominantly annual Delta Smelt life cycle can lead to low abundance even in a wet year. Identifying the reason(s) for low abundance in a wet year may give important insights into key habitat attributes and environmental drivers that could be managed in a way that would improve the likelihood of abundance increases in all wet years.
The two wettest years after the onset of the POD were 2006 and 2011 (Fig. 58). Delta Smelt abundance increased substantially in 2011, but not in 2006 (Fig. 3). The failure of the Delta Smelt population to increase in the wet year 2006 and the increase of Delta Smelt in the wet year 2011 provides an opportunity to compare and contrast habitat attributes in these two years and possibly identify new options for management actions. As stated in Chapter 3, our working assumption is that different Delta Smelt abundances in 2006 and 2011 should be attributable to differing environmental conditions, in some cases attributable to management actions, and subsequent ecological processes influencing the Delta Smelt population.

preceding habitat conditions may have important implications for the response of a population to the environmental conditions present during a wet year; therefore, we also consider data from 2005 and 2010. Further, we also consider adult and larval abundance in 2012 following the wet year of 2011. We did not include any years predating the POD period in this analysis. This was done to prevent the possibly more subtle, but management-relevant, environmental changes occurring during the POD period from being overwhelmed by effects of the strong POD step changes in the early 2000s as well as similarly strong changes that occurred before the POD (e.g., after the invasion of the clam \textit{Potamocorbula amurensis}).

For the purpose of this report, we call 2005, 2006, 2010, and 2011 our “study years.” We use “year” rather loosely because the Delta Smelt life cycle does not follow the calendar year. As already explained, life stages can overlap and can be observed during different months in different years. Mature adults of a cohort produced in one year are generally not observed until the following year. Similarly, the life cycle does not strictly follow the water year type. We do our best to explain these mismatches when they occur and keep the presentation focused on the life cycle and the conceptual models.

Note that we do not examine the complex interactions that may occur when more than one hypothesis is true (or false), nor do we rule out that a hypothesis may be true in some years and false in others. Therefore, it is important to recognize that data contrary to a hypothesis may indicate that the habitat attribute was not controlling in the selected years, or that complex interactions among multiple habitat attributes (and corresponding hypotheses) contributed to the observed effects. Addressing such complexities is more appropriate for quantitative models as discussed in Chapter 9.

**Hydrological Conditions**

According to annual water year indices and classifications for overall hydrological conditions in the Sacramento and San Joaquin Valleys that provide the freshwater inflow into the Delta, 2005, 2006 and 2011 were the wettest years of the POD period (Fig. 58, see also \url{http://cdec.water.ca.gov/cgi-progs/iodir/WSIHIHST}). In the San Joaquin Valley, 2010 was the fourth wettest year of this period. In the Sacramento Valley, 2003 and 2004 were wetter than 2010. Specifically, water year 2010 was classified as “below normal” in the Sacramento Valley and “above normal” in the San Joaquin Valley and 2011 was classified as wet in both areas, according to the water year index classifications. Water year 2005 was classified as “above normal” in the Sacramento Valley and “wet” in the San Joaquin Valley and 2006 was classified as wet in both areas. (Fig. 58). Water year 2012 was classified as “below normal” in the Sacramento Valley and “dry” in the San Joaquin Valley.
Figure 58. Annual water year indices for the a) Sacramento and b) San Joaquin Valleys since the initiation of the Summer Townet Survey in 1959. Horizontal dashed lines: threshold levels for water year type classifications as wet (W), above normal (AN), below normal (BN), dry (D) and critically dry (C). Darker grey bars indicate the four study years (2005, 2006, 2010, 2011) examined in Chapter 7 of this report. (Data are from http://cdec.water.ca.gov/cgi-progs/iodir/WSIHIST).

The overall wet hydrological conditions in the Sacramento and San Joaquin Valleys in 2005-6 and 2010-11 resulted in relatively prolonged periods of high Delta inflow and outflow and low X2 values in the winter and spring months of the four study years (Fig. 59). In the first half of the year, 2006 had the highest outflow and lowest X2 values followed by 2011, 2005, and 2010. In the second half of 2011, outflow was higher and X2 values were lower than in the second half of 2006 and of all other years during the POD period. In spite of having the lowest spring X2, 2006
had the highest fall X2 (September to October) of all study years, followed by 2005, 2010, and 2011 (Fig. 60).

The overall high flows during these four years allowed for periods of very high fresh water exports from the Delta (Fig. 59). This led to record high volumes of fresh water exported in water year 2011 (6.7 maf) and in water year 2005 (6.5 maf) and a somewhat lower export volume in water year 2006 (6.3 maf). The total water export volume was substantially lower in water year 2010 (4.8 maf) because 2010 immediately followed a three-year drought and the below normal hydrological conditions in the Sacramento Valley (Fig. 58) were not sufficient to rapidly replenish reservoirs and allow for greater exports.

**Hypotheses**

Individual hypotheses are indicated in the life stage transition conceptual model diagrams next to the arrows depicting each hypothesized linkage or outcome (figs. 46-49). While all linkages are considered important, we only developed hypotheses for selected linkages. We developed hypotheses for linkages with sufficient data for quantitative assessments and where there is disagreement or uncertainty regarding the outcome resulting from a driver. We also developed hypotheses for linkages considered important but where we found critical information was missing; thus, highlighting topics where new work is needed. For each of these hypotheses, we then considered the available data to examine whether the Delta Smelt response expected under the hypothesis was consistent with the observed trends in habitat attributes or population dynamics. While we would have liked to test hypotheses about the linkages between habitat attributes and the specific life stage transition processes shown in the life stage transition conceptual model diagrams, the available data often only allowed us to test “lower tier” hypotheses about the linkages between ecosystem drivers and habitat attributes.

Note that we have not examined the complex interactions that may have occurred when more than one hypothesis was true (or false), nor have we ruled out that a hypothesis may be true in some years and false in others. Therefore, it is important to recognize that data contrary to a hypothesis may indicate that the habitat attribute was not controlling in the selected years, or that complex interactions among multiple habitat attributes (and corresponding hypotheses) contributed to the observed effects. Addressing such complexities is likely more appropriate for quantitative models as discussed in Chapter 9. Our overall objective in this Chapter is to provide a demonstration of how the conceptual model can be used to generate and test hypotheses and highlight data gaps while addressing a specific topic of management interest—the increased Delta Smelt abundance index in 2011.

**Adult Hypotheses**

**Hypothesis 1: Hydrology and water exports interact to influence entrainment risk for adult Delta Smelt.**

As discussed earlier, we do not currently have a reliable measure of actual entrainment of fishes by the SWP and CVP export pumps. We also do not have actual population abundance estimates for Delta Smelt. As discussed by Kimmerer (2008, 2011) and Miller (2011), it is thus difficult to estimate proportional population losses due to entrainment. We consider the published
Figure 59. Net daily flows in cubic feet per second for a) Delta inflow from all tributaries, b) Delta outflow into Suisun Bay, and d) total freshwater exports from the Delta. Also shown are daily values for c) X2 (see Chapter 4 for explanation). Flow data are from Dayflow (http://www.water.ca.gov/dayflow/). X2 values are calculated from daily Delta outflow with the equation in Jassby et al. (1995.)
proportional loss estimates for adult Delta Smelt entrainment losses for the two years for which they are available (2005 and 2006; Kimmerer 2008). However, we otherwise restrict our analysis – and this hypothesis – to an assessment of entrainment risk based on salvage and OMR flow data. Note that high entrainment risk for an individual fish does not automatically lead to a high proportion of the population lost to entrainment mortality. For example, in wetter years when large numbers of fish are present but most of the population is distributed farther away from the pumps, a large number of fish can be entrained but only a small percentage of the entire population.

Adult (December-March) Delta Smelt salvage was highest in 2005 followed by 2006 and 2010 and lowest in 2011 (Fig. 61). In 2005, most salvage occurred in January, while in the other three years it occurred in February and March (Fig. 62). Overall, adult Delta Smelt salvage in the four comparison years was on the very low end of the historical time series starting in 1980 (Fig. 26). On the other hand, the ratio of adult salvage divided by the previous year’s FMWT index was high in 2005 (6th highest on record since 1979), but much lower in 2006 and 2010, and lowest in 2011 (Fig. 26).

Low salvage levels in these years and especially in 2010 and 2011 were not particularly surprising due to the low FMWT levels of the POD years along with more active management of OMR flows for Delta Smelt and salmonid protection after 2008 in accordance with the USFWS (2008) and NMFS (2009) BioOps. For management purposes, the onset of increased

Figure 60. Daily X2 values in January to December for each of the four study years. Seasonal X2 averages are indicated by horizontal lines for spring X2 (February to June), summer X2 (July and August), and fall X2 (September to December). See Fig. 15 for seasonal X2 in other years.
adult Delta Smelt entrainment risk is inferred from distributional patterns of Delta Smelt detected by the SKT survey, Delta Smelt salvage and, more recently, consideration of Delta conditions, including turbidity patterns. Since 2009, net OMR flows during periods of increased adult Delta Smelt entrainment risk are now always less negative than they were in years prior to the BioOps. Prior to 2008, net OMR flows often reached -8,000 to -10,000 cfs (see Fig. 31, Kimmerer 2008, Grimaldo et al. 2009), when outflow was low. An exception to these strongly negative flows occurred during April-May export curtailments associated with the Vernalis Adaptive Management Program (VAMP, 2000-2012). These curtailments were especially pronounced in the first half of the VAMP period (2000-2005). During the four comparison years, winter (December-March) net OMR flows were least negative in 2006 followed by 2011 and 2010 with the most negative net OMR flows in 2005 (Fig. 63). High inflows particularly from the San Joaquin River during 2005, 2006 and 2011 moderated effects of negative OMR flows, while export pumping generally remained high. In 2010 at the end of a three-year drought, there was little water in storage to provide for Delta exports prior to the first substantial inflows in mid-January. Subsequently, export levels had to be curtailed to achieve the desired OMR flows. Average winter-time net flows past Jersey Point on the San Joaquin River were positive in all four study years and greatest in 2006 followed by 2011, 2005, and 2010 (Fig. 63).

Kimmerer (2008) used salvage, OMR flows, and fish survey data to estimate proportional population losses due to entrainment for the years 1995-2006. The years 2005 and 2006 represent some of the lower loss estimates in the years examined by Kimmerer (2008); mean population losses reached up to 22% of the adult population in some years when OMR flows were more negative than -5000 cfs (Kimmerer 2008). Even if Kimmerer’s estimation method provides a potential overestimate of loss (Miller 2011), proportional losses of the adult population were less than 10% in the two years that coincide with our comparison years (2005 ≈ 3%, 2006 ≈ 9%; from Fig. 12 in Kimmerer 2008). These types of proportional loss estimates are not available for

**Figure 61.** Annual adult (December-March) Delta Smelt salvage at the CVP (blue bars) and SWP (green bars) fish protection facilities for 2005-2012.
2010 and 2011, but would likely be even smaller than for 2005 due to less negative OMR flows and fish distributions away from the CVP and SWP pumps. Salvage was also lower in these two years than in 2005 and 2006.

In summary, we conclude that hydrology and water exports do interact to influence entrainment risk for adult Delta Smelt and that adult Delta Smelt entrainment risk during the four comparison years was perhaps higher in 2005 than in the other years, but was low relative to historical levels in all four years.

Hypothesis 2: Hydrology interacting with turbidity affects predation risk for adult Delta Smelt.

At present, we do not have information about differences in actual predation mortality between the comparison years. As with entrainment, we thus limit this hypothesis and our analysis to a general discussion of predation risk. Fully characterizing predation risk is exceptionally complicated, making it difficult to generate simple hypotheses that describe associated losses of all life stages of Delta Smelt. We thus limit our hypotheses about predation risk to a few factors for each life stage. For adults, we consider hydrology and turbidity as well as overlap with predators (next hypothesis).

Because Delta Smelt migrate during higher flow conditions when the water is generally turbid, it is assumed that losses to visual predators are lower or at least not substantially higher during the migration period than during other periods. First flush studies led by the USGS and UC Davis
suggest that Delta Smelt aggregate in the water column away from channel edges during daytime flood tides during upstream migration events (Bennett and Burau 2014), but it is not known if Striped Bass or Sacramento Pikeminnow *Ptychocheilus grandis*, the most likely predators of Delta Smelt in the water column, can detect and exploit these aggregations.

In the winters of 2005, 2006, 2010, and 2011 the highest Secchi depths (lowest turbidity) were found in the freshwater regions of the estuary (< 1 salinity), except for the Cache Slough region in the north Delta which was as turbid as the saltier regions of the estuary (Fig. 64). Winter-time Secchi depths in the freshwater region recorded during the SKT surveys (Fig. 64) were often higher (water clearer) than the average Secchi depths across all IEP EMP monitoring sites during these months since 2003 (about 60 cm) and especially when compared to pre-POD winter Secchi depths (around 50 cm on average) recorded by the EMP (Fig. 25). Winter-time Secchi depths in the other salinity regions were generally lower (water more turbid) than the EMP Secchi depth averages for the POD years and more similar to historical averages. In all four comparison years, predation risk associated with turbidity levels was thus likely not different from the historical risk in the more saline regions and the Cache Slough complex, but possibly higher in the freshwater regions, except for the Cache Slough region.

The salinity region differences were much more pronounced than the interannual differences between the four comparison years. Based on these data, it is not clear that higher flows in 2006 and 2011 contributed to higher turbidity in the winter months. The exception might be near the end of the Delta Smelt spawning season in early April when Secchi depths in the freshwater

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**Figure 63.** Annual average daily net flows for December through March in cubic feet per second (cfs) in Old and Middle River (OMR), past Jersey Point on the lower San Joaquin River (QWEST) and total exports in millions of acre feet (MAF), 2005-2013. Error bars are 1 standard deviation.
region were often substantially lower in the two wetter years 2006 and 2011 than in the two drier years 2005 and 2010 (Fig. 64). This will be discussed further in the report section about larval Delta Smelt. For adults, we conclude that interannual differences in turbidity between the wetter and drier of the four comparison years did not likely contribute substantially to reduced predation risk and increased survival in the two wetter years.

**Hypothesis 3: Predator distribution affects predation risk of adult Delta Smelt**

Spatial and temporal overlap with predators is a likely factor contributing to predation risk for all life stages. At present, we do not have information about how predator distribution varied between our comparison years but it is recognized that adult Delta Smelt could be vulnerable to predation if the distributions of predators and Delta Smelt populations overlapped. As already mentioned, Striped Bass and Sacramento Pikeminnow are the most likely open-water predators of adult Delta Smelt. If Delta Smelt utilize littoral habitats to a greater extent than presently assumed, then increased overlap with the distributions of Largemouth Bass and other centrarchid populations is possible. Results of field studies (Feyrer et al. 2013, Bennett and Burau 2014), described for Adult Hypothesis 2, found that adult Delta Smelt did move nearshore on a tidal basis to avoid displacement or move upstream during the “first flush.” Such movements would increase proximity to shoreline predators like Largemouth Bass, albeit during periods of increased turbidity when such visual predators would be at a disadvantage. Clearly, Hypothesis 2 and Hypothesis 3 are closely linked because predation risk is a function of predator presence and prey vulnerability. More information about predator presence is needed to evaluate this aspect of predation risk.

**Hypothesis 4: Variability in prey availability during winter and spring affects growth and fecundity (eggs per clutch and number of clutches) of female Delta Smelt.**

The hypothesis is that increased food availability leads to not only increased adult survivorship, but also growth, which in turn increases reproductive output (number of eggs per female increases with size; Bennett 2005). In addition, with cooler temperatures and lower metabolic rates, sufficient food resources during winter can contribute to energetically demanding multiple spawning events (three spawns possible in wild fish; L. Damon, CDFW, written communication 2012).

For adult females, the ability to meet the bioenergetic demands of reproductive development with sufficient food consumption may be particularly important for fish that spawn multiple times in a year. Preliminary findings from January through April 2012 indicated that adult Delta Smelt are indeed consuming large prey items, such as amphipods, mysids, and larval fish during their spawning period (Fig. 44) with feeding incidence near 98% for the period (Table 2). For this report, we cannot address whether food limitation is a relevant factor during the late winter-spring spawning period because we do not have sufficient data about adult Delta Smelt feeding, but we hypothesize that it may be a critical issue for spawners that need energy for multiple egg clutches. Evidence in support of this hypothesis comes from the modeling simulation experiment by Rose et al. (2013b) who found that food availability along with water temperature affected fall and winter growth and egg production prior to spawning and ultimately population success.
Based on trajectories in adult fork lengths, it appears that adult growth may have been somewhat higher in 2005 and 2011 than in 2006 and 2010, although differences were not pronounced (Fig. 17) and as noted in Chapter 6, annual fork lengths of Delta Smelt collected in the SKT were similar in the four study years (Fig. 55). From these data we infer that environmental conditions were generally good, supporting both continued growth in length and maturation of eggs, except perhaps in 2010. In 2011, only 13 mature females were collected, so growth estimates are uncertain. In general, the number of mature females collected each year reflected year-class strength as measured by the SKT (Fig. 3), except in 2011 when only 13 ripe or ripening females were collected. Adults may use more energy for egg production than for continued somatic growth, but we do not have data on clutch sizes to evaluate this for the four study years.

Data on prey availability for current IEP sampling locations is also limited. Adult Delta Smelt diet is varied (Fig. 44) and includes pelagic and demersal invertebrates, as well as larval fish. Current mesozooplankton (copepod and cladoceran) and mysid sampling by the EMP
Zooplankton Study and invertebrate sampling by the EMP Benthic Monitoring Study does not sample the full geographic range occupied by adult Delta Smelt, including Cache Slough and the Sacramento River Deep Water Ship Channel. In addition, epibenthic cumaceans and amphipods consumed by Delta Smelt might not be effectively sampled with current methods (substrate grabs using a Ponar dredge), which are more suited to sampling organisms in or attached to the substrate. Amphipods found in stomachs of adult Delta Smelt collected January 2012-May 2012 (Fig. 44) were 95% *Corophium* spp., and of those, 90% were juveniles ranging 0.8 to 1.3 mm in body length. These amphipods are believed to be mostly juvenile *Americorophium spinicorne* and *A. stimpsoni*, which as adults are tube building amphipods (Hazel and Kelley 1966). Dirt, substrate debris, and tube pieces were not found in Delta Smelt stomachs with the amphipods, so it is possible these juveniles amphipods are epibenthic or pelagic prior to settling and building tubes. Size distribution of amphipods collected by the DWR EMP Benthic Monitoring Study is not currently available. The IEP Smelt Larva Survey does collect larval fish data during winter (January-March) over a wide section of the estuary, but comparisons with larval fish consumption by adult Delta Smelt are limited because this survey is still new; it was initiated in 2009.

Data were insufficient to conclusively test the hypothesis that variability in prey availability affects growth and fecundity of adult Delta Smelt. More data are needed on growth, clutch number and size, and prey availability.

**Larval Hypotheses**

**Hypothesis 1: Delta Smelt larvae numbers are positively affected by increased duration of the temperature spawning window**

To evaluate this hypothesis, we developed two water temperature measures. The first is the number of days in the temperature spawning window as indexed by mean daily water temperatures at Rio Vista between 12 and 20 °C. This temperature range was selected as representing a reasonable balance between the various temperature ranges observed in laboratory
and field studies (Wang 1986, Baskerville-Bridges et al. 2004b, Bennett 2005) and reviewed in earlier sections of this report. Presumably, a longer duration spawning window would result in more repeat spawning for individual females and greater total fecundity. The second water temperature measure is the number of days in the optimal temperature for egg survival to hatch. We referred to Fig. 10a in Bennett (2005) and selected the temperature range of 12-17 °C as optimal for egg survival. As explained in previous sections, adult abundance, based on SKT sampling, peaked in 2012 as the 2011 year-class of Delta Smelt reached maturity (Fig. 3). In contrast, the spawning stock (i.e., 2011 SKT) that produced the 2011 year-class ranked second lowest to 2006 (Fig. 3, Adults). Despite this low level, the 2011 spawning stock produced the highest adult abundance observed to date in 2012. This suggests that adult stock size has not limited subsequent adult recruitment from rebounding to levels comparable to those of immediate pre-POD years (see Fig. 3, Subadult). As mentioned in Chapter 6, this suggests that even a severely depleted adult stock can still produce a substantial number of larvae and a rebound in the Delta Smelt population, albeit with potentially lower genetic variability than before (Fisch et al. 2011). It also suggests that factors acting on the survival of larval, juvenile and later stages have a substantial effect on recruitment of adults, because relatively low larval abundance in 2011, was associated with the high 2012 adult abundance (Fig. 3).

As mentioned in the adult section, mature adult female Delta Smelt appeared to grow throughout the spawning seasons of the years compared, except 2010 (Fig. 17). We used water temperatures at the Rio Vista Bridge as a surrogate for temperatures experienced by spawning Delta Smelt (Fig. 65) and calculated the duration of the spawning window and of optimal temperatures to hatch. We calculated each as the number of days between the date of first achieving the lower temperature and the date of first achieving the upper temperature. The onset of the spawning window occurred earliest in 2010, followed by 2005 and 2011 (Fig. 65; Table 3). The spawning window occurred latest in 2006 (Fig. 65; Table 3). The spawning window was broad in both 2005 and 2010 at 128-129 days, intermediate in 2011 at 113 days (20 °C not achieved until July 4, not shown), and was shortest in 2006 at 85 days (Fig. 65; Table 3). Assuming that female Delta Smelt undergo a 35-day refractory period, based on a 4-5 week refractory period (J. Lindberg, U.C. Davis, personal communication, 2013) between each spawning, even in 2006 three spawning events were possible, assuming fish were mature and ready to spawn at the initiation of the spawning window. In all other years, four spawning events were possible, so this measure does not discriminate among years well. The duration of optimal hatch temperature was also lowest in 2006, but other durations ranked differently across years than did spawning window duration (Table 3).

The data for the four study years do not provide conclusive support for the hypothesis that the duration of the spawning window or duration of optimal hatching temperature affected larval production. Relatively high larval abundance in 2005 was consistent with a long spawning window and moderate duration of optimal hatch temperatures (129 days and 68 days, respectively; not shown). However, 2006 with the shortest spawning window (85 days) and shortest optimal hatch duration among the 4 study years also had relatively good larva abundance (Fig. 3). In contrast, larval abundance was low in 2010 although the spawning window and optimal hatch duration were both relatively long. Other factors likely contributed to poor larval abundance in 2010, because ripening and ripe females were not detected after early April 2010 and female growth through the winter was poor (Fig. 17). Finally, both the spawning window and optimal hatch duration were fairly long in 2011 as compared to 2006, so slightly lower larval production in 2011 is inconsistent with these durations. This hypothesis was not supported.
Hypothesis 2: Increased food availability results in increased larval abundance and survival.

This hypothesis focuses on seasonal changes in phytoplankton biomass and the zooplankton community and resulting changes in abundances of food items most often consumed by Delta Smelt larvae. Phytoplankton biomass data (chlorophyll-a) collected at 10 stations by the IEP.
EMP show that the highest spring biomass levels were observed in May of 2010 and 2011 (Fig. 66). Median biomass levels were lower in April and May of 2005 and 2006 than in April and May of 2010 and 2011. This suggests that more food was available for zooplankton growth in the spring of 2010 and 2011 than in 2005 and 2006. In all four years, however, chlorophyll concentrations were lower than 10 ug/L at almost all stations, suggesting that zooplankton may have generally been food limited in these years (see Chapter 4). Nevertheless, greater phytoplankton biomass in late spring of 2010 and 2011 may have contributed to overall greater food availability and better survival of late larvae and early juveniles in these years.

Juvenile and adult calanoid copepods, particularly *E. affinis* and *P. forbesi*, comprise most of the larval diet through June (Nobriga 2002, Slater and Baxter 2014). *E. affinis* is moderately abundant only during winter and spring and rare in summer and fall, whereas *P. forbesi* is abundant only in summer and fall (Durand 2010, Hennessy 2010, 2011, Winder and Jassby 2011). It is not clear whether the seasonal decline in abundance of *E. affinis* is related to temperature, potential competitive interactions with *P. forbesi*, differences between the species in vulnerability to consumption by *P. amurenensis* (Miller and Stillman 2013), or a combination of such factors. The transition between high abundances of the two species, may create a seasonal “food gap” during late spring or early summer. This food gap has been hypothesized to be an important period for Delta Smelt larval survival (Bennett 2005, Miller et al. 2012).

To assess whether a gap in prey availability existed between periods of high abundance of *E. affinis* and *P. forbesi*, we evaluated abundance patterns in 20 mm Survey copepod data for stations with and without Delta Smelt. The food gap hypothesis was only weakly supported by the data. The density of *E. affinis* (in the presence of Delta Smelt larvae) typically reached 100 m$^3$ by week 16 (Figs. 67 and 68). Assuming 100 m$^3$ as a baseline density for *E. affinis*, this baseline was generally maintained until about week 22, when they declined at about the same time that *P. forbesi* densities increased to 100 m$^3$ (Figs. 67 and 68). After combining the densities of both *E. affinis* and *P. forbesi* and tracking them through time, we detected a gap in food during week 22 (late May – early June) of 2005 (Fig. 67), which is inconsistent with 2005 exhibiting the highest larva abundance among our comparison years (Fig. 3). Such density gaps were not observed in the other three comparison years (Figs. 67 and 68), which exhibited lower abundance than 2005 (Fig. 3). Survival of larvae to juveniles was very low in 2005, but was also low in 2006 (Fig. 51) with no evidence for a food gap in 2006. Survival of larvae to juveniles was relatively high in 2010 and 2011 (Fig. 51). This analysis does not support the hypothesis that differences in zooplankton availability affected larval abundance and survival in the four study years, but higher phytoplankton biomass in April and May of 2010 and 2011 could have contributed to overall greater food availability and better survival of late larvae and early juveniles in these years.

**Hypothesis 3: Distributional overlap of Mississippi Silverside with Delta Smelt and high abundance of Mississippi Silverside increases predation risk/rate on larval Delta Smelt, whereas, increased turbidity, decreases predation risk/rate on larval Delta Smelt.**

Silversides are ubiquitous within the Delta (Brown and May 2006) and have long been proposed (Bennett 1995) and more recently confirmed as a predator of Delta Smelt larvae (Baerwald et al. 2012). We do not have estimates of predation losses to Silversides during the four study years and thus focus on assessing predation risk by evaluating fish distributions, predator and prey sizes, and prey growth, which is related to temperature.
Silversides large enough to consume fish larvae are present in the Delta during spring and are likely to prey upon Delta Smelt larvae. Silverside habitat has been characterized as open water shoals and shoreline (Brown and May 2006, Grimaldo et al. 2012); however, the species also occurs in low density in deep open water primarily in summer (Grimaldo et al. 2012). Catches in the SKT confirm silverside presence in open water in spring as well, though catches tended to be low. However, SKT sampling does not occur at night when offshore Silverside densities may be higher, if foraging patterns follow those observed in Clear Lake, California (see Wurtsbaugh and Li 1985). Compared to the open embayments, SKT Silverside catches were higher in channels such as Montezuma Slough, Cache Slough, the San Joaquin River, and especially the Sacramento Deepwater Ship Channel (Table 4). This Silverside distribution matched higher March through May regional catches of Delta Smelt larvae (Table 4, see http://www.dfg.ca.gov/delta/data/20mm/CPUE_map.asp), except that larvae catches in Suisun Bay and the lower Sacramento River were occasionally high and Silversides catches were usually low. Delta Smelt larvae were found in significantly higher densities in offshore-open water habitats (Grimaldo et al. 2004), which corresponds to the habitat where Silversides consuming Delta Smelt larvae were captured (Baerwald et al. 2012). As discussed above, the relatively large-sized silversides present in the Spring Kodiak Trawl indicates some offshore movement and overlap of predator-sized foraging silversides with Delta Smelt larval habitat.

The frequency and magnitude of Silverside catches by the Spring Kodiak Trawl increased as Secchi depths approached and dropped below 50 cm (Fig. 69), suggesting that Silversides may venture offshore more frequently and in higher numbers in turbid water. This might also represent a displacement effect resulting from high flows, but high catches were most common in Montezuma Slough and the Sacramento Deepwater Ship Channel (Table 4) where displacement by flow should not have been a factor.

The hypothesis is somewhat supported in that: 1) Silversides are captured in Spring Kodiak Trawl in March and April (Fig. 70), when early stage Delta Smelt larvae are common; 2) Silverside...
catches offshore increase with increased turbidity (i.e., declining Secchi depth; Fig. 69), and 3) there is regional overlap in Cache Slough and the Sacramento Deepwater Ship Channel, and some in Montezuma Slough (cf. Table 4 and http://www.dfg.ca.gov/delta/data/20mm/CPUE_
map.asp), known larval rearing regions. It is also possible the nighttime offshore foraging by silversides is a more common strategy (Wurtsbaugh and Li 1985), but one that goes undetected by current sampling. Silverside catch per trawl (Table 4) indicates low offshore densities and the same turbidity that facilitates offshore movement may also inhibit predation effectiveness.
Overall, the conclusion regarding the effects of species distributions and abundances on predation risk is unclear. If there is an effect, it is most likely to occur in smaller channels, such as Montezuma Slough and those in the Cache Slough and the Sacramento Deepwater Ship Channel where Silversides are present in high numbers along the shoreline and larval Delta Smelt occur offshore.

Hypothesis 4: Hydrology and water exports interact with one another to influence direction of transport and risk of entrainment for larval Delta Smelt.

As for adults, we do not have proportional entrainment estimates for all four study years, so the entrainment portion of this hypothesis cannot be directly evaluated. Also, larvae (< 20 mm fork length) entrained in the State and federal water export systems are generally not quantified. To test this hypothesis we use data for the distribution and density of larvae (≥ 20 mm fork length)
in the central and south Delta and estimates of channel flows to infer risk of entrainment. Among the study years only 2005 larval entrainment was estimated by Kimmerer (2008), and loss to the population was relatively low. However, Delta Smelt density and distribution in the central and south Delta were greater in 2005 than in the three other study years (Table 5). This simple analysis suggests that in our 4-year comparison, entrainment risk for larval Delta Smelt may have been highest in 2005. Hardly any larval Delta Smelt were caught in this region in the two wet years, 2006 and 2011.

As for adults, we also used OMR flows (Fig. 31) to assess larval entrainment risk. Mean March through May OMR flows were positive during the two wet years 2006 and 2011 (8,221 cfs and 3,560 cfs respectively) and negative during the two dry years 2005 and 2010 (-417 cfs and -1,302 cfs, respectively). These OMR values suggest little if any risk during 2006 and 2011, and at most moderate risk in 2005 and 2010. Grimaldo et al. (2009) found that juvenile salvage was a function of abundance in the 20 mm Survey (positive) and OMR flows (negative). Looking more closely at various net daily flows from March to June of 2005, we find that OMR flows were moderately negative (i.e., toward the export pumps) only in March, and were zero to weakly positive in April and May, except for a brief period in mid-April (Fig. 31); also in 2005, Qwest was strongly positive from late March through early June, promoting downstream transport in the San Joaquin River, and exports were low from late April through late May (Fig. 31). The other dry year, 2010 exhibited a similar pattern, but lower inflows resulted in the magnitude of exports more directly influencing OMR flows (Fig. 31), and leading to moderately negative OMR flows.
in March and again in June, but only weakly negative flows in April and most of May coincident with positive Qwest. In the high outflow years 2006 and 2011, few larvae were detected in the central or south Delta (Table 5) and Qwest flows were strongly positive from March through at least early June, while OMR flows were near zero or weakly negative in March and positive to strongly positive by April and continuing to early June of both years (Fig. 31). Thus, for our comparison years, it appears that the available data generally support our hypothesis, but entrainment of larvae was unlikely to be an important factor during either wet year and was probably not a substantial factor in either dry year.
### Table 5. Mean monthly catch of Delta Smelt per 10,000 m$^3$ by station for stations in the south and central Delta for the 20 mm Survey, 2005, 2006, 2010, 2011. Non-zero values are bolded.

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| Year = 2006 |
|-------------|---------|---|---|---|---|
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| 812         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 815         | 0.00   | 0.00  | 1.24| 0.00 | 0.00 |
| 901         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 902         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 906         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 910         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 912         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 914         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 915         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 918         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
| 919         | 0.00   | 0.00  | 0.00| 0.00 | 0.00 |
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|-------------|--------|--------|--------|--------|
| STATION     | MARCH  | APRIL  | MAY    | JUNE   | JULY   |
| 809         | 0.00   | 0.00   | **1.62** | 0.00   | 0.00   |
| 812         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 815         | 0.00   | **1.77** | **1.72** | 0.00   | 0.00   |
| 901         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 902         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 906         | 0.00   | **3.36** | 0.00   | **1.64** | 0.00   |
| 910         | 0.00   | **5.24** | 0.00   | 0.00   | 0.00   |
| 912         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 914         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 915         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 918         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 919         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| Year = 2011 | Months | | | | |
| STATION     | MARCH  | APRIL  | MAY    | JUNE   | JULY   |
| 809         | 0.00   | 0.00   | 0.00   | **1.73** | 0.00   |
| 812         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 815         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 901         | 0.00   | 0.00   | **3.69** | 0.00   | 0.00   |
| 902         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 906         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 910         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 912         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 914         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 915         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 918         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
| 919         | 0.00   | 0.00   | 0.00   | 0.00   | 0.00   |
Juvenile Hypotheses

Hypothesis 1: High water temperatures reduce juvenile Delta Smelt growth and survival through lethal and sublethal (bioenergetic stress; reduced distribution) effects.

High water temperatures have a strong effect on juvenile Delta Smelt survival (Swanson et al. 2000, Komoroske et al. 2014). In addition to the obvious potential for lethal effects, temperature can have sub-lethal effects such as reduced habitat area, higher food requirements, increased susceptibility to disease and contaminants, and increased predation. The potential for increased prey requirements and increased predation is described below for other hypotheses.

As noted in the adult section, spring water temperature was generally coolest in 2006 and 2011, but warmed up more rapidly toward the end of spring 2006 (May) than in spring 2011. Spring water temperature was overall warmest in 2005 (Fig. 71). Following the high late-spring water temperatures in 2005 and 2006, summer temperatures in 2005 and 2006 tended to be higher than in 2010 and 2011 during July and August (e.g. TNS surveys 3-5; Fig. 72). Temperatures during surveys 4 and 5 may have been particularly important as they exceeded lethal levels in freshwater at some sites, suggesting the potential for mortality. Note that this does not mean that temperatures were universally cooler in 2010 and 2011 than in 2005 and 2006; for example the region around Cache Slough had relatively high temperatures in August 2011. Larval to juvenile survival (ratio of TNS index to 20 mm index) was highest in 2011 followed by 2010, 2006, and 2005, suggesting that the cooler late spring and summer temperatures in 2011 and 2010 may have been beneficial for Delta Smelt. However, juvenile to subadult survival (ratio of FMWT index to TNS index) was highest in 2011 and lowest in 2010 (Fig. 51). While relatively high water temperature in late spring and early to mid summer of of 2005 and 2006 may thus have contributed to low survival of late-stage larvae and early juveniles, water temperature may have been less important to survival in the late summer and early fall. Overall, the results of this analysis of temperature and survival data support our hypothesis that high water temperatures reduce juvenile Delta Smelt growth and survival.

At this point, our data and analyses are inadequate to address temperature effects on juvenile Delta Smelt growth. Although there are some data for Delta Smelt growth during several of the target years, it is difficult to separate the relative effects of improved bioenergetics (see below) versus simple ontogenetic changes in fish size. Juvenile fish growth rates are typically not constant and change with size (“allometric effects;” Fuiman 1983). Specifically, daily growth rates (e.g., mm/day) are often faster for smaller fish and slower for older fish. Hence, cooler years may delay Delta Smelt transitions from faster to slower growth phases, yielding a relatively fast measured growth rate at a specific point in time (e.g., September) because at that specific time the fish are still relatively young and still on the “steepest” part of an idealized growth curve.

Hypothesis 2. Distribution and abundance of Striped Bass, temperature, and turbidity influence predation risk/rate on juvenile Delta Smelt

We hypothesize that subadult (age 1-3) Striped Bass are the major predator on juvenile Delta Smelt and that losses are likely affected by temperature and turbidity patterns. However, other factors likely affect predation risk (e.g., other predators such as centrarchids) and several factors
may interact. As noted above for temperature and below for food, high temperatures and low prey density likely lead to bioenergetics problems and increased foraging activity, which might reduce predator avoidance behavior (e.g., Marine and Cech 2004) in Delta Smelt. These effects may be compounded by low turbidity, which makes Delta Smelt more visible to predators in their habitat. Although higher Striped Bass abundance could theoretically result in greater consumption of prey including Delta Smelt (Loboschefsky et al. 2012), changes in habitat variables for both species such as food, temperature, and turbidity mean that predation rates on Delta Smelt periodically may be independent of predator abundance. Although there has been substantial progress in modeling (Loboschefsky et al. 2012, Nobriga et al. 2013) and genetic methods (Baerwald et al. 2012), there is not yet a standardized way to assess the effects of predation on Delta Smelt. Moreover, there are no effective surveys to assess age 1-3 Striped Bass abundance or distribution. Therefore, we are unable to directly evaluate this hypothesis. Lacking this information, we can
at least examine turbidity and temperature patterns for the four years. Temperature responses were described for Hypothesis 2. In general, summer 2005 and 2006 temperatures were relatively higher than 2010 and 2011 during key summer months (e.g. TNS surveys 3-5; Fig. 72). We expect that cooler temperatures in 2010 and 2011 may have contributed to reduced predation on Delta Smelt. Turbidity data are limited to 2010 and 2011 (Fig. 73). There were no consistent differences between the two years. Secchi depth data did not suggest major differences among the 4 years except at salinities > 6 when 2005-2006 had higher values in some months (Fig. 74).
Hypothesis 3. Juvenile Delta Smelt growth and survival is affected by food availability.

As for Hypothesis 1, we are currently unable to evaluate the growth data because water temperature affects development time, and because growth curves are complicated by allometric effects. The general conceptual model is that higher food abundance results in faster growth rates and larger, healthier fish. In addition, larger, healthier Delta Smelt are presumably less vulnerable to predators because of increased size making them difficult for smaller predators to capture and consume. In general, the median abundance of some of the key prey for juvenile Delta Smelt such as calanoid copepods is highest in summer months (Fig. 75), when juvenile Delta Smelt are present; however, the range of observed densities is broad in all months. As noted previously, Kimmerer (2008) found that Delta Smelt survival from summer to fall was positively associated with calanoid copepod biomass in the low salinity zone.
Interpretation of the field data is complicated because there are no long-term IEP EMP study stations located in some of the core habitats for Delta Smelt, for example, Cache Slough and the Sacramento River Deep Water Ship Channel. Moreover, densities of calanoid copepods vary among regions based on differing habitat (temperature and salinity) requirements of each species (Fig. 76).

Summer-time phytoplankton data (chlorophyll-a) suggest that the base of the food web was most enhanced in July and August 2011 and relatively depleted in 2005 (Fig. 66). There is some evidence that these changes may have affected zooplankton abundance. For example, summer densities of calanoid copepods in the LSZ and <1 ppt regions also tended to be highest in 2011 as compared to the other years (Fig. 76). This pattern generally held when individual taxa are considered including two of the most important food sources for Delta Smelt, *Eurytemora affinis* (Fig. 33) and *Pseudodiaptomus forbesi* (Fig. 34).
As mentioned above (Hypothesis 1), juvenile to subadult survival was highest in 2011 followed by 2006 and 2005 and lowest in 2010 (Fig. 51). If food availability was the primary habitat attribute driving juvenile survival, our expectation was that summer prey abundance would have been higher in 2011 than 2010. Figure 69 suggests that while differences were not very pronounced, prey levels were indeed somewhat higher in July and August of 2011 than 2010. Calanoid copepod levels varied across the different salinity ranges, but generally followed the same pattern (Fig. 76). In addition, calanoid copepod densities in June and August were higher in 2006 than in 2005 (Fig. 75), which may have contributed to higher juvenile to subadult survival in 2006 compared to 2005 (Fig. 51).

Fish bioenergetics are affected by both food and temperature. As mentioned above, both summer 2010 and 2011 had relatively cool temperatures as compared to 2005 and 2006, which may have affected bioenergetics. In addition, recent studies (S. Slater, CDFW, unpublished data) indicate that Delta Smelt consumption was not just limited to calanoid copepods, so our assessment does not reflect the full dietary range.

In conclusion, our analyses provide some support for the hypothesis that juvenile Delta Smelt growth and survival is affected by food availability; greater food availability may have contributed to greater juvenile survival in 2011 and 2006 compared to 2010 and 2005. However, differences in prey availability among years were not very pronounced and our analyses were limited to calanoid copepods; other species may also be important prey items for Delta Smelt.
Figure 76. Trends in calanoid copepods (number/m³ for all types combined) collected by the IEP Environmental Monitoring Program (EMP) in three salinity ranges (> 6 ppt; 1-6 ppt; < 1 ppt) during each the four study years (2005, 2006, 2010, and 2011). See Chapter 3: Data Analyses for explanation of boxplots.
Hypothesis 4. Juvenile Delta Smelt survival and growth is reduced by harmful algal blooms (HAB) because of direct (habitat quality and toxic effects) and indirect (food quality and quantity) effects.

The appearance of late-summer HAB, especially *Microcystis*, is thought to be another component of the decline in habitat quality for Delta Smelt (Baxter et al. 2010, Lehman et al. 2010). Direct effects may include toxicity to Delta Smelt and a reduced area of suitable habitat. There also may be indirect effects on food quantity and quality, particularly with respect to their zooplankton prey (Ger et al. 2009, 2010a,b, Lehman et al. 2010).

The growth responses of Delta Smelt during the four target years are still unclear (see below), but there is evidence that Delta Smelt juvenile to subadult survival was highest in 2011 and lowest in 2010 (Fig. 51). If HABs have a negative effect on survival, we would expect that lower *Microcystis* (or other HAB) abundance would be associated with higher survival in 2011. This seems to have been the case for 2010 and 2011. Densities of *Microcystis* near the water surface were qualitatively assessed (visually ranked) at all TNS stations in these years. In agreement with our expectation, observed levels were low during the TNS in 2011 as compared to 2010 across a range of salinities (Fig. 77).

Unfortunately, we do not have data about other HAB species and more quantitative estimates, nor is similar data available for 2005 and 2006. In general, our expectation is that 2006
Microcystis levels would have been relatively low as a result of higher flow levels that discourage blooms (Lehman et al. 2005). Based on the available qualitative data for 2010 and 2011, this analysis supports the hypothesis that juvenile Delta Smelt survival and growth is better when Microcystis does not bloom as intensely, but more data is needed to more conclusively assess this relationship.

Subadult Hypotheses

Hypothesis 1. Subadult Delta Smelt abundance, growth, and survival is affected by food availability.

Similar to juveniles, the general conceptual model is that higher food abundance results in faster growth rates and subsequently, lower predation loss and greater survival (e.g., Houde 1987, Sogard 1997, Takasuka et al. 2003); however the opposite situation in which the fastest growing fishes are most vulnerable to predators has also been observed in at least one east coast estuary (Gleason and Bengston 1996). Fall abundance of Delta Smelt was highest in 2011 followed by 2006, 2010, and 2005 (Fig. 3) while survival of subadults to adults was highest in 2010 followed by 2006 and equal in 2011 and 2005 (Fig. 45). In spite of the lower subadult survival in 2011, the relatively large number of subadults in 2011 gave rise to the highest adult abundance on record in 2012.

In general, fall calanoid copepod abundance and cladocera abundance were higher in 2011 in freshwater and the low-salinity zone compared to the other years, particularly 2005 and 2006 (Fig. 71). However, these data are highly variable, so this conclusion does not apply to each region in every month. With that caveat, the data generally support the hypothesis that food availability affects Delta Smelt abundance and survival; on average, prey density was higher for subadult Delta Smelt in 2011. This may have contributed to the high FMWT abundance index in 2011, although it did not contribute to an equally high survival to adults relative to the other three years. Nevertheless, it seems likely that the relatively good food availability in 2011 also contributed to the high number of adults in 2012. As noted above, we are currently unable to evaluate whether Delta Smelt grew faster in 2011 because water temperature affects spawning and hatch dates, which complicates the interpretation of growth rates.

Hypothesis 2. Distribution and abundance of Striped Bass, temperature, and turbidity influence predation risk/rate on subadult Delta Smelt

As already described for other life stages, predation risk is exceptionally complicated, making it difficult to generate simple hypotheses that describe associated losses of Delta Smelt. The data are not currently available to test this hypothesis (Nobriga et al. 2013). Thus, no firm conclusion can be made.
Hypothesis 3. Subadult Delta Smelt abundance, survival and growth are reduced by harmful algal blooms (HAB) because of direct (habitat quality and toxic effects) and indirect (food quality and quantity) effects.

The appearance of late-summer harmful algal blooms (HAB), especially *Microcystis*, is thought to be another detriment to habitat quality for Delta Smelt (Baxter et al. 2010, Lehman et al. 2010). Direct effects may include toxicity to Delta Smelt and a reduced distribution if the fish try to limit their overlap with the bloom. There also may be indirect effects on food quantity and quality, particularly with respect to their zooplankton prey (Ger et al. 2009; 2010a,b, Lehman et al. 2010).

The growth responses of Delta Smelt during the four target years are still unclear (see above), but there is evidence that summer juvenile to subadult survival was highest in 2011, while juvenile survival to adults was highest in 2010 (Fig. 45). Our expectation is therefore that HAB were less prevalent in the summer of 2011 compared to 2010, but more prevalent in fall 2011. As already described for juveniles, the hypothesis that summer *Microcystis* bloom would be less intense in 2011 compared to 2010 was generally supported (Fig. 77). In fall, *Microcystis* levels were also overall lower in 2011 than in 2010, except in September 2011 when a high level of *Microcystis* was observed in the LSZ (Fig. 78). This may be an indication that the higher outflow in September-October 2011 displaced *Microcystis* produced in the Delta seaward into the LSZ. The comparatively high 2011 Delta Smelt FMWT index that coincided with this shift in *Microcystis* distribution is not consistent with the hypothesis; however, the occurrence of fairly high levels of *Microcystis* in the LSZ in 2011 may help explain the lower subadult to adult survival in 2011 compared to 2010. It is also important to remember that the visual survey results presented here are only qualitative and do not necessarily reflect the potential for differences in actual toxicity among years. Overall, these results are inconclusive, although they may provide limited support for the hypothesis that high *Microcystis* levels may have a negative effect on subadult to adult survival; this may help explain the lower subadult survival in 2011 compared to 2010.

Hypothesis 4. Subadult Delta Smelt abundance, survival and growth are affected by the size and position of the low salinity zone during fall.

We do not address this hypothesis in detail because it is the subject of an adaptive management experiment (FLaSH) described earlier (Reclamation 2011, 2012; see also Brown et al. 2014, http://deltacouncil.ca.gov/science-program/fall-low-salinity-habitat-flash-studies-and-adaptive-management-plan-review-0). According to the FLaSH conceptual model, conditions are supposed to be favorable for Delta Smelt when fall X2 is approximately 74 km or less, unfavorable when X2 is approximately 85 km or greater, and intermediate in between (Reclamation 2011, 2012). Surface area for the LSZ at X2s of 74 km and 85 km were predicted to be 4000 and 9000 hectares, respectively (Reclamation 2011, 2012). The data generally supported the idea that lower X2 and greater area of the LSZ would support more subadult Delta Smelt (Table 6). The greatest LSZ area and lowest X2 occurred in September and October 2011 and were associated with a high FMWT index which was followed by the highest SKT index on record, although survival from subadults to adults was actually lower in 2011 than in 2010 and 2006. There was little separation between the other years on the basis of X2, LSZ area, or FMWT index (Table 6). The position and area of the LSZ is a key factor determining the quantity and quality of low salinity rearing habitat available to Delta Smelt and other estuarine species (see Chapter 4 for more detail.
and Chapter 8 for additional analysis results). In addition, the complex hydrodynamics produced during higher outflows may alter the lateral mixing environment of the Estuary (especially in shallower areas like Suisun Bay) in ways that improve the quality of Delta Smelt habitat in general (Monismith, personal communication). The limited amount of available data provides some evidence in support of this hypothesis, but additional years of data and investigations are needed.

Chapter 8: Conclusions

As with all reports focusing on conceptual models, this report is intended as a working document, not as the final word on Delta Smelt ecology, because our knowledge will continue to increase. We intend the conceptual model to be used as a framework and tool to further improve our understanding of Delta Smelt ecology and to explore and test management options for improving conditions for the Delta Smelt population. In essence, the updated conceptual model represents a synthesis of our current thinking on the factors affecting vital rates of the Delta Smelt population. We fully expect a wide range of opinion about the relevance of the conceptual models presented here and about the degree of certainty regarding many of its component dynamics and linkages. We have clearly acknowledged that we lack information on many important factors and processes that likely affect Delta Smelt, such as predation and toxicity and their functional relationships.

![Figure 78. Fall Midwater Trawl mean visual rank of Microcystis spp. (ranks 1-5 possible; 1 = absent) observed at all stations during monthly surveys in various salinity regions (> 6, 1-6, and < 1 ppt) and in the CS-SRDWSC during September through December 2010 and 2011.](image)
Table 6. Mean and standard deviation (SD) for X2, surface area of low salinity zone (M. McWilliams, Delta Modeling Associates, unpublished data), and values of the Fall Midwater Trawl index (FMWT) for abundance of subadult Delta Smelt.

<table>
<thead>
<tr>
<th>YEAR</th>
<th>X2 (km)</th>
<th>Surface area LSZ (hectares)</th>
<th>FMWT index</th>
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<tr>
<td></td>
<td>MEAN</td>
<td>SD</td>
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<tr>
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<td>2</td>
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<tr>
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<td>8366</td>
</tr>
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</table>

with survival and growth. The conceptual model incorporates many hypotheses that should be tested via new research, modeling, and ongoing analysis and synthesis of new and previously collected data. This is how science advances.

Conceptual models are increasingly used as tools to develop questions or hypotheses about specific mechanisms through which stressors or other environmental factors drive ecological outcomes. Conceptual models can be used as a basis for communication among managers and scientists to plan research activities and assess outcomes of management actions (Ogden et al. 2005). Because of their broad utility, conceptual models are viewed as a critical element of adaptive management programs (Thom 2000). In the SFE, conceptual models have become common and even required as the community moves toward adaptive management and collaborative science. A primary outcome of conceptual models is the identification of key areas of uncertainty due to lack of information, or areas of disagreement due to different interpretations of the available data and information. Careful examination of these areas often identifies critical data and information gaps, which if filled, would allow a more robust evaluation of the major hypotheses derived from conceptual models. In this way, conceptual models can guide the research community to the topics critical for understanding Delta Smelt biology and formulating effective management actions.

The development of our conceptual model, based on assessment of recent information, identified some key points about conceptual models that are worth highlighting, including the following:

1. Nested and linked conceptual models of increasing specificity provide a useful framework for capturing the dynamics of ecosystem drivers and habitat attributes over a large range of temporal and spatial scales and for providing a comprehensive picture about their effects.

2. Our knowledge about Delta Smelt and the SFE is constantly growing and conceptual models about them have to be regularly updated and revised to properly reflect this knowledge.

3. Construction of our conceptual model and the formulation and evaluation of hypotheses greatly benefitted from the large amount of high-quality ecological data and information available about Delta Smelt and the SFE. The most critical data about Delta Smelt dynamics came from four long-term IEP fish monitoring surveys. Other monitoring
and studies provided key data and information about habitat attributes and ecosystem drivers.

4. Our conceptual model is also useful for identifying important data and information gaps. More data and information is especially needed about predation risk and toxicity, two potentially important attributes of Delta Smelt habitat.

Conceptual models are meant to be useful tools for scientists, managers, and others. But just how useful are the new conceptual models in this report? To find out, we used them to generate and test hypotheses and highlight data gaps while addressing a specific topic of high management interest—the increased Delta Smelt abundance index in 2011.

We found that our conceptual model allowed us to formulate a variety of testable hypotheses about individual components and the linkages among them. Our hypotheses and the analyses we conducted to test them had some clear limitations (discussed below), but highlighted some key points about Delta Smelt and their habitat. In many respects, the points about Delta Smelt seem self-evident from basic biology and earlier conceptual models, but they warrant reinforcement because they are crucial to understanding Delta Smelt and to developing and assessing habitat management actions. Key points about Delta Smelt include the following:

1. Environmental conditions occurring in all four seasons contribute to year-class strength of Delta Smelt—"it takes a year to make a mature Delta Smelt."

2. Survival and recruitment are affected by many factors that interact in complex ways and the importance of these factors and interactions varies from season to season and year to year.

3. Recovery of Delta Smelt depends on better than average larval production (recruitment) and survival in all seasons. The number of eggs and larvae sets an upper limit for the production of mature adults. Low survival between any two life stages can substantially reduce the actual production of mature adults. Success of Delta Smelt in 2011 was related to a high level of larval production (recruitment) followed by moderate to high stage-to-stage survival over the entire year. In contrast, the high level of larval production (recruitment) in 2006 was followed by very low survival from larvae to juveniles which led to low abundance of mature adults.

4. Throughout 2011, Delta Smelt may have benefitted from a combination of favorable habitat conditions: 1) adults and larvae benefitted from high winter 2010 and spring 2011 outflows which reduced entrainment risk and possibly improved other habitat conditions, prolonged cool spring water temperatures, and possibly good food availability in late spring; 2) juveniles benefitted from cool water temperatures in late spring and early summer as well as from relatively good food availability and low levels of harmful Microcystis; 3) subadults also benefitted from good food availability and from favorable habitat conditions in the large, westward low salinity zone.

Our hypothesis tests were carried out with the simple comparative approach used in the FLaSH investigations (Brown et al. 2014). Specifically, we compared differences in Delta Smelt responses and in individual habitat attributes during the two most recent wet years and the two years immediately preceding the two wet years. Using this approach allowed us to put the FLaSH results into a year-round context as recommended by the FLaSH Panel (FLaSH Panel 2012).
It also provided an opportunity to further assess the utility of this approach for evaluating the outcome of adaptive management actions such as the fall outflow action.

As with the FLaSH investigations (Brown et al. 2014), we restricted our analyses to simple comparisons among four recent years after the 2002 POD decline for several reasons including the following:

1. Using a comparative approach similar to that in the FLaSH investigation allowed us to place the results of the FLaSH investigation in a year-round, life cycle context as recommended by the FLaSH Panel (FLaSH Panel 2012).

2. This report is intended for a broad audience. Simple comparisons are easily replicated and understood by all.

3. More pertinent data is available for recent years than for earlier years. For example, adult Delta Smelt monitoring began in 2002 with abundance index values available starting in 2003.

4. The POD regime shift (Baxter et al. 2010) changed ecological relationships and the strong pre-POD signals would have likely overwhelmed more subtle, yet meaningful, signals in the period after the POD. For example, it appears that high larval recruitment may now be positively associated with wet hydrology, but that this may not have been the case before the onset of the POD.

5. Clear differences in habitat conditions among years might point to new or refined management strategies aimed at improving specific habitat conditions.

6. More complex modeling approaches take much more time and effort than was available to produce this report. A complex life cycle modeling effort is currently underway (see Chapter 9).

As noted above, our analytical approach yielded some interesting results, but it also raised more questions than it could answer. In many cases this was due to critical data and information gaps; these will be described in more detail in Chapter 9. It also illustrates, however, several limitations of our simple comparative approach as well as difficulties associated with posing and testing hypotheses about ecological phenomena in general. Examples of specific limitations and difficulties include the following:

1. Our hypotheses focused on individual habitat attributes and were tested with a series of separate univariate analyses even though we know that Delta Smelt are affected by multiple interacting habitat attributes. We did not conduct multivariate tests or examine the complex interactions that may have occurred when more than one hypothesis was true (or false), nor did we consider or rule out that a hypothesis may be true in some years and false in others.

2. Our simple comparisons of differences in individual habitat attributes among different years cannot conclusively establish whether these differences are indeed mechanistically linked to the observed differences in Delta Smelt dynamics. In addition, an absence of observed differences does not prove that there is really no effect because actual effects can be masked or counteracted by interactions with other causal factors that differ among years. For example predation in the South Delta may mask actual entrainment
effects and toxicity of anthropogenic contaminants may counteract the effects of abundant food in some years, but not in others.

3. Results contrary to our observations may simply indicate different outcomes in other years or that complex interactions among multiple habitat attributes (and corresponding hypotheses) contributed to the observed effects.

4. We restricted our analyses to observational data collected in a small number of moderately and very wet years during the POD period; including data from additional, more historical, and drier years may have provided more conclusive results.

5. Data available for our analyses were not necessarily collected to test hypotheses similar to the ones in this report; targeted data collections are needed in addition to routine status and trends monitoring.

Many of these difficulties and limitations were expected because hypothesis testing in an ecological context is nearly always problematic. For example, Quinn and Dunham (1983) warned that attempts to follow a strictly hypothetico-deductive scheme (Popper 1959, Platt 1964) to draw “strong inference” from a series of univariate tests aiming to falsify hypotheses about the ecological effects of individual causal factors often lead to inconclusive or even erroneous results. One reason for this is that by design, they generally do not consider non-additive interactions among causal factors. While we did not necessarily set out to strictly follow such a scheme, we nevertheless treated habitat attributes as largely independent from each other and formulated a series of distinct hypotheses about their univariate effects on Delta Smelt. But habitat attributes are not necessarily additive and habitat is indeed more than the “sum of its parts.” A more inductive, multivariate modeling approach with hypotheses about interactive effects and evaluations of the relative contributions of multiple interacting habitat attributes to these effects would have likely been more appropriate, but would have required analyses beyond the scope of this report.

We give some examples of multivariate approaches in Chapter 9, but note that even with the most sophisticated modeling techniques, ecological responses to management manipulations and other changes of the SFE have been notoriously difficult to assess and interpret. Reasons for this persistent difficulty include limited opportunities for experimental control, multiple interacting causal factors, multiple ecological response pathways, and changing environmental conditions due to species invasions, species declines, and the many physical and chemical changes and management manipulations described in this report. In other words, the signal to noise ratio of management actions to environmental variation tends to be low in the SFE because of its size and complexity. The fact that Delta Smelt is now a rare species adds another considerable difficulty. Together, these difficulties are part of the reason why adaptive management actions such as those described in the ongoing Fall Outflow Adaptive Management Plan (Reclamation 2011, 2012) and the now concluded Vernalis Adaptive Management Plan (VAMP, San Joaquin River Group Authority 2013) are planned for a minimum of 10 years, allowing accumulation of data, development of appropriate interpretation of these data, and comparison of observations across as broad a range of conditions as is possible given a 10-year time frame. But even after such a relatively long period of manipulation and observation, questions will likely remain about how some factors interact to affect Delta Smelt abundance.

In summary, we conclude that our new conceptual models can be used successfully to derive testable hypotheses about Delta Smelt responses to changing habitat conditions. Our hypotheses
and the analyses we conducted to test them highlighted some key points as well as critical data gaps and the challenges associated with formulating and testing hypotheses in complex ecological contexts. The key points about Delta Smelt and their habitat generally agree with basic biological principles and earlier conceptual models, but warrant reinforcement because they are crucial to understanding Delta Smelt and to developing and assessing habitat management actions. Other results are less conclusive because of data limitations and the shortcomings of our largely univariate hypotheses and simple comparative analysis approach. Next steps should include addressing critical data gaps, modeling that more fully considers the effects of interacting factors on Delta Smelt, and applications of the information in this report in support of management actions. Examples of such efforts are provided in Chapter 9.

Chapter 9: Recommendations for Future Work and Management Applications

The conceptual model in this report can be viewed as a collection of hypotheses. These hypotheses are not limited to the hypotheses posed in Chapter 7 of this report; essentially, each component and linkage in the conceptual models can give rise to meaningful questions and hypotheses by itself or together with other components and linkages. This is one of the main functions of conceptual models.

Some of the hypotheses that can be derived from our conceptual model have already been addressed in the published research reviewed in Chapter 4 of this report. These results provide the knowledge base used to construct our conceptual model as well as previous conceptual models. They also provide the knowledge base for current Delta Smelt management efforts. The results and conclusions in this report add to this knowledge, but they also emphasize the need for additional monitoring, focused studies, and/or additional analysis and synthesis of existing data. These are the information gaps that can be used to guide future research activities to enhance our understanding of how factors interact to control Delta Smelt abundance.

Filling these information gaps is critically important for improving management strategies for Delta Smelt and for constantly adapting them to expected and unexpected future changes. It is clear that ecological changes due to continued growth of California’s human population, climate change, new species invasions, and other natural and anthropogenic factors will increase the challenges associated with Delta Smelt management. Moreover, as discussed in the previous Chapter, we will likely never be able to correctly detect or predict all effects of management actions and other changes in an ecosystem as complex and constantly changing as the San Francisco estuary. Science and management have to go hand in hand to constantly identify, implement, evaluate, and refine the best management options for this ever-changing system. In this Chapter, we provide examples of next steps in three major areas where additional work is needed: 1) filling critical data and information gaps; 2) mathematical modeling; and 3) applications to support adaptive management actions. We conclude this report with recommendations for future analysis and synthesis efforts.
Critical Data and Information Gaps

A short list of the most critical data and information gaps identified by the updated conceptual model is given below. It is important to note that this is not an exhaustive list of the potentially productive research questions that could be addressed for Delta Smelt. Instead, these are primary research topics that emerge as major data and information gaps in multiple places within the updated conceptual model. This indicates that additional monitoring and research on these topics may be particularly urgently needed and filling these gaps would provide immediately useful results. The list of critical data and information gaps is organized around the environmental drivers and habitat attributes identified in our conceptual models.

Contaminants and Toxicity

There is a general awareness that exposure to contaminants can impair the health of Delta Smelt and other fishes. A few studies have documented adverse effects, but little is known regarding the thresholds at which most contaminants would be toxic to or otherwise adversely affect Delta Smelt (or their prey). Even less is known about how various contaminants may interact when they co-occur, or how their effects may be enhanced or suppressed by these interactions or by other environmental factors.

1. Focused laboratory studies may provide the most efficient way to assess effects of metals, pesticides, pharmaceutical products, or mixtures of contaminants as long as field-relevant concentrations are used. However, translating results of laboratory tests to the field remains a challenging problem (Scholz et al. 2012).

2. Significant work to understand the effect of nutrient loading from municipal sources on the food web has been done (Weston et al. 2014) (e.g., Sacramento Wastewater Treatment Plant, Parker et al. 2012). A logical next step is to conduct manipulative experiments in which effluent is reduced or shut off. This type of work has recently begun (T. Kraus, USGS, personal communication), but may require multiple iterations during a variety of seasons and environmental conditions in order to understand how such manipulations or future treatment upgrades could be used to provide desired food web responses. Monitoring should continue after any such upgrades to determine if they have the expected outcomes.

Entrainment and Transport

Evaluation of differences in entrainment among years could not be critically evaluated from salvage data; better ways to estimate, monitor, and evaluate entrainment losses due to south Delta exports are needed. Such improved estimates could be derived from experimental research on Delta Smelt and other species along with hydrodynamic modeling. Besides the need to improve the estimates of direct proportional population losses due to entrainment, similarly relevant or more important needs include assessing the influence of entrainment on key population attributes (e.g., genetics, demographics, population dynamics and viability effects).
**Predation Risk**

The majority of the hypotheses regarding predation risk could not be fully evaluated due to a lack of data regarding co-occurring predator and prey biomass and predation rates of predators on Delta Smelt.

1. The distribution and diet of major predators with respect to the distribution of Delta Smelt needs further investigation. For some predator species, data may already be available that describe distributions over multiple years and one data synthesis effort has already begun (Mississippi Silversides, USFWS Beach Seine Survey; analysis initiated by B. Schreier, DWR). However, data are lacking for several Striped Bass and Largemouth Bass life stages and focused studies are necessary to understand how these species’ distributions overlap with the distribution of larval, juvenile, sub-adult, and adult Delta Smelt.

2. The distributional overlaps of Delta Smelt with their predators need to be described over varying conditions of turbidity, salinity, temperature, and hydrology. Linking predation risk to key environmental drivers and habitat attributes will shed light on how Delta Smelt may experience varying degrees of predation across seasons and years.

**Food**

Food availability is a critical aspect of Delta Smelt habitat throughout the conceptual model. However, many of the hypotheses about effects of food availability in the conceptual model could not be fully evaluated with available observational data due to incomplete information on prey densities and Delta Smelt feeding behavior throughout Delta Smelt habitat.

1. An extension of the IEP EMP into the Cache Slough complex and possibly other areas around the margins of the estuary would allow a fuller regional comparison of prey densities.

2. Another option is to make concurrent zooplankton sampling a routine part of the four major surveys monitoring Delta Smelt (SKT, 20 mm, TNS, FMWT). To varying degrees, this has been ongoing since 2005, but lack of trained staff has resulted in delayed processing of many samples and concurrent zooplankton samples have never been collected during the SKT survey. Adding appropriate zooplankton sampling and sample processing capacity to the fish monitoring surveys would allow for broader and more timely comparisons of pelagic food availability between monitoring stations with and without Delta Smelt present, similar to the analysis conducted in this report for the larvae collected during the 20mm survey (Larval Hypothesis #2).

3. Studies of Delta Smelt growth (from otoliths) and feeding habits (from stomach contents) concurrent with zooplankton sampling would maximize the utility of the concurrent prey sampling by allowing the refinement of functional response models.

4. Studies of Delta Smelt feeding behavior and prey availability with regard to amphipods and other prey that are not well sampled by any of the existing monitoring surveys could help determine the importance of these types of prey to the Delta Smelt population.
Harmful Algal Blooms

While recent research has resulted in improved understanding of the factors influencing the quantity, toxicity and location of HABs, there are still many uncertainties about their direct and indirect effects on Delta Smelt relative to other factors and about what can be done to prevent them. Furthermore and in spite of their importance to ecosystem and human health, there is still no routine quantitative monitoring program in place that specifically targets harmful algae. The TNS and FMWT surveys now include qualitative, visual assessment of Microcystis, but more quantitative techniques and techniques that detect additional harmful species and their toxicity would likely provide greater insights. Such techniques are increasingly available (e.g., solid phase adsorption tracking; Wood et al. 2011) and some focused studies that quantify and provide distributions of HABs have been conducted or are underway. These studies should be continued in order to address hypotheses related to the effects of HABs in the conceptual model and evaluate the utility of these techniques for routine monitoring applications.

Delta Smelt Responses

To fully evaluate the interactions of various stressors on Delta Smelt population biology, a quantitative life cycle population model is needed. While such models exist, they can be refined based on research into important aspects of Delta Smelt reproductive biology, including the reproductive output of individual Delta Smelt and the population as a whole, and how it varies with environmental conditions.

In particular, fecundity data on adult female Delta Smelt caught in the SKT have only recently been collected. This is a critical parameter, necessary to assess the reproductive potential of the population in any given year. Continued collection of fecundity data over multiple years and hydrological conditions is crucial to understanding the population response to environmental conditions in the seasons preceding reproduction. In addition, an understanding of variables controlling the number of spawning events in a year for wild Delta Smelt is necessary to understand the full reproductive potential of the population. An exploration of whether spawning events are discernible on otoliths is ongoing (Hobbs group, UC Davis); if so, retrospective analyses relating multiple spawning events to concurrent conditions (e.g., tidal phase, food availability, water temperature) may be possible.

Finally, efforts to better characterize spawning habitat and habitat attributes needed for successful egg hatching should also continue. This is needed to more fully evaluate and understand linkages between environmental drivers such as hydrology and larval recruitment. Of all the life stages of Delta Smelt, we know the least about the egg stage; Delta Smelt eggs have never been found in the wild. Because of this, we were not able to construct a life stage transition conceptual model that specifically focused on eggs. More information about spawning and egg hatching habitat is needed to fill this gap in our conceptual models and to identify management actions that would promote beneficial habitat attributes.

Mathematical Modeling

As demonstrated in this report and by others, conceptual models are useful tools for identifying and understanding key ecosystem components and relationships, but they do not quantify them and cannot be used to quantitatively define functional responses to environmental drivers or make
quantitative predictions. Furthermore, as discussed above, the simple univariate and comparative analysis approaches employed throughout this report cannot capture the effects of multiple and often interacting drivers on the Delta Smelt population as a whole and on specific processes such as growth, mortality, and reproduction. The influences of interspecific interactions and abiotic forcing factors on populations and communities in complex ecosystems such as estuaries are also difficult to directly measure in any practical way. Only mathematical models can deal with such complexities and provide quantitative assessments and predictions.

Fortunately, the number of scientific publications about Delta Smelt that include various types of increasingly sophisticated mathematical models is growing rapidly. Recent examples include mathematical models based on statistical approaches (e.g., Bennett 2005, Manly and Chotkowski 2006, Feyrer et al. 2007, Nobriga et al. 2008, Kimmerer 2008, Kimmerer et al. 2009, Feyrer et al. 2010, Thomson et al. 2010, Mac Nally et al. 2010, Miller et al. 2012, Sommer and Mejia 2013, Kimmerer et al. 2013). These efforts generally focused on habitat associations using presence/absence data from the various monitoring surveys or on changes in Delta Smelt abundance based on abundance indices generated by the monitoring surveys and the effects of multiple habitat attributes (covariates) on these changes.

There is also a rapidly developing body of population life cycle models for Delta Smelt and other SFE fish species (e.g., Blumberg et al. 2010, Maunder and Deriso 2011, Massoudieh et al. 2011, Rose et al. 2011, Rose et al. 2013a, b). These models use either a statistically-based “state–space” multistage life cycle modeling approach or a spatially explicit, individual-based simulation modeling approach. Both approaches allow for analysis of the importance of drivers that affect different life stages of Delta Smelt and vary in space and time.

Not surprisingly, results of mathematical modeling efforts to date agree strongly that no single factor can explain the observed Delta Smelt population dynamics and long-term changes in abundance. There is less agreement, however, about which factors are most important (see for example Rose et al. 2013b) and about the exact sequence and nature of their interactions that led to the 2002-3 Delta Smelt POD decline. It is possible, perhaps even likely, that the natural complexity of the estuarine ecosystem coupled with multiple human impacts will prevent definitive answers to these types of questions, especially when they are sought through overly rigid application of formal hypothetico-deductive reasoning and methods (Quinn and Dunham 1983). We agree with Rose et al. (2013b) that the inherent complexity of the system and the challenges it presents for scientists and managers alike “is perhaps the best reason to develop and compare alternative modeling approaches.” Even the most sophisticated modeling oversimplifies complex systems and includes many assumptions. This means that instead of a single modeling approach, multiple alternative conceptual and mathematical modeling approaches, from the simple to the complex, are needed to understand how complex systems work and to predict future changes with sufficient confidence to allow for effective management interventions. The following sections give a brief overview of some of the alternative mathematical modeling efforts currently underway or proposed for the future.

A comprehensive state-space modeling effort that takes advantage of available Delta Smelt abundance data from all monitoring surveys and the even larger monitoring data set about habitat attributes is currently underway (Ken Newman, FWS, personal communication) and future analyses using the individual-based model developed by Rose et al. (2013a) have been proposed (Rose et al. 2013b). As mentioned above, a full description or application of mathematical models is outside of the scope of this report, but to illustrate the utility of additional alternative approaches and further explore some of the linkages and interactions in our conceptual model,
we give three additional examples of alternative mathematical modeling approaches that may be used to further test some of the hypotheses in the conceptual models in this report. The first is a qualitative modeling approach, the second a multivariate statistical modeling approach, and the third a numerical simulation modeling approach. Each of these approaches was explored by one of the co-authors of this report. Importantly, these approaches are meant to complement, not replace state-space, individual-based, and other modeling approaches for Delta Smelt. **Furthermore, results are preliminary and included for illustrative purposes only; peer-reviewed publications of these analyses need to be completed before they can be used to draw any conclusions.**

### Qualitative Models

Qualitative modeling provides a theoretical foundation for understanding system behavior by minimizing the loss of generality and realism at the expense of model precision (Levins 1974, Levins 1975, Puccia and Levins 1991). Qualitative modeling is based on a mathematically rigorous approach that can be used to gain insight on community level process and to examine the consequences of intended or inadvertent human-induced perturbations in managed systems. Questions often addressed through qualitative modeling include the resilience and stability of the system and the direction of population change (Puccia and Levins 1991), the role of system structure on stability (Dambacher et al. 2003, Fox 2006) and the degree of predictability in the response of populations to perturbations (Montaño-Moctezuma et al. 2007, Hosack et al. 2009). Such questions have strong implications in terms of stability-complexity relations (May 1972, Pimm 1984, Haydon 1994) and the persistence of populations and communities following regime shifts (Baxter et al. 2010, Brook and Carpenter 2010, Capitán and Cuesta 2010, Cloern and Jassby 2012).

The increased ecological understanding of the upper SFE and the potential drivers and mechanisms underlying the interannual population responses of Delta Smelt reviewed by the FLaSH and MAST syntheses provide a strong rationale to further refine and integrate our knowledge on community level interactions and ecological drivers in this highly altered system. Towards that goal, we envision qualitative modeling as a complementary approach to other types of models to evaluate the response of Delta Smelt and other populations in the upper SFE over several temporal and spatial scales. Qualitative modeling for Delta Smelt can address some relevant system-level knowledge gaps which are usually less amenable to analyses using other modeling approaches, namely, the influence of species interactions and multiple feedback levels on community stability and population changes in response to perturbations on one or more species. For example, understanding the mechanisms leading to Delta Smelt population responses under different hydrological conditions is an area of significant interest.

Signed-digraphs are a useful representation of the structure of a system, as defined by the community matrix, and have been used in qualitative models exploring food webs (Liu et al. 2010), extinction events in communities (Vandermeer 2013), and other ecological topics of theoretical and conservation relevance. Castillo (unpublished data) used this approach to evaluate the predicted response of Delta Smelt to a sustained change in fall outflow as required in the 2008 FWS Biological Opinion. Recognizing that outflows can control $X_2$ and the size and location of the LSZ (see Chapter 4), and affect other segments of the aquatic community supporting Delta Smelt, Castillo (unpublished data) modeled the response of subadult Delta Smelt to low (5,000 cfs; $X_2 = 85$ km), intermediate (8,000 cfs; $X_2 = 81$ km) and high (11,400 cfs; $X_2 = 74$ km) fall outflow scenarios. Community composition for each outflow scenario was determined relative
to the geographical distribution of species expected to occupy the LSZ. The high outflow model included six community components: phytoplankton, zooplankton, Delta Smelt, predators of Delta Smelt, the overbite clam *Potamocorbula amurensis*, and outflow. The intermediate outflow scenario included two additional community components: the Asian clam *Corbicula fluminea* and the cyanobacteria *Microcystis aeruginosa*. The low outflow scenario included the same variables as in the intermediate flow scenario, except that the overbite clam was excluded and the Brazilian waterweed, *Egeria densa* was added. For each of these communities, community components could exhibit positive or negative feedbacks and positive or negative interactions with other community components. For each of the assumed flow conditions, the four alternative types of community interactions were assumed and each met the stability criteria, as defined by Puccia and Levins (1991). The predicted response of the Delta Smelt population was: 1) predominantly positive under the high outflow community scenario, 2) ambiguous under the intermediate outflow community scenario and 3) very ambiguous under the low outflow community scenario. According to these preliminary results, both outflow and outflow-induced changes in community composition and structure seem to play a critical role in determining the population response of Delta Smelt. These model predictions supported the hypothesis that a shift in the LSZ towards $X_2 = 74$ km is a necessary condition for the fall outflow action to exert a positive influence on the Delta Smelt population. Qualitative models like these can provide useful assessments when the general direction of community interactions are understood but the data are insufficient to support a quantitative model.

**Multivariate Statistical Modeling**

In this report we reviewed results from many multivariate statistical modeling efforts such as the multivariate autoregressive modeling (MAR) conducted by MacNally et al (2010) to discern the main factors responsible for the POD declines and the hierarchical log-linear trend modeling by Thomson et al. (2010) that used Bayesian model selection to identify habitat attributes (covariates) with the strongest associations with abundances of the four POD fish species and determine change points in abundance and trends. The state-space life cycle modeling by Maunder and Deriso (2011) is also based on multivariate statistical modeling; an extension of this work is currently underway by Newman and others (Ken Newman, USFWS, unpublished data). We anticipate that insight from the current conceptual model may be used to facilitate additional multivariate statistical models. As an example, we present preliminary results (Mueller-Solger, USGS, unpublished data) of univariate and multivariate statistical analyses of $X_2$ relationships with annual Delta Smelt abundance indices that follow the approach in Jassby et al. (1995). The purpose is to further explore some of the hypotheses related to hydrology and the size and position of the LSZ included in our conceptual model and to illustrate the importance of considering more than one factor when trying to understand Delta Smelt dynamics. We include this brief exploration in this report because it serves as a useful and relevant example, but as noted above, we advise readers that these are preliminary results from an analysis that has not yet undergone peer review and should be viewed with caution. Moreover, individual and interactive effects of additional factors were not considered in this analysis, but are likely also important (see Chapter 8). As noted in Chapter 7, we recognize that “hydrology” by itself does not affect Delta Smelt, nor does the “$X_2$” index which is used in this analysis as an index of general hydrological (outflow) conditions in the estuary. As shown in our conceptual model (Fig. 38), hydrology affects Delta Smelt through the combined effects of its interactions with other dynamic drivers and stationary landscape attributes (tier 1) on habitat attributes (tier 3). Many of
these interactions have been described in this report; others should be explored further in future studies.

This analysis is intended to evaluate the effects of prior abundance, step changes, and concurrent and prior hydrological conditions in the estuary on the relative abundance of larval to early juvenile Delta Smelt (20 mm index, Fig. 3; hereafter referred to as “larval” Delta Smelt). It also considers prior hydrological conditions and the entire available abundance index time series for larval Delta Smelt provided by the 20 mm survey. The 20 mm survey, one of the newest IEP monitoring surveys, was started in 1995. Delta Smelt distribution data from this survey is heavily used to assess and manage entrainment risk. Similar to prior analyses of TNS and FMWT data (Feyrer et al. 2007, Nobriga et al. 2008), Kimmerer et al. (2009, 2013) and Sommer and Mejia (2013) used a generalized additive modeling (GAM) approach to examine the associations between Delta Smelt occurrence or catch per trawl at 20 mm survey stations and habitat attributes (salinity, temperature, turbidity, and calanoid copepod density) measured concurrently at the same stations. There have, however, been few analyses of annual abundance data from this survey. After 19 years, the 20 mm survey now provides barely enough annual abundance data points (indices) to conduct multiple regression analyses with up to two predictor variables. Clearly more years of data collection and more in-depth analyses are needed and the analyses presented here are merely a starting point.

This analysis uses annual abundance indices for larval Delta Smelt (20 mm survey, 1995-2013), adult Delta Smelt (SKT survey, 2003-2013), and subadult Delta Smelt during the previous year (FMWT survey, 1995-2013) (Fig. 3). It also uses larval recruitment indices calculated from the annual abundance indices (20 mm to SKT ratio and 20 mm to FMWT, Year-1 Ratio, Fig. 46; see previous chapters for caveats regarding index ratios). Data from the SKT survey was only used for univariate analyses because the SKT index time series only has 11 data points at this time. Spring and fall X2 values were obtained by first calculating mean monthly X2 values calculated from daily X2 values provided by the DWR Dayflow database and then averaging the mean monthly X2 values for the “spring” months February to June and the “fall” months September to December. The 2002-2003 step decline in Delta Smelt abundance (Thomson et al. 2010) was introduced as a before/after factor (“Step”). Details about the data sources are provided in Chapter 3 of this report.

The multivariate analyses presented here were conducted with generalized linear modeling (GLM) following the approach of Jassby et al. (1995) and followed with a classical linear modeling (LM) approach guided by the GLM results. For the GLM, model parameters were estimated with a Poisson error distribution, a log link function describing the relationship between the predictor variables(s) and the mean, and a natural spline to represent non-linearities. The degrees of freedom for the splines were restricted to only 2 (i.e. one interior knot) because of the low number of available data points. Models requiring estimation of more than two independent parameters (aside from the intercept) were not considered for the same reason. Applying the GLM approach avoids the need for log-transforming the abundance data and using natural (quadratic) splines as smoothers allows a more natural representation of non-linearities than using polynomials.

The responses predicted by these models have a fairly high degree of precision as indicated by low values of SE/Mean and residuals were consistent with model assumptions. The results show significant univariate relationships at the P < 0.05 level (Table 7) between the 20 mm abundance index and spring X2, prior fall X2, and prior FMWT abundance index. The relationship is strongest with prior fall X2, followed by spring X2 and prior FMWT abundance index (Table
The relationship with spring X2 appears unimodal with maximum 20 mm indices associated with spring X2 values between about 55 and 70 km (Fig. 79a). The relationship with prior fall X2 appears negative (Fig. 79b), and the relationship with the prior FMWT abundance index (Fig. 79c) appears positive. Each of these univariate relationships was improved by the inclusion of one of the other predictor variables (Table 7). Relationships with spring and prior fall X2 were also improved by including the 2002-3 step change. As mentioned above, multivariate analyses with more than two predictor variables were not conducted because of the relatively small amount of available data (n = 19, Table 7). Based on AIC comparisons (Table 7), including the 2002 step change (introduced as a before/after factor, “Step”) somewhat improved the relationship of the 20 mm index with spring X2 (Fig. 73a) and with prior Fall X2 (Fig. 79b), but not with the prior FMWT index because that index was the basis for the analyses that detected the step change and thus already includes the step change in the actual data (Fig. 79c, model not included in Table 7). Including the prior FMWT abundance index improved the relationships with spring and fall X2 more substantially, but the model combining the effects of spring X2 and prior FMWT fit the 20 mm index data nearly as well as the model combining the effects of spring X2 and prior FMWT (Table 7).

It is interesting to note that while prior fall X2 by itself was a stronger predictor of the 20 mm index than spring X2, spring X2 was the stronger predictor when the step change or previous fall abundance were taken into account. Baxter et al. (2010) hypothesized that the shift toward higher prior fall X2 values (Fig. 17) may have contributed to an ecological “regime shift” associated with the step decline in Delta Smelt and other species. This means that prior fall X2 and the “step” factor and FMWT decline in this analysis may be related, which could explain the very similar outcomes for the two models combining spring X2 with either prior fall X2 or the prior FMWT index.

Partial residual plots show the relationship between a predictor variable and the response variable given that other independent variables are also in the model; in other words, they show the effect of one predictor variable given the effect of one or more additional predictor variables. Partial residual plots for the relationships of the 20 mm index with the combinations of spring X2 and prior fall X2 (Fig. 80 a and b) and spring X2 and prior FMWT abundance index (Fig 80 c and d) show that the general shape and direction of the relationships of the 20 mm index with each of the individual predictor variables (Fig. 79) remains intact in the models with combined predictors, but the partial residuals do not closely follow the fitted lines. This indicates that while each variable has its own, distinct effect on the 20 mm index that is maintained in the presence of the other variables, interactive effects among these variables are quite strong. In summary, low values of prior fall X2, high prior FMWT abundance, and intermediate values of spring X2 have positive associations with the abundance of larval/postlarval Delta Smelt, but the effects of individual variables are mediated by the presence of the other variables.

Because the spline degrees of freedom were strongly restricted in this GLM analysis, the results are quite similar to the results of classical linear models (LM) with log-transformed abundance data and a quadratic term to represent the unimodal non-linearity in the relationship between the 20 mm index and spring X2 (Fig. 81). We include these models here because they are more easily reproducible than the GLM models and offer simple equations for making predictions about larval abundance that can be used in adaptive management applications. As for the GLM analysis (Table 7), the best fits overall were achieved by combining spring X2 with either the step change or the prior FMWT abundance index (Table 8). All predictor combinations improved the models compared to the univariate relationships (Table 8). Based on a comparison of regression...
coefficients and P-values, the LM relationships were statistically weaker (Table 8) than in the GLM analysis (Table 7).

Another way of including prior abundance in statistical relationships of abundance with habitat attributes and environmental drivers is to use abundance indices that are proportional to prior abundance indices, in other words, ratios of present to prior abundance indices. In this report, we used the ratios of 20 mm to SKT and 20 mm to FMWT\textsubscript{year-1} abundance indices (Fig. 46; see also caveats about these indices in Chapter 3) as larval recruitment indices from adults and subadults, respectively. We found that recruitment of larvae from adults was linearly related to spring X\textsubscript{2} for the entire available time series (2003-2013, Fig.82a and Table 9). The recruitment index for 2013 was higher than expected based on the other data points. The relationship of the recruitment index from subadults to next year’s larvae with winter-spring X\textsubscript{2} was also linear for the POD period after the abundance step decline in 2002 (Thomson et al. 2010), but with more scatter at higher X\textsubscript{2} values. Interestingly, no relationship was apparent at all before the 2002 step decline when the proportional larval recruitment from then more abundant subadults was generally low (Fig. 82b and Table 9). In the current POD regime, larval recruitment from parental stock appears to be highest when flows through and out of the Delta are high and the interface between fresh and brackish water is located to the west (i.e. low X\textsubscript{2}), although it can occasionally also be high at lower flows, as was the case in 2013.

In late winter and spring 2013, CVP and SWP exports were reduced to comply with OMR flow requirements in the 2008 USFWS Biological Opinion aimed at reducing the risk of adult and

Table 7. Summary of relationships between the 20 mm abundance index for Delta Smelt (response variable) and one or more predictor variables: n, number of observations (years); SE/Mean, model standard error (square root of mean squared residual) as proportion of mean response, P, statistical significance level for the model; R\textsuperscript{2}, coefficient of determination; adjusted R\textsuperscript{2}, adjusted for the number of predictors in the model; AIC, Akaike information criterion; Δ AIC, AIC differences; w (AIC), AIC weights. All relationships modeled with generalized linear models (GLM) with a Poisson error distribution, log link function, and a natural cubic spline with two degrees of freedom as a smoother for all predictor variables except “Step.”

<table>
<thead>
<tr>
<th>Predictor Variable(s)</th>
<th>n</th>
<th>SE/Mean</th>
<th>P</th>
<th>R\textsuperscript{2}</th>
<th>Adjusted R\textsuperscript{2}</th>
<th>AIC</th>
<th>Δ (AIC)</th>
<th>w (AIC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spring X\textsubscript{2}, FMWT\textsubscript{year-1}</td>
<td>19</td>
<td>0.119</td>
<td>&lt;0.001</td>
<td>0.791</td>
<td>0.731</td>
<td>39.5</td>
<td>0.00</td>
<td>0.53</td>
</tr>
<tr>
<td>Spring X\textsubscript{2}, Fall X\textsubscript{2,year-1}</td>
<td>19</td>
<td>0.120</td>
<td>&lt;0.001</td>
<td>0.787</td>
<td>0.726</td>
<td>40.1</td>
<td>0.60</td>
<td>0.39</td>
</tr>
<tr>
<td>Fall X\textsubscript{2,year-1}, FMWT\textsubscript{year-1}</td>
<td>19</td>
<td>0.126</td>
<td>&lt;0.001</td>
<td>0.764</td>
<td>0.697</td>
<td>43.2</td>
<td>3.78</td>
<td>0.08</td>
</tr>
<tr>
<td>Spring X\textsubscript{2}, Step (Factor)</td>
<td>19</td>
<td>0.143</td>
<td>&lt;0.001</td>
<td>0.677</td>
<td>0.612</td>
<td>53.6</td>
<td>14.12</td>
<td>0.00</td>
</tr>
<tr>
<td>Fall X\textsubscript{2,year-1}, Step (Factor)</td>
<td>19</td>
<td>0.135</td>
<td>&lt;0.001</td>
<td>0.712</td>
<td>0.655</td>
<td>55.8</td>
<td>16.35</td>
<td>0.00</td>
</tr>
<tr>
<td>Fall X\textsubscript{2,year-1}</td>
<td>19</td>
<td>0.145</td>
<td>&lt;0.001</td>
<td>0.646</td>
<td>0.601</td>
<td>56.0</td>
<td>16.53</td>
<td>0.00</td>
</tr>
<tr>
<td>Spring X\textsubscript{2}</td>
<td>19</td>
<td>0.176</td>
<td>0.006</td>
<td>0.476</td>
<td>0.411</td>
<td>79.9</td>
<td>40.43</td>
<td>0.00</td>
</tr>
<tr>
<td>FMWT\textsubscript{year-1}</td>
<td>19</td>
<td>0.187</td>
<td>0.015</td>
<td>0.408</td>
<td>0.334</td>
<td>89.4</td>
<td>49.98</td>
<td>0.00</td>
</tr>
</tbody>
</table>
Figure 79. Plots of the Delta Smelt 20 mm survey abundance index as a function of a) spring (February-June) X2, b) previous year fall (September-December) X2, and c) Delta Smelt fall midwater-trawl abundance index in the previous year. Details of general linear models (GLM) used to fit the lines are in Table 7.
Figure 80. Plots of partial residuals for the relationships of the 20 mm index with the combinations of spring X2, prior fall X2, and prior FMWT abundance index summarized in Table 1 (panels a, b, d, and e). The plots shown here also include partial fit lines and their 95% confidence intervals. Values for the time period of analysis are shown for: c, X2; and f, the fall midwater trawl abundance index from the previous year.
larval Delta Smelt entrainment into the water export pumps. This was the first time since the 2008 USFWS Biological Opinion was issued that exports were specifically reduced to lower Delta Smelt entrainment risk. In other years, flows were high enough to allow for higher export levels or export reductions to protect salmon were deemed sufficiently protective for Delta Smelt. It is possible that the intentional reduction in Delta Smelt entrainment risk in 2013 contributed to the high larval recruitment from adults during relatively low flow conditions, but additional years with similar conditions and targeted management actions as well as better estimates of entrainment and more in-depth analyses with other flow variables and flow averaging periods.

**Figure 81.** Plots of the Delta Smelt 20 mm survey abundance index as a function of a) spring (February-June) X2, and b) previous year fall (September-December) X2. Lines are either simple linear least squares regression (lines) or quadratic regression (curves). Details of linear models (LM) used to fit the 1995-2013 lines are in Table 8.
are needed to test this hypothesis and obtain a better understanding of flow effects on larval recruitment.

Overall, these preliminary findings suggest that abundance of the larval to early juvenile life stages of Delta Smelt may respond quite strongly to spring and prior fall outflow conditions. The relationships of the 20 mm index with spring X2 shown in this analysis were much stronger than relationships of the TNS and FMWT indices with spring X2 (Table 1, Fig. 17. Similarly, hydrological conditions in the fall seem to have a greater impact on subsequent abundance of larvae than on subsequent juvenile abundance (TNS index; Mount et al. 2013). This is consistent with the findings by Kimmerer et al. (2009) who noted more pronounced relationships of spring X2 with earlier than with later life stages of Delta Smelt and explained that this was “probably because the earlier life stages occupy areas that are fresher and therefore more responsive to changing flow than the more brackish regions.” While the size and location of the LSZ itself may be important for maturing adults in the fall, its interface with fresh water may be important to larvae and spawning adults. A more westward interface means a larger freshwater habitat for spawning and larval rearing that reaches into the shallow eastern region of Suisun Bay and is well connected with Suisun Marsh sloughs and, in wetter years, the Napa River. It also means a larger distance to the export pumps in the southern Delta and thus a reduced risk of entrainment for spawning adults and larvae. Interactions of flow with other drivers and habitat attributes as shown in the conceptual models in this report are likely also important. This suggests that at least

### Table 8. Summary of relationships between the log-transformed 20 mm abundance index for Delta Smelt (response variable) and one or more predictor variables. All relationships modeled with simple least-squares linear models (LM). For explanation of column headings see Table 6.

<table>
<thead>
<tr>
<th>Predictor Variable(s)</th>
<th>n</th>
<th>SE/Mean</th>
<th>P</th>
<th>R²</th>
<th>Adjusted R²</th>
<th>AIC</th>
<th>Δ (AIC)</th>
<th>w (AIC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spring X2, (Spring X2)², log FMWT year⁻¹</td>
<td>19</td>
<td>0.237</td>
<td>0.000</td>
<td>0.745</td>
<td>0.694</td>
<td>2.1</td>
<td>0.00</td>
<td>0.85</td>
</tr>
<tr>
<td>Spring X2, (Spring X2)², Fall X2 year⁻¹</td>
<td>19</td>
<td>0.274</td>
<td>0.001</td>
<td>0.661</td>
<td>0.593</td>
<td>7.5</td>
<td>5.42</td>
<td>0.06</td>
</tr>
<tr>
<td>Fall X2 year⁻¹, log FMWT year⁻¹</td>
<td>19</td>
<td>0.280</td>
<td>0.000</td>
<td>0.621</td>
<td>0.574</td>
<td>7.7</td>
<td>5.54</td>
<td>0.05</td>
</tr>
<tr>
<td>Spring X2, (Spring X2)², Step (Factor)</td>
<td>19</td>
<td>0.292</td>
<td>0.002</td>
<td>0.616</td>
<td>0.540</td>
<td>9.9</td>
<td>7.78</td>
<td>0.02</td>
</tr>
<tr>
<td>Fall X2 year⁻¹, Step (Factor)</td>
<td>19</td>
<td>0.307</td>
<td>0.002</td>
<td>0.544</td>
<td>0.487</td>
<td>11.2</td>
<td>9.06</td>
<td></td>
</tr>
<tr>
<td>Fall X2 year⁻¹</td>
<td>19</td>
<td>0.318</td>
<td>0.001</td>
<td>0.479</td>
<td>0.449</td>
<td>11.7</td>
<td>9.58</td>
<td>0.01</td>
</tr>
<tr>
<td>Spring X2, (Spring X2)²</td>
<td>19</td>
<td>0.329</td>
<td>0.006</td>
<td>0.473</td>
<td>0.407</td>
<td>13.9</td>
<td>11.83</td>
<td>0.00</td>
</tr>
<tr>
<td>log FMWT year⁻¹</td>
<td>19</td>
<td>0.333</td>
<td>0.002</td>
<td>0.430</td>
<td>0.397</td>
<td>13.4</td>
<td>11.29</td>
<td>0.00</td>
</tr>
</tbody>
</table>
at present, increased Delta outflow and a more westward LSZ in fall, winter, and spring may have important beneficial effects on early life stages of Delta Smelt, but other factors (possibly including summer flows which were not included in this analysis) may be more important for their survival to adults.

Finally, similar to previously published analyses, this analysis strongly suggests that previous life stage abundance should always be taken into account in statistical explorations of habitat effects.

**Figure 82.** Adult (panel a, SKT) and subadult (panel b, FMWT the previous year) to larvae (20 mm Survey) recruitment indices (abundance index ratios) as a function of spring X2 (February-June). For 20 mm/SKT a linear regression was calculated with and without 2013, which appears to be an outlier. For 20 mm/FMWT the previous year separate regressions were calculated for the POD period (2003-2013), the period before the POD (1995-2002), and the entire data record (not shown). See Table 9 for regression results.
Table 9. Summary of relationships of larval recruitment indices (abundance index ratios) for Delta Smelt (response variable) and spring X2 (predictor variable; spring: February-June): n, number of observations (years); SE/Mean, model standard error (square root of mean squared residual) as proportion of mean response, P, statistical significance level for the model; R², coefficient of determination. All relationships modeled with least-squares linear models (LM).

<table>
<thead>
<tr>
<th>Index Ratio</th>
<th>Period</th>
<th>n</th>
<th>SE/Mean</th>
<th>P</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>20-mm/ SKT</td>
<td>2003-2013</td>
<td>11</td>
<td>0.556</td>
<td>0.006</td>
<td>0.588</td>
</tr>
<tr>
<td>20-mm/ SKT</td>
<td>2003-2012</td>
<td>10</td>
<td>0.270</td>
<td>0.000</td>
<td>0.918</td>
</tr>
<tr>
<td>20-mm/ FMWT_Year-1</td>
<td>2003-2013</td>
<td>11</td>
<td>0.469</td>
<td>0.003</td>
<td>0.648</td>
</tr>
<tr>
<td>20-mm/ FMWT_Year-1</td>
<td>1995-2002</td>
<td>8</td>
<td>1.012</td>
<td>0.771</td>
<td>0.015</td>
</tr>
<tr>
<td>20-mm/ FMWT_Year-1</td>
<td>1995-2013</td>
<td>19</td>
<td>0.981</td>
<td>0.321</td>
<td>0.058</td>
</tr>
</tbody>
</table>

on Delta Smelt. Prior abundance can be introduced into these relationships as actual abundance data (e.g. abundance indices or catch per trawl data), periods of relatively constant abundance (here introduced as a “step” factor), or by combining it with present abundance in proportional abundance indices such as the index ratios used here as recruitment indices. Similar to the relationships of juveniles with spring X2 discussed in Chapter 4, the overall depressed abundance of larval Delta Smelt during the POD period that started in 2002 leads to less substantial larval abundance increases with increasing outflows and decreasing X2 values than before the onset of the POD. However, the association of high larval recruitment with high spring outflow suggests that winter and spring hydrology, through its effects on habitat attributes, may be an important driver of larval recruitment during the current POD period, although it may be less important at higher abundance levels.

In summary, this preliminary analysis provides an example of how relatively simple multivariate modeling can yield interesting insights, in this case about how prior conditions (prior fall X2), prior abundance (prior FMWT), step changes in abundance, and concurrent environmental conditions (spring X2) may all have important effects on Delta Smelt abundance in the spring. While further analyses, more sophisticated life cycle modeling, and publication in a peer-reviewed journal are needed to draw firm conclusions, these preliminary results support the idea discussed throughout this report that neither scientific understanding nor management effectiveness can be improved by only considering a single effect, or a single season or life stage. High larval recruitment is essential for setting the stage for a strong year class, but higher growth and survival through subsequent life stages are also needed to achieve and sustain higher population abundance levels.

**Numerical Simulation Modeling**

Quantitative simulations of the multiple factors and processes that affect Delta Smelt life stage transitions in our conceptual model are an obvious next step in the exploration and synthesis
of the information presented in this report. The purpose of simulation modeling is to represent a phenomenon or process in a way that allows users to learn more about it by interacting with the simulation (Alessi and Trollip 2001). In particular, simulations allow users to easily control experimental variables and test hypotheses. Guidance from simulation model “dry runs” can make actual laboratory and field experimentation much more efficient and effective. Simulations are also valuable in visualizing outcomes, thus further promoting learning and understanding.

The individual-based Delta Smelt model by Rose et al. (2013a, b) is an example of a complex simulation model specifically created for Delta Smelt. Another simulation modeling option is to utilize “off-the-shelf” simulation software such as the “STELLA” (Structural Thinking and Experiential Learning Laboratory) simulation construction kit (http://www.iseesystems.com/softwares/Education/StellaSoftware.aspx). STELLA is designed to let users easily create their own simulations using system dynamics including positive and negative causal loops, and flows, accumulations and conversions of materials.

Culberson (USFWS, unpublished data) created a simple quantitative simulation model in STELLA that includes several life stages of Delta Smelt and is based on seasonal environmental conditions and stage to stage estimates of survival. While this simulation modeling approach appears to be feasible, it remains to be seen how such an approach will approximate actual population dynamics encountered in the field and how results compare to those of other simulation models such as the individual-based life cycle model by Rose et al. (2013a,b). A user-friendly STELLA-based model can be useful in the interim, however, to explore the relative contribution of lifecycle stage and environmental covariates to the overall status of Delta Smelt abundance from year to year and to test hypotheses derived from the conceptual model. In its fullest expression, this MAST-associated lifecycle model will be useful for illustrating how multiple suites of plausible co-variates can allow for different Delta Smelt abundance outcomes. For example, it may be possible to find high abundance under degraded conditions given low entrainment losses across successive winters and springs. Conversely, it is possible to encounter low Delta Smelt abundance given otherwise good environmental and outflow conditions with significantly warmer temperatures during fall pre-adult maturation periods. Moreover, simulated changes in survival can provide a useful frame of reference to evaluate alternative outcomes of cohort size or population size attained at different life stages. For example, given the reported levels of larva, juvenile and sub-adult Delta Smelt in IEP surveys, what levels of daily survival between life stages would be required to attain the relative abundances corresponding to each of the four years being compared? Could the small anticipated differences in assumed daily survival among those four years be attributed to some combination of habitat attributes? Or, could stage-to-stage survival (e.g., percent of individuals surviving from one stage to the next) provide a more useful frame of reference to address that question? Our proposed STELLA simulation model and associated modeling exercises will comfortably allow exploration of these questions and related ideas.

This type of modeling will best be used iteratively with emerging data and within synthesis reports to identify where important gaps exist in the Delta Smelt lifecycle understanding and demonstrate how disparate information sources might be brought together to inform our smelt population estimates through time. Importantly, our model can be used in combination with the narrative description of “a year in the life” of the Delta Smelt population from the conceptual model to more effectively describe environmental and management effects on population status in the SFE. We are especially interested in using such a model to avoid single-factor outcome discussions where smelt populations are seen as the result of “one versus another” environmental
or management-related trade off, particularly when single factor analysis is aggregated over decades of data collection efforts in what we know is a constantly-changing estuary.

Figure 83 shows how output from such a model might be useful for keeping track of the variable influence of factors on overall Delta Smelt abundance across seasons within three hypothetical years. Six factors are plotted according to their sensitivity rank (their relative influence on simulated population outcomes). Specific sensitivity levels can then be identified according to the combinations of factors that emerge as important across succeeding seasons and years. Models built to simulate these influences can then be closely examined to discern how different years, year types, or management practices influence simulated abundance, and to detect where potential data gaps or inconsistencies are among the alternative conceptual models or model modes. The basis for using such an approach is a comparative one, and an absolute resolution of the size or behavior of the real Delta Smelt population is not anticipated – but remains the overall objective. Of real interest here is providing a way to interpret our emerging conceptual model within potential regime-shifts, and to capitalize on previous specifications of this model to organize our ever-improving understanding. Of additional benefit is the ability to use these models easily in “learning sessions,” where users interact with the modelers and species experts to deepen understanding of Delta Smelt biology and its relationship to Delta ecology and management.

**Applications to Support Delta Smelt Management**

We have shown that the conceptual models in this report provide a reasonable and up to date conceptual framework that can be used to analyze and synthesize existing data and knowledge about Delta Smelt, identify critical data and information gaps, and guide new field and laboratory studies as well as mathematical modeling efforts. We have also discussed many challenges that limit our ability to reach firm conclusions and make highly confident predictions about the effects of management actions and other changes on Delta Smelt. And we have noted that science and management have to go hand in hand to constantly identify, implement, evaluate, and refine the best management options for Delta Smelt in the highly altered and ever-changing estuarine ecosystem that represents the entire range of this species.

Adaptive management is a well-established approach for systematically integrating science and management. As mentioned earlier in this report, it is increasingly required in plans for management of the San Francisco estuary, but to date, the Vernalis Adaptive Management Program (VAMP) and the Fall Outflow Adaptive Management Plan are among the few clear examples of systematically planned and implemented adaptive management in the estuary.

We end our report with examples of how our conceptual models can be used to adaptively manage and improve Delta Smelt habitat. We conclude with several recommendations for the next analysis, synthesis, and modeling efforts. These efforts are a key ingredient for the more widespread adoption and success of adaptive management strategies; without the conceptual and mathematical models provided by these efforts adaptive management of ecosystems simply cannot proceed.

Table 10 gives examples of adaptive management goals and associated uncertainties to address habitat deficiencies (“habitat problems”) identified and discussed in this report. This table is intended as an illustration of how our conceptual models can be used to inform the first three steps of the nine-step adaptive management framework developed by the DSC Delta Science Program (DSP 2013). These three steps are: 1) definition of the problem; 2) establishment of
management goals and actions to address the problem; and 3) modeling of linkages between management goals and actions. The third step specifically requires conceptual or quantitative models for the purpose of evaluating outcomes of alternative management actions and identification of uncertainties and data gaps. Conceptual models are also important in the other six adaptive management steps, for example to design effective adaptive management experiments and appropriate monitoring and to analyze, synthesize and evaluate results.

Table 10 is organized around the habitat attributes identified in the conceptual models. For each habitat attribute, we describe some example categories of management actions that could be considered to improve the status of Delta Smelt. In essence, these actions represent an example “tool box” for the management of Delta Smelt.

Note that the tool box identified in Table 10 is not meant to be exhaustive. Rather, the list is intended as an example set of adaptive management actions suggested by the conceptual models. As such, the list provides no insight into the cost-effectiveness or feasibility of any of the potential actions. Moreover, we acknowledge that there is substantial uncertainty about the potential benefits of actions in the tool box. As mentioned above, identification of uncertainties about the feasibility and benefits of proposed management actions is an important step in adaptive management that can only be accomplished with the help of conceptual or quantitative models. A key point is that these studies are somewhat different than the critical data and information gaps presented earlier in this Chapter. Specifically, Table 10 emphasizes information gaps that are most relevant to specific management questions, while the earlier list focuses on needs to improve the overall scientific understanding that provides the basis for our conceptual models for Delta Smelt. Clearly, efforts to resolve uncertainties and gaps in understanding are needed in both categories. Overlapping uncertainties may highlight especially urgent data and information needs. For Delta Smelt, this includes uncertainties related to contaminants, predation, and entrainment along with interactions of physical habitat attributes with other factors.
Table 10. Example tool-box for applying the conceptual model to Delta Smelt management.

<table>
<thead>
<tr>
<th>Habitat Attribute</th>
<th>Management Actions</th>
<th>Example Study Efforts</th>
</tr>
</thead>
</table>
| Physical Features      | Increase habitat area & quality                                                     | -Identification of key microhabitats for each life stage and attributes.  
|                        |                                                                                    | -Effects of flow/LSZ position on habitat quality, particularly key biotic habitat elements (access to prey, evasion of predators).  
|                        |                                                                                    | -Approaches to maintain & expand high turbidity habitat (e.g. supply, habitat design, SAV management).  
|                        |                                                                                    | -Approaches to maintain and expand habitat with moderate temperatures (e.g. channel configuration, water depth and velocity).  
|                        |                                                                                    | -Evaluation of whether targeted restoration meets habitat needs (e.g. temperature, substrate, turbidity)  
| Chemical Features      | Reduce toxicity                                                                     | -Identification of chronic effects of contaminants.  
|                        |                                                                                    | -Identification of effects of Harmful Algal Blooms.  
|                        |                                                                                    | -Approaches to reduce toxicity from contaminants and HABs  
| Food                   | Increase pelagic production                                                         | -Role of tidal wetlands as subsidy habitats (not necessarily occupied by smelt)  
|                        | Increase access to alternative foods (e.g. epibenthic).                             | -Ammonia-bivalve interactive effects on diatom, copepod, mysid, amphipod production.  
|                        | Reduce sources of loss                                                              | -Relative importance (contribution to smelt growth) of epibenthic foods (e.g., mysids, amphipods, aquatic insects).  
|                        | Manage towards higher quality foods                                                  | -Effect of bathymetry, vegetation type (and density) on access to epibenthic and pelagic foods.  
|                        | Prevention and control of non-native species                                        | -Role of tidal wetlands and wetland/open-water complexes.  
|                        |                                                                                    | -Approaches to reduce losses to benthic grazing (e.g. invasive clams) and/or to the suppression of bivalve populations  
|                        |                                                                                    | -Value of different food types to Delta Smelt nutrition.  
|                        |                                                                                    | -Effects of habitat conditions (e.g. ammonia, flow) on food quality.  
|                        |                                                                                    | -Identification of nutrient sources and sinks.  
|                        |                                                                                    | -Improved detection methods for invasive species  
|                        |                                                                                    | -Studies to evaluate alternative control methods.  
| Entrainment            | Avoid entrainment region                                                             | -Identification of factors that lead to increased occupancy of South Delta.  
|                        | Adjustments to timing and magnitude of exports                                       | -Improved measurement of entrainment and its environmental correlates  
|                        |                                                                                    | -Effects of exports and entrainment on viability (e.g. abundance, genetics, demographics).  
|                        |                                                                                    | -Approaches to reduce entrainment and enhance emigration success.  
| Predation risk         | Reduction of predator population                                                    | -Studies on delta smelt responses (behavior, distribution, abundance) to variation in predator abundance.  
|                        | Reduction of predation rate                                                         | -Identify habitat features that reduce predation rate (e.g. depth, turbidity, food, lower water temperatures).  

**Recommendations for future analysis and synthesis**

Efforts to resolve the management issues listed in Table 10 or carry out the modeling and fill the critical science gaps discussed earlier in this Chapter will not succeed without an organizational commitment to continued systematic and long-term collection, synthesis and evaluation of data and information about Delta Smelt, its habitat, and important drivers of habitat and abundance changes. The importance of Delta Smelt for ecosystem and water supply management in and far beyond the SFE is widely recognized. The impressive rate at which we are learning about Delta Smelt and the estuarine ecosystem and the large amount of existing information about them is less widely recognized by many managers and even by many scientists. Part of the reason for this is that it is difficult to track the large quantity of new (since 2010) information documented in this report and even more difficult to integrate it with the previously existing information in a meaningful way. But without this integration, identification of priorities for additional scientific investigations is ad hoc and piecemeal at best and the value of new information cannot be fully realized in management applications such as those listed in Table 10.

Moreover, comprehensive adaptive management efforts simply cannot succeed without adequate conceptual and mathematical models and important science and management opportunities will be missed. Such efforts currently include the ongoing fall outflow adaptive management for Delta Smelt and new efforts called for by the new “Collaborative Science and Adaptive Management Program” (CSAMP), the California Delta Stewardship Council’s Delta Plan, and the multi-agency Bay Delta Conservation Plan (BDCP). The fact that even the incomplete draft version of our report released for public review in June 2013 already played a central role in CSAMP work planning, court documents, and elsewhere bears clear testimony to the fact that there is a great and urgent policy and management need for analysis, synthesis and conceptual models such as those provided in this report.

In consequence, we strongly recommend that there be a continued management, analysis, and synthesis effort, whether carried out by the IEP, the Delta Science Program, or some other scientist, group or agency. While it is possible for individual scientists to take on such efforts (e.g., Bennett 2005), the amount, diversity, and rapid growth of pertinent data and information suggests that team efforts may usually be a more feasible and possibly also a more effective option. Collaborative, multidisciplinary analysis and synthesis teams are also at the core of the National Center for Ecological Analysis and Synthesis in Santa Barbara, CA (NCEAS, [http://www.nceas.ucsb.edu/](http://www.nceas.ucsb.edu/)), the newer National Socio-Environmental Synthesis Center in Annapolis, MD (SESYNC, [http://www.sesync.org/](http://www.sesync.org/)) and the Delta Collaborative Analysis and Synthesis (DCAS) approach promoted by the Delta Science Program’s Delta Science Plan (DSP 2013). Important IEP POD and MAST lessons for future synthesis teams are that the role and responsibilities of all team members need to be very clear, that lines of communication need to always be open and available to all, and that there needs to be strong and fully engaged team leadership with a clearly dedicated lead author and/or lead editor for all major team products. In addition, to complete analyses and reports on schedule, it is necessary for team members to prioritize synthesis efforts for sustained periods of time, without being tasked with additional projects that may be urgent for short-term needs.

Another consideration is the type of publication that results from analysis and synthesis efforts. The IEP MAST and POD teams have written comprehensive agency reports, but would have preferred writing peer-reviewed books or monographs (e.g., published by the American Fisheries Society or by U.C. Press) had the time and resources been available to do so. Such books would be considered better scientific products with greater scientific standing and a longer life span.
and would reach a much larger audience. Another approach would be to write a series of shorter articles that could be published in a special issue of a peer-reviewed scientific journal. This too would take more time and effort and would also somewhat restrict the types of topics that could be covered. Journal articles are, however, the main target for national analysis and synthesis centers such as NCEAS and SESYNC because they have the greatest scientific standing and are the most widely accepted and well established method of written science communication.

Regardless of which analysis, synthesis, and communication approach is chosen, none of these efforts can succeed without commitment of adequate funding, staffing, and other resources. The IEP MAST team that developed and wrote this report was formed in 2012 for IEP science synthesis and work planning, but it has remained a pilot-level effort that was never adequately supported. MAST work remained a part-time effort for all co-authors of this report, and for most it was an “on the side” task compared to their “regular” agency duties. There is no doubt that completion of this report could have proceeded much more rapidly with greater allocation of resources. Public and independent peer reviews of a draft version of this report (see http://www.water.ca.gov/iep/pod/mast.cfm) greatly improved the structure and content, but were not an original part of the MAST planning. Preparing and conducting the reviews as well as responding to the 355 specific and many more general review comments took considerable time (see also Appendix A). Other MAST tasks also added to the delays. In addition to this report, the MAST completed a synthesis report for the Fall Low Salinity Habitat (FLaSH) investigation component of the Fall Outflow Adaptive Management Program (Brown et al. 2014) and prepared a solicitation package for research proposals, which it then also reviewed.

We strongly recommend that adequate, long-term support for these types of efforts be among the highest science and adaptive management priorities for the region and the entire State of California. Given its pivotal role in adaptive management and the increasingly large amounts of new scientific data and information that are produced every year, the authors of this report, individually and as a team, cannot think of any science activity that is more urgently in need of greater support than analysis, synthesis, and communication of scientific results.

For additional analysis and synthesis efforts about Delta Smelt, we recommend that the next individual or team to take this on should:

- Build on this report by evaluating the conceptual model with more rigorous analyses that include more years of data, developing lifecycle and numerical models as discussed above, and/or using the conceptual model to develop a comprehensive list of data and information gaps and approaches to addressing these gaps in order to inform management strategies;
- Early in the process, make clear decisions about the analytical/modeling approaches to be used, the scope of the synthesis to be done, and approaches for review and communication of results;
- Evaluate additional data and information needs concerning Delta Smelt;
- Consider approaches to understand the effects of the wide variety of management actions targeting Delta Smelt, including adaptive management of fall outflow, entrainment, habitat restoration, etc (e.g., Table 10);
- Develop key “indicator” variables that can be used to track and predict the status of Delta Smelt and its habitat and serve as “performance metrics” to evaluate the success of management actions. Such variables, and a “report card” to summarize them, were considered for this report, but the MAST decided that developing them was beyond the scope of
An additional recommendation is that an ultimate goal of these efforts should be the integration of conceptual and mathematical models such as those described in the previous section of this Chapter and the routine use of both types of models in adaptive management. Neither the recently published mathematical models nor existing conceptual models for Delta Smelt have been applied to management issues in a consistent manner. This is likely at least partially due to unfamiliarity of managers with the models and the need for specialists (model developers) to apply the mathematical and in some cases even the conceptual models to management issues in the absence of easy to use and understandable model interfaces and specifications. We also recommend a comprehensive biological modeling forum and/or more specific biological modeling teams and “summits” as recommended by the IEP Science Advisory Group (2010, available at http://www.water.ca.gov/iep/docs/IEPModelWorkshopReview.pdf) and, more recently, the Delta Science Plan (DSP 2013). Such groups would not only facilitate communication among modelers, but could also help make the connection from model development to model applications of interest to managers and policy makers. They would complement and could (and likely should) be integrated with the existing, California Water and Environmental Modeling Forum (CWEMF, see http://www.cwemf.org), which tends to focus on modeling physical processes. As with the overall analysis and synthesis teams, these groups could be implemented by the IEP, The Delta Science Program, CWEMF, or others. The chosen organizational umbrella is less important than actual implementation and involvement of appropriate local and outside scientific and management expertise. Some possible topics for these groups include:

1. Reviews and updates to existing conceptual and mathematical models

2. Further development of mathematical models of Delta Smelt population abundance drawn specifically from the conceptual models described in this report; applications and extensions of recently published models to help make management decisions and guide new modeling efforts; additional modeling efforts and future research projects to improve resolution and understanding of the particular factors identified as critical to reproduction, recruitment, survival, and growth.

3. Review and refinement of new models such as the emerging comprehensive state-space population model (Newman, personal communication); development of additional models or modules of models specifically aimed at estimating effects of inadequately monitored or difficult to measure and evaluate habitat attributes such as predation risk and toxicity; development of new “nested” and/or “linked” mathematical modeling approaches that can accommodate multiple drivers and their interactive effects across temporal and spatial scales.

4. Collaboration among physical and biological modelers, experimental and other scientists, managers, and stakeholders to develop and model management scenarios and strategies that move beyond the current focus on relatively crude distinctions among “water year types” toward a more integrative ecosystem and landscape-based management approach.

We end this report with the hope that the conceptual models and information presented will be used for achieving better management outcomes for Delta Smelt and the estuarine ecosystem on which it depends. These precious natural resources are owned by no one, but are held in public
trust by the California and U.S. governments for the benefit of all the people. We are grateful for
the opportunity to serve our State and nation in the collaborative manner afforded by working
under the interagency umbrella of the Interagency Ecological Program for the San Francisco
Estuary.

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Appendix A: How the Delta Smelt MAST Report was Written

The report titled “An updated conceptual model for Delta Smelt: our evolving understanding of an estuarine fish” (hereafter referred to as Delta Smelt MAST report) was written in 2013-2014 by the IEP Management, Analysis, and Synthesis Team (MAST). The Delta Smelt MAST report was developed through a series of report drafts and a public technical review and followed a set of general report guidelines. This report appendix describes the Delta Smelt MAST report guidelines, the report review and revisions, and report milestones.

Delta Smelt MAST Report Guidelines

Report Purpose and Approach

The Delta Smelt MAST report is a technical report intended to synthesize the latest scientific data and information on Delta Smelt, a topic of particularly high relevance to agency managers and decision makers in California. Specifically, it provides an up to date assessment and conceptual model of factors affecting Delta Smelt throughout its primarily annual life cycle and demonstrates how the conceptual model can be used in science and management. The Delta Smelt MAST report updates and redesigns previous conceptual models for Delta Smelt with new data and information since the release of the last synthesis report about the “Pelagic Organism Decline” (POD) by the Interagency Ecological Program (IEP) in 2010. It then uses the conceptual model to generate hypotheses about the factors that may have contributed to the 2011 increase in

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Delta Smelt abundance and evaluate them using a simple comparative approach. The Delta Smelt MAST report ends with key conclusions, a discussion of our hypothesis testing approach, and recommendations for future work and adaptive management applications, with examples.

1. **Report Development.** The 2014 MAST report is a synthesis report developed and written by the IEP Management, Analysis, and Synthesis Team (MAST). The MAST is co-chaired by the IEP Lead Scientist and IEP Program Manager and includes senior scientists from IEP member agencies tasked with data analysis, synthesis, and work planning. The MAST report is the collective product of a dynamic and collaborative interagency team process involving focused team discussions at monthly MAST meetings, intensive conceptual model and report development at additional multi-day off-site meetings, presentations and discussions with other scientists, stakeholders, and the public (e.g., at the annual IEP workshop, meetings of the IEP Stakeholder Group and IEP Project Work Teams), and data analysis and synthesis as well as writing, integration, and revisions of report sections by MAST members with written communication via email and the MAST wiki. MAST report authors were expected to follow the MAST report guidelines described here. They were also expected to consider all internal review comments by other MAST members and members of the IEP Management and Coordinators teams as well as external technical review comments received during a 40-day public review period. Details about the public review process are given in II.

2. **Report Authorship.** The “author of record” for the 2013 MAST report is the entire IEP MAST, and the responsibility for authorship lies with the entire MAST as well. Individual authorship of report sections is not credited; the report is a product of the IEP MAST and not of any individual author or an individual IEP member agency. All current MAST members are MAST report authors and are listed alphabetically in the initial pages of the report (see III. below). Former MAST members will not be listed as authors, but will be noted as contributors. Each report section had a lead author who had primary responsibility for writing and revising the section. One designated MAST member (Larry Brown, USGS) functioned as report lead editor who compiled and integrated all sections and sent full draft report versions to the MAST for review by all MAST members. All MAST members sent their edits and comments back to Larry Brown and the section authors for revisions. The report went through multiple draft versions before its finalization.

3. **Report Organization.** The 2014 MAST report is an IEP technical report and follows the same basic organization as other IEP technical reports, including a title page, list of all authors, acknowledgements, table of contents, executive summary, an introductory section with background information and report objectives, and concise sections detailing the analysis and synthesis approach, models and hypotheses, findings, and conclusions as well as illustrative tables, figures, and full references for all citations. In response to reviewer recommendations received during the public technical review (see II.), the report was restructured and expanded from originally six to nine Chapters.

4. **Supporting Evidence.** The 2014 MAST report follows the conventions of IEP and other technical reports regarding supporting evidence, which includes the following. The rationale for any findings, conclusions, and recommendations should be fully explained in the report. Whenever possible, conceptual models and hypotheses should be evaluated through analysis of the available data. Additional supporting information should be obtained from the peer-reviewed literature or from publicly accessible reports. Related or competing hypotheses and models that have been previously published in the peer-
reviewed literature should be acknowledged and discussed in the report and conclusions should be based on even-handed, dispassionate consideration of all available evidence. Sources for all supporting data and information should be clearly identified and cited. Citation of personally communicated unpublished results (e.g. emails, memos) is permissible, but should be used sparingly.

**Delta Smelt MAST Report Review and Revisions**

1. **What was the purpose of the review?** The purpose of the public technical review of the draft Delta Smelt MAST report was to ensure its scientific credibility, relevance to managers and decision makers, and a transparent and legitimate process that welcomed and considered input and recommendations from other scientists, managers, stakeholders, and the public.

2. **What was expected of draft Delta Smelt MAST report reviewers?** MAST report reviewers were asked to provide written comments on any and all technical aspects of the draft report, but to pay particular attention to review criteria outlined in the MAST report review guidelines.¹

3. **Who reviewed the draft Delta Smelt MAST report?** The draft Delta Smelt MAST report released for public review on July 23, 2014, was reviewed by invited IEP staff and colleagues as well as by invited external peer reviewers and other scientists who submitted comments during the 40-day public review period, as follows.
   
   a. IEP Coordinators (1 Reviewer, IEP management review)
   b. Former MAST Members (2 Reviewers, IEP colleague scientific peer review)
   c. Invited Subject Area Expert (1 Reviewer, IEP colleague review of contaminants sections)
   d. Independent Scientific Peer Reviewers (3 Reviewers, external independent scientific peer review facilitated by the Delta Science Program)
   e. Other Scientists, Stakeholders and the Public (7 Reviewers, external public review)

   In addition, the IEP Coordinators were asked to review the revised, near-final version of the Delta Smelt MAST report and the executive summary and to approve the final version. The IEP Directors were briefed and invited to comment on the direction and progress of the Delta Smelt MAST report on a quarterly basis.

4. **How were external draft Delta Smelt MAST report reviewers identified, invited, and informed?** Independent Scientific Peer Reviewers for the draft Delta Smelt MAST report were identified by the Delta Stewardship Council’s Delta Science Program (DSP) and Delta Lead Scientist. In accordance with the DSP “Procedures for Independent Scientific Peer Review,”² the Delta Lead Scientist determined and invited the independent scientific peer reviewers using the following selection criteria: standing in the scientific community, expertise relevant to the documents being reviewed, and free of conflict of interest.

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² [http://deltacouncil.ca.gov/docs/2012-11-06/delta-science-program-procedures-conducting-independent-scientific-peer-review](http://deltacouncil.ca.gov/docs/2012-11-06/delta-science-program-procedures-conducting-independent-scientific-peer-review)
All other review was invited by email and in a notice posted on the IEP website. A draft of the 2013 MAST report, associated figures, and MAST report review guidelines were posted on July 23, 2013, for public technical review. The draft report release for review did not include an executive summary and conclusions. The public review period closed on August 31, 2013.

5. **How many review comments were received and where can they be accessed?** The MAST received 14 sets of review comments on the July 2013 draft MAST report. They included many general comments as well as 355 comments that referred to specific lines in the report, see table A1. All comments by external reviewers (public review comments and the review comments by the three independent scientific peer reviewers) were posted on the IEP website.

6. **How were the review comments addressed?** All review comments received during the 40-day review period were compiled in an Excel spreadsheet and summarized numerically (Table A1). Review comments and procedures for addressing them were discussed by the MAST at its regular monthly meetings and during a one-day offsite meeting in November 2013. The process for addressing review comments included the following:

   a. The lead author for each report section had the primary responsibility for addressing review comments pertaining to that section and for revising the section.

   b. Secondary revision leads were also assigned and assisted the primary revision lead.

   c. For each review comment in the Excel spreadsheet, it was noted whether the comment: (1) Did not suggest a revision and no revision was made; (2) Suggested a revision and a revision was made; or (3) Suggested a revision, but no revision was made, for example because it was outside of the report scope, explained elsewhere, or the lead author did not agree with the recommended revision.

   d. Revised sections and the annotated excel spreadsheet were sent by email to the entire MAST. MAST members were alerted to all major revisions.

   e. Major revisions were discussed with all MAST members during MAST meetings and via email.

   f. Decisions about major revisions were made by the whole MAST; no comment implied consent.

   g. Decisions about more minor revisions were made by the section revision leads and the report lead editor, often in consultation with some or all other MAST members.

   h. The report lead editor (Larry Brown, USGS) compiled, further revised, and integrated all revised report sections and sent full draft report versions to the MAST for review by all MAST members. The final draft versions of the report and executive summary were also sent to the IEP coordinators for their review and approval.

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3 http://www.water.ca.gov/iep/pod/mast.cfm
4 http://www.water.ca.gov/iep/pod/mast.cfm
7. **What major changes were made to the draft report in response to review comments?** The draft Delta Smelt MAST report underwent several major changes in response to review comments. Changes include the following:

   a. The report purpose and goals were reconsidered, clarified, and somewhat expanded. Specifically, the four-year comparison of factors that may have contributed to the Delta Smelt abundance increase in 2011 was deemphasized in favor of a broader assessment and conceptual model of factors affecting Delta Smelt throughout its primarily annual life cycle and demonstrations of how the conceptual model can be used in science and management.

   b. The report structure was substantially changed to better fit the revised report purpose and goals and to improve the organization of the large amount of information included in the report. Four new Chapters were added to describe the updated conceptual model (Chapter 5), provide a more thorough overview of Delta Smelt life history and population dynamics (Chapter 6), summarize and discuss findings and conclusions (Chapter 8), and provide recommendations and examples of future work and management applications (Chapter 9). An executive summary was also added, along with this appendix.

   c. The content of the report was expanded to accomplish the somewhat expanded report purpose and goals, reflect previously missing information pointed out by reviewers as well as new information from the latest scientific publications, and provide conclusions and recommendations for future work and management applications.

   d. Several reviewers commented that the simple four-year comparative approach that was used to evaluate factors that may have contributed to the Delta Smelt abundance increase in 2011 was too limited and that more years of data and more in-depth analyses and modeling were needed for this evaluation. The MAST agreed, but decided that these types of analyses would require additional

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### Table A1. Numerical summary of review comments for the July 2013 draft MAST report.

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<th>Total Number of References and Attachment Pages</th>
<th>Total Number of Pages</th>
<th>Total Number of Specific Comments (by Line)</th>
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<td></td>
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<tr>
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<td>Public: Water Supply</td>
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time and resources and were outside the scope of this report which emphasized synthesis of existing information over new data analyses. Instead, the MAST decided to discuss some of the benefits and limitations of analysis and synthesis approaches used in the report in Chapter 8 and existing and ongoing analyses and modeling efforts along with additional, analysis, synthesis, modeling, and other science needs and potential management applications in Chapter 9. Three examples of additional mathematical modeling approaches are also included in Chapter 9. These approaches were explored by individual co-authors of this report. Preliminary results of these analyses are given for illustrative purposes only; peer-reviewed publications of these analyses need to be completed before they can be used to draw firm conclusions.

**Delta Smelt MAST Report Milestones**

**Note:** The time line for the development, review, revision and completion of the Delta Smelt MAST report had to be adjusted repeatedly because of numerous new work assignments for individual MAST members, the large number and depth of review comments, the federal government shut-down, personnel changes, etc.

**2012**

March 13-16 Initial MAST off-site meeting (Marconi Center, CA) to discuss MAST products and direction and start MAST work on the 2012 IEP proposal solicitation\(^5\), the “FLaSH” report\(^6\), and the Delta Smelt MAST report (hereafter MAST report)

Sep 13-14 MAST off-site meeting (Yolo Wildlife Area, CA)

Dec 4-5 MAST off-site meeting (Clarksburg, CA)

**2013**

March 29 First draft MAST report completed

April 24 MAST presentation (talk) at annual IEP Workshop (Larry Brown, USGS)

May 20 Second draft MAST report completed

June 6 Third draft MAST report completed

July 23 – Aug 31 Fourth draft MAST report completed and posted on the IEP website for a 40-day review period

August 14 Draft MAST report discussion with IEP Stakeholder Group

Sep 11 Special IEP Stakeholder Group meeting about the draft MAST report

Oct 30 MAST report poster presentation at 2013 State of the Estuary Conference

Nov 14 MAST off-site meeting (UC Davis, CA)

Dec 8 Fifth draft MAST report completed

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\(^5\) [http://www.water.ca.gov/iep/archive/2012/solicitations.cfm](http://www.water.ca.gov/iep/archive/2012/solicitations.cfm)

2014

Feb 3  Sixth draft MAST report completed

Feb 11  MAST presentation (talk) at DSP-SWRCB “Delta Outflows” workshop (Larry Brown, USGS)

Feb 20  MAST presentation (talk) at a meeting of the IEP Resident Fishes Project Work Team (Larry Brown, USGS)

Feb 26  MAST presentation (talk) at annual IEP Workshop (Larry Brown, USGS)

April 16  Seventh draft MAST report completed

April 17  First draft MAST report executive summary completed

April 24  Second draft MAST report executive summary completed and sent to IEP Coordinators for review

May 15  Eight draft MAST report completed and sent to IEP Coordinators for a one-week “red flag” review. This draft includes the executive summary and a description of how the MAST report was written and revised with a list of major report revisions in response to review comments (Appendix A)

June 2  Ninth draft MAST report completed and sent to IEP Coordinators for review and IEP Directors briefings

June 11  IEP Coordinators briefed on MAST report including a review of the major changes.

June 17  Agencies and stakeholders of the CAMT Delta Smelt Scoping Team briefed about the MAST report including major findings and changes since 2013.

July 2  IEP Stakeholder Group meeting to discuss MAST report revisions and completion

July 3  Coordinators approve the final draft MAST report for publication as an IEP Technical Report; when ready the draft final report will be posted on the MAST webpage\(^7\) until the IEP Technical Report publication is completed and report is posted on the IEP Technical Reports webpage\(^8\)

July 14  MAST model presented to IEP Wetlands Conceptual Model Team.

July 29  IEP Directors meeting with presentation and discussion of final MAST report

July 30  MAST model presented to IEP Wetlands Project Work Team.

August 6  MAST briefing to Drought Operations Plan Team

Appendix B: Calculation of Annual Abundance Indices

This Appendix describes the data and methods used by 4 long-term fish monitoring surveys for calculating annual abundance indices for Delta Smelt (*Hypomesus transpacificus*). Descriptions are arranged sequentially beginning with the Spring Kodiak Trawl, which calculates an index of abundance for adult Delta Smelt, followed by the 20 mm Survey, which calculates an index

\(^7\) http://www.water.ca.gov/iep/pod/mast.cfm

\(^8\) http://www.water.ca.gov/iep/products/technicalrpts.cfm
for late-stage larvae and small juveniles; the Summer Townet Survey calculates an index for juveniles and the Fall Midwater Trawl Survey calculates an index for sub-adults. As mentioned in the main document, abundance indices are not population estimates, but they are believed to increase monotonically with increases in true population size.

**Spring Kodiak Trawl**

The Department of Fish and Wildlife (DFW) initiated the Spring Kodiak Trawl Survey (SKT) in 2002. The SKT replaced the Spring Midwater Trawl and provided a more effective means to monitor the distribution and reproductive status of adult Delta Smelt. Survey results provide near real-time information on the proximity of adult Delta Smelt to south Delta export facilities and can provide an indication of likely spawning areas.

The SKT includes 5 monthly Delta-wide surveys, January through May (Figure 84). Only the first 4 surveys contribute to the annual abundance index. No index exists for 2002, when only 3 surveys were conducted. The index is calculated after all data have been verified for accuracy.

Field crews tow the net at the surface between 2 boats once for 10-min at each station per survey; 5-min surface tows are used at stations with historically high catch to limit excessive Delta Smelt take; a second 5-min surface tow is completed if Delta Smelt catch in the first tow did not exceed 50. A flow meter deployed at the start of the tow and retrieved at the end provides information on distance towed through the water. To calculate fish density, survey personnel assume that the SKT net fishes with the mouth fully opened, an area of 13.95 m$^2$ (7.62 m wide by 1.83 m deep). Volume filtered is the product of distance towed and mouth area. Volume filtered varies and by convention researchers expand catch per volume filtered (number per m$^3$) for juvenile and adult fish to catch per 10,000 m$^3$.

Annual abundance index calculations use adult Delta Smelt data from 39 of the 40 stations (Fig. 84). For each of the first 4 monthly surveys, adult catch per 10,000 m$^3$ values from each station are grouped into 3 distinct regions based on geographic location: 1) the confluence and Suisun region (sites 340, 405, 411, 418, 501, 504, 508, 513, 519, 520, 602, 606, 609, 610, 801); 2) the Sacramento River and Cache Slough region (sites 704, 706, 707, 711, 712, 713, 715, 716, 719, 724); and 3) the San Joaquin River and Delta region (804, 809, 812, 815, 902, 906, 909, 910, 912, 914, 915, 919, 920, 921, 922, 923). A monthly mean is calculated for each region and the sum of the regional means is the monthly or survey index. The sum of the 4 survey indices is the annual index.

**20 mm Survey**

DFW initiated the 20 mm Survey in 1995 to monitor the distribution and relative abundance of larval and juvenile Delta Smelt throughout their historical spring range in the upper San Francisco Estuary (Fig. 85), and provide near real-time information on the relative densities and proximities of these young fish to south Delta export pumps. The 20 mm Survey includes sampling on alternate weeks from mid-March through early July, typically resulting in 9 surveys per year. During each survey, field crews complete 3 oblique tows at each of the 47 stations (Fig. 85). The 20 mm Survey added stations over time, but not all contribute to annual abundance index calculation. The survey added 5 Napa River stations in 1996 for a total of 41 core stations, which are included in the annual abundance index calculations (Fig. 85, circles). In 2008, 6 non-
core stations were added, which are not included in the annual abundance index calculations, including Barker Slough (site 720), Lindsey Slough (site 718), Miner Slough (sites 724 and 726), and the Sacramento Deep Water Shipping Channel (n = 2; sites 719 and 723) (Fig. 85, triangles).

The 20 mm net includes a flow meter located within the mouth of the net to measure distance traveled by the net during the tow. This value is then multiplied by the fixed mouth area of the net (1.51 m²) to provide total volume filtered. The tows are then standardized to catch of Delta Smelt per 10,000 m³.

As already noted, the annual abundance index calculation uses only catch per 10,000 m³ values from the 41 index stations. For each survey, the mean fork length of Delta Smelt is calculated from measurements of the fish captured during each survey. The two surveys just before the average fork length reached 20 mm and the 2 surveys just after the average fork length reached 20 mm are included in the annual abundance index calculation. For these 4 surveys the geometric
mean of the catch of Delta Smelt per 10,000 m$^3$ is calculated across the 41 core stations. The geometric mean for each survey is calculated as the arithmetic mean of log$_{10}(x+1)$-transformed values of Delta Smelt catch per 10,000 m$^3$ across the 41 core stations. The resulting value is then back-transformed (including subtraction of 1) for the calculation of the annual abundance index. The annual abundance index is calculated as the sum of the geometric means of the 4 selected surveys.

**Summer Townet Survey**

The Summer Townet Survey (TNS) was started by DFW in 1959 to produce an annual index of summer abundance for age-0 Striped Bass (*Morone saxatilis*). In the mid-1990s, DFW staff developed an abundance index calculation for Delta Smelt. Annual abundance indices for Delta Smelt have been calculated for the period 1959 through the present, except for 1966-1968. The

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**Figure 85.** Map of 20 mm survey stations showing all currently sampled stations. Data from all core stations are used in abundance index calculation.
TNS Survey samples 32 historic stations, 31 of which contribute to index calculation (labeled as “core stations,” Fig. 86). Currently sampled TNS stations range from eastern San Pablo Bay to Rio Vista on the Sacramento River and to Stockton on the San Joaquin River (Fig. 86). In 2011, TNS added 8 supplemental stations in the Cache Slough and the Sacramento River Deepwater Ship Channel region to increase spatial coverage and better describe Delta Smelt range and habitat (Fig. 86). Historically, TNS sampling began when age-0 Striped Bass achieved a mean fork length of 20 mm based on larval sampling, typically in mid-June to early July, and ended when age-0 Striped Bass surpassed a mean size of 38.1 mm fork length. Since 2003, TNS has consistently included 6 surveys annually, running on alternate weeks from early June through mid- to late August.

Field crews perform at least two 10-min oblique tows at most stations. A third tow is conducted when any fish were caught during either of the first 2 tows. At least 1 tow is completed at each of the new Cache Slough and Sacramento River Deepwater Ship Channel stations. To reduce Delta Smelt take, field crews only perform a second tow at these stations if Delta Smelt catch from the first tow is less than 10. Delta Smelt catch per tow data are used for index calculation.

The annual abundance index for Delta Smelt is the arithmetic mean of the abundance indices from the first 2 surveys conducted each year. Delta Smelt abundance indices for each biweekly survey are calculated by summing catch across all tows for each index station, multiplying the summed catch by a station weighting factor representing the water volume of that station (Table B1); then the volume-weighted catches are summed across all 31 index stations and the sum divided by 1000.

The annual abundance index for age-0 Striped Bass is calculated using similar methods, except the first two surveys are not used. Instead, abundance indices from the 2 surveys that bound the date when the fish reach a mean fork length of 38.1 mm are used; this frequently occurs after several surveys have been completed in a field season.

**Fall Midwater Trawl Survey**

DFW began the Fall Midwater Trawl Survey (FMWT) in 1967 to provide an annual index of relative abundance and information on the distribution of age-0 Striped Bass for the fall period. Later, DFW staff developed abundance and distribution information for other upper-estuary pelagic fishes, including Delta Smelt. Surveys have been conducted in all years from 1967 to present, except 1974 and 1979. The FMWT survey currently samples 122 stations monthly (Fig. 87), from September through December. Station locations range from San Pablo Bay to Hood on the Sacramento River, and from Sherman Lake to Stockton on the San Joaquin River (Fig. 87). Currently, annual abundance index calculations use catch data from 100 of the 122 stations sampled monthly, but the number of stations used for the index has varied through time. Table 12 contains the complete list of stations used for abundance index calculation for FWMT (n = 117), including historical stations (underlined) that must be included for proper calculation of past indices, but are not included in calculations for recent years. The remaining 22 stations were added in 1990, 1991, 2009, and 2010 to improve our understanding of Delta Smelt habitat use (Fig. 87). At each sampling station, field crews perform a single, 12-min oblique tow monthly.

Delta Smelt catch per tow data are used for calculation of the annual abundance index. Individual survey indices are calculated by first grouping the 100 core stations (Fig. 87) into 14 regions based on their location (Table 12). Survey indices are calculated by averaging Delta Smelt catch
across index stations within each region, multiplying these regional means by their respective weighting factors (i.e. a scalar based on water volume; Table 12), and summing the weighted values. Annual abundance indices are calculated as the sum of the 4 survey abundance indices (i.e. September through December).
Table B1. Station weighting factors for stations used in calculations of the summer townet survey annual abundance indexes. Regions are geographic areas designated by the California Department of Fish and Wildlife. See fig. 86 for station locations.

<table>
<thead>
<tr>
<th>Region</th>
<th>Station</th>
<th>Station weighting factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>MONTEZUMA SLOUGH</td>
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<td>20</td>
</tr>
<tr>
<td></td>
<td>609</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>610</td>
<td>4</td>
</tr>
<tr>
<td>SAN PABLO BAY</td>
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<td>213</td>
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<tr>
<td>SUISUN BAY</td>
<td>405</td>
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<tr>
<td></td>
<td>411</td>
<td>46</td>
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<td>SACRAMENTO RIVER</td>
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<td>SAN JOAQUIN RIVER</td>
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<tr>
<td>EAST DELTA</td>
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</table>
Figure 87. Map of fall midwater trawl survey stations showing all currently sampled stations. Data from core stations are used in abundance index calculation.

Fall midwater trawl sampling sites

- = Core stations
- = Non-core stations (began in 1990)
- = Non-core stations (began in 1991)
- = Non-core stations (began in 2009)
- = Non-core stations (began in 2010)
Table B2. Area-regions, weighting factor for each area-region, and stations included within each area-region. Bolded station numbers indicate the current 100 core stations used in calculation of annual abundance indexes. Underlined station numbers indicate stations previously included in calculations but subsequently dropped.

<table>
<thead>
<tr>
<th>Area-region</th>
<th>Weighting factor</th>
<th>Stations included</th>
<th>8-San Pablo Bay</th>
<th>10-Napa River</th>
<th>11-Carquinez Strait</th>
<th>12-Suisun Bay</th>
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<tbody>
<tr>
<td>1-San Pablo Bay</td>
<td>8.1</td>
<td>336, 337, 338, 339</td>
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<tr>
<td>4-San Pablo Bay</td>
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<td>5-San Pablo Bay</td>
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<td>8-San Pablo Bay</td>
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<tr>
<td>10-Napa River</td>
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<tr>
<td>11-Carquinez Strait</td>
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<td>12-Suisun Bay</td>
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Interagency Ecological Program: Management, Analysis, and Synthesis Team

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