Zooplankton Monitoring 2019

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Introduction 3 Zooplankton are a vital trophic link between aquatic primary producers and 4 higher-level consumers of the San Francisco Estuary (SFE). As primary 5 consumers of phytoplankton, zooplankton facilitate the flow of carbon into a large 6 and complex food web, historically supporting abundant fisheries in the region 7 8 (Schroeter et al. 2015; Kimmerer et al. 2018). Many fishes, including Striped 9 Bass (Morone saxatilis) and Chinook Salmon (Oncorhynchus tshawytscha) feed on zooplankton while rearing in the estuary as larvae and juveniles (Goertler et 10 al. 2018; Heubach et al. f1963), while others like Tule Perch (Hysterocarpus 11 traski) and Prickly Sculpin (Cottus asper) feed on zooplankton throughout their 12 lifetimes (Kimmerer 2006Feyrer et al. 2003)). Zooplankton in the SFE are also a 13 key food source for several endangered and threatened species, notably the 14 15 Delta Smelt (Hypomesus transpacificus) and Longfin Smelt (Spirinchus thaleichthys) (Hobbs et al. 2006; Slater and Baxter 2014). 16 This importance of zooplankton prompted the implementation of the Zooplankton 17 Study in 1972 to assess fish food resources in the upper SFE. Mandated by the 18 State Water Resources Control Board's Water Right Decision D-1641, the study 19 is conducted jointly by the California Department of Fish and Wildlife and the 20 California Department of Water Resources under the guidance and management 21 of the Interagency Ecological Program. For nearly 5 decades, this study has 22 monitored the zooplankton community in the region, tracking abundance trends 23 and distributional patterns, detecting and monitoring introduced species, and 24 documenting the dramatic shifts in the community's composition. Changes in 25 zooplankton abundance and composition have since been linked to major 26 declines of the pelagic fishes in the upper estuary (Sommer et al. 2007; Winder 27

and Jassby 2011). This report presents zooplankton annual and seasonal abundance indices and distribution trends from 1974 through 2019 for the most common copepods, cladocera, rotifers, and mysids of the upper estuary.

31 Methods

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Zooplankton sampling has been conducted since 1974 at least once a month 32 March-November through 1995, monthly from 1995 on. at 20 fixed stations in the 33 upper SFE (Figure 1). Three gear types are used for each sampling event: a 34 pump with a 43-micron mesh net for micro-zooplankton (rotifers, nauplii, and 35 small cyclopoid copepods); a Clarke-Bumpus (CB) net with a 160-micron mesh 36 for sampling meso-zooplankton (cladocera and most juvenile and adult calanoid 37 copepods); and a mysid net with a 505-micron mesh for sampling mysid shrimp 38 and other macro-zooplankton. Both the mysid and CB nets are attached to a sled 39 and towed obliquely from the bottom through the surface for a 10-minute tow. 40 Volume is measured using a General Oceanics flowmeter placed in the mouth of 41 each net so that: $V = (end\ meter - start\ meter) * k * a$; where V is the volume 42 of water sampled, k is a flowmeter correction value, and a is the area of the 43 44 mouth of the net. The Teel Marine 12V utility pump is also used at each station to sample approximately 19.8 gallons from the entire water column, which is filtered 45 through a 43-micron mesh net to concentrate the pump sample. Samples are 46 preserved in 10% formalin with Rose Bengal dye before being processed in the 47 laboratory for identification and enumeration of organisms using a dissecting 48 microscope. More information about the sampling and processing methods can 49 be found in the metadata at ftp://ftp.wildlife.ca.gov/IEP_Zooplankton/. 50 51 Abundance indices are calculated for each organism based on the gear type most effective at its capture and reported as the mean catch-per-unit-effort 52 (CPUE). CPUE is calculated as the number of each organism collected per cubic 53 meter of water sampled, so that: $CPUE = s * V^{-1}$; where s is the estimated count 54

- of the target organism in the sample. Copepod abundance indices reported here
- only include adults, as juveniles were not always identified to species. Annual
- and seasonal abundance indices were calculated using 14 fixed stations
- sampled consistently since 1974 (Figure 1) and 2 non-fixed stations sampled
- 59 where bottom specific conductance was roughly 2 and 6 millisiemens per
- 60 centimeter (approximately 1 and 3 psu).
- To analyze long-term trends (1974 to present), annual abundance indices were
- calculated as the mean CPUE for samples collected from March through
- November, as winter sampling was inconsistent before 1995. Seasonal
- abundance indices were calculated as the mean CPUE for samples collected
- during each season: winter (previous December to February), spring (March to
- 66 May), summer (June to August), and fall (September to November). Long-term
- seasonal trends for winter are only shown for 1995 to present. Spatial distribution
- indices for organisms is described as seasonal mean CPUE for by region.
- 69 Estuary regions were defined as San Pablo Bay (stations D41 and D41A),
- Suisun Bay (stations D6, 28, 54, and 48), Suisun Marsh (stations 32 and S42),
- West Delta (stations 60, 64, and 74), Central Delta (stations D16, 86, and D28),
- and the East Delta (92 and M10).

73 Results and Discussion

- Since the implementation of the Zooplankton Study in 1974, a significant
- decrease in the overall abundance of zooplankton has been detected in the
- estuary (Figure 2). Only the abundance of cyclopoid copepods increased in the
- region during this period, driven by the invasion and spread of *Limnoithona*
- 78 *tetraspina*. The overall decrease in zooplankton abundance in the estuary can be
- attributed to a series of invasions into the estuary, most notably that of the Asian
- clam Potamocorbula amurensis in the mid-1980s (Kimmerer, Gartside, and Orsi
- 1994; Carlton et al. 1990). The spread of *P. amurensis* throughout SFE has had

- significant impacts on planktonic abundance in the upper estuary due to its high
- filtration feeding rates on phytoplankton and copepod nauplii. Not only has
- 84 abundance decreased for most of the zooplankton groups, but dramatic shifts in
- the composition of these communities have been detected during the study
- period. These changes have been driven by the introduction and spread of non-
- native zooplankton species throughout the estuary, compounded with changes to
- 88 the abiotic and biotic environments.
- 89 Calanoid copepods
- 90 While overall calanoid copepod abundance has declined slightly over the study
- period, community composition has shifted dramatically (Figure 2A). When the
- study began in the early 1970s the copepods *Eurytemora affinis* and *Acartia* spp.
- dominated the calanoid community. The non-native *E. affinis* was once the
- 94 primary prey item of the endangered Delta Smelt, but its abundance has declined
- to a fraction of what it once was, forcing fish to prey switch to more recently
- 96 introduced calanoids like *Pseudodiaptomus forbesi* (Moyle et al. 1992; Slater and
- 97 Baxter 2014).
- One of the first recorded introduced calanoid copepods was Sinocalanus doerrii,
- a freshwater species native to China that invaded the estuary in 1978 and
- became the most dominant calanoid species in the estuary for a decade (Orsi et
- al. 1983). Then in 1987, after the invasion of *P. amurensis*, the calanoid
- 102 Pseudodiaptomus forbesi was introduced and first detected in the region,
- competing with the abundant E. affinis (Orsi and Walter 1991). P. forbesi quickly
- became the numerically dominant calanoid in the upper estuary and remains the
- most abundant to this day. Another introduction occurred in 1993, when the
- predatory calanoid copepod *Acartiella sinensis* quickly became the second most
- abundant calanoid in the upper estuary, dominating the low-salinity zone in
- Suisun and the West Delta (Orsi and Ohtsuka 1999). This invasion was

- hypothesized to have narrowed the range of *P. forbesi* towards the freshwater zone of the estuary, as predation on *P. forbesi* nauplii by *A. sinensis* has been
- recorded (Kayfetz and Kimmerer 2017).
- In general, calanoid copepod abundance is highest in the estuary during the
- summer and fall months, with lowest abundance during the winter (Figure 3A).
- 114 While calanoid copepod abundance peaked in the summer of 2017 at a nearly 20
- year high, 2018 and 2019 abundance returned to levels comparable to the
- previous two decades. In 2019 the distribution of calanoids throughout the
- estuary was similar to 2018, with *P. forbesi* the most abundant in summer and fall
- in most of the upper estuary (Figure 4A). The predatory *Acartiella sinensis* was
- seen in highest densities in the summer and fall mostly in the Suisun Bay and
- West Delta regions, similar to 2018. In fall 2019, A. sinensis was the most
- abundant calanoid in Suisun, where it co-occurred with high densities of one of
- its primary prey items *Limnoithona tetraspina* (Figure 4B), while *P. forbesi* was
- found to the east of Suisun Bay (Hennessy 2018). Acartia spp. was the only
- native calanoid copepod commonly found in 2019, but it was restricted to Suisun
- 125 Bay and San Pablo Bay in the lower reaches of the estuary. *Eurytemora affinis*
- was the most dominant calanoid throughout the upper estuary in the spring of
- 2019, similar to seasonal distribution patterns of the last two decades.
- 128 Cyclopoid copepods
- 129 While overall zooplankton abundance has declined over the study period, the
- abundance of cyclopoid copepods exploded (Figure 2B). The native *Oithona* spp.
- and Acanthocyclops copepods were at low abundances when the study began,
- but with the introduction of *Limnoithona sinensis* in the early 1980s, and the later
- identification of the invasive *Limnoithona tetraspina* in 1993, cyclopoid indices
- have increased dramatically(Ferrari and Orsi 1984; Orsi and Ohtsuka 1999)
- Abundance indices for the two species of *Limnoithona* were reported together

- from 1980 through 2006 as *Limnoithona* spp., then separately since 2007 when
- they were identified and enumerated as *L. sinensis* and *L. tetraspina*. In 2019, *L.*
- 138 tetraspina abundance was the highest observed for all copepods (Figures 2A and
- 139 **2B)**.
- Much smaller than calanoid copepods collected in the CB net, the *Limnoithona*
- cyclopoids are best retained in pump samples, which use a smaller mesh. Since
- the early 1990s, *Limnoithona* spp. abundance has been higher than calanoid
- 143 copepod abundance, and the small *L. tetraspina* has become the most common
- 144 copepod in the upper estuary. This increase in *L. tetraspina* abundance is likely
- due to a decline of Northern Anchovy in the upper SFE and subsequent
- decreased predation (Kimmerer 2006), as well as the cyclopoid's small size, high
- growth rate, and motionless behavior, making it very difficult for visual feeders to
- capture (Bouley and Kimmerer 2006; Greene et al. 2011). These characteristics
- may increase its ability to escape predation in a region where visual predation is
- most dominant among fish (Kimmerer 2006). The introduction of *L. tetraspina* is
- also linked to the reduction of the range of *P. forbesi* out of the low-salinity zone
- of the estuary, as high *L. tetraspina* densities may have fed and sustained larger
- populations of the predatory *A. sinensis*, which also fed on *P. forbesi* nauplii
- 154 (Kayfetz and Kimmerer 2017).
- Seasonally, *Limnoithona tetraspina* peaks in summer and fall (Figure 3B), with
- lower abundance in winter and spring. As in prior years, this cyclopoid was most
- abundant in the low-salinity zone of the estuary in Suisun Bay and the West
- Delta (Figure 4B). Oithona davisae, a native cyclopoid, was the most abundant
- cyclopoid in the higher-salinity San Pablo Bay in summer and fall (Figure 4B).
- 160 Cladocera
- The cladoceran community of the upper estuary is composed of Bosmina,
- 162 Daphnia, Ceriodaphnia, and Diaphanosoma species, whose populations have

also substantially declined since the onset of the study (Figure 2C). These 163 164 cladocera tend to be herbivorous, feeding primarily on phytoplankton, and were likely hard hit by the invasion of P. amurensis (Baxter et al. 2008; Kratina and 165 Winder 2015). Cladocerans make up a significant portion in the diets of Delta 166 167 Smelt, juvenile Chinook Salmon, and young-of-the-year Striped Bass throughout the upper estuary (Heubach et al. 1963; Slater and Baxter 2014; Goertler et al. 168 2018). The invasion and increase of available copepod prey such as *P. forbesi* 169 and the decline in cladocera has created a shift in the nutritional content of the 170 171 plankton community available for fish, with yet to be determined consequences 172 (Kratina and Winder 2015). While Cladocera abundance has declined overall, recent years summer 173 abundance has been increased and in 2018, summer cladocera abundance was 174 175 the highest observed since the *P. amurensis* invasion (Figure 3C). However, in 2019, abundances dropped back down to the lowest seen in almost a decade, 176 with summer abundance slightly higher than other seasons. In the high outflow 177

year of 2017 some cladocera, namely *Bosmina*, were found downstream in
Suisun and the West Delta, while in 2019 the highest densities of cladocera were

found in the East Delta, with trace concentrations found in other regions of the

estuary (Figure 4C).

182 Rotifer

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While they are the most abundant zooplankton in the estuary, long-term sampling of rotifers using the pump system shows a dramatic decrease in their annual abundance in the estuary since the beginning of this study (Figure 2D). Several species of rotifers make up the community: most abundant being the *Polyarthra*, *Synchaeta*, and *Keratella* genera. Interestingly, the decline of rotifer abundance beginning in the late 1970s preceded the invasion of *P. amurensis* in the estuary (Cloern and Jassby 2012).

The distribution and abundance of rotifers were similar between 2018 and 2019. 190 191 with abundances similar to those found since the invasion of *P. amurensis*. Rotifers were the most abundant zooplankton sampled during 2019 (Figure 2) 192 and were found throughout the estuary (Figure 4D). Overall rotifer abundance 193 peaked in the summer and spring (Figure 3D). Keratella and Polyarthra tend to 194 be most abundant in the freshwater and low-salinity zone of the estuary, while 195 Synchaeta species occur most in the higher-salinity areas of San Pablo Bay and 196 Suisun (Figures 3D and 4D)(Winder and Jassby 2011). A spatial and temporal 197 split was discernable between Synchaeta and other rotifers, with Synchaeta 198 199 having highest densities in the low-salinity zone Bay during the spring, and other 200 rotifers being most abundant in the East Delta in summer.

201 Mysida

202 Not only have mysid abundances declined significantly since the 1970s, but the community has also shifted from being composed almost entirely by the native 203 204 *Neomysis mercedis*, to being dominated by the non-native *Hyperacanthomysis* longirostris (formerly Acanthomysis bowmani) (Figure 2E). The first significant 205 decline in *N. mercedis* occurred during the 1976-1977 drought, likely caused by 206 food limitation from an absence of diatoms due to very low river discharges 207 (Siegfried et al. 1979; Cloern et al. 1983). The populations of N. mercedis were 208 209 able to rebound after the years of drought and stayed at high densities in the Suisun Bay region of the upper estuary until the introduction of *P. amurensis* in 210 the mid-1980s, after which their numbers crashed. 211

In 1993 the introduced *H. longirostris* was first detected by this study, shortly
after the decline of *N. mercedis*, and it quickly became the most common mysid
in the system. However, overall mysid abundances have not returned to their
pre-clam invasion levels (Modlin and Orsi 1997, Figure 2E). Mysids have always
peaked in the spring and summer months, fluctuating with the higher productivity

- in the estuary during those seasons (Figure 3E). Historically mysids have been of
- critical importance in the diets of many fish species in the SFE including Delta
- Smelt, Longfin Smelt, Striped Bass, and Chinook Salmon (Moyle et al. 1992;
- Feyrer et al. 2003; CDFG 2009; Goertler et al. 2018). However, the decline of
- mysids in the upper estuary has resulted in a significant decrease in their
- presence in the diets of fishes of the region (Feyrer et al. 2003).
- This general decline in abundance continued in 2019, even though 2016 and
- 224 2017 had a modest peak in mysid abundances (Figure 2E), and the distribution
- 225 and timing of peaks stayed similar over the last two decades (Hennessy 2018).
- 226 Hyperacanthomysis longirostris was again the most common mysid in the
- estuary during all seasons, while the once common and native *Neomysis*
- 228 mercedis continued to be almost imperceptible in the region. This has been the
- overall trend in the estuary's mysid communities since 1994. As in prior years,
- mysids in 2019 were most abundant during the summer and fall (Figure 3E), and
- 231 highest concentrations occurred in the low-salinity zone of Suisun Bay and Marsh
- 232 (Figure 4E).
- 233 2010 2019 A Decade in Review
- 234 2019 marks the end of the fifth decade of operations for the Zooplankton Study.
- The previous 10 years have been a period of extremes in the San Francisco
- 236 Estuary, with the historic drought of 2012 to 2016, immediately followed by
- record levels of precipitation in the winter and spring of 2017 (Figure 5). We
- briefly analyzed some of the trends in abundance and distribution of key species
- 239 during these periods of extreme climate events to better understand their impacts
- on zooplankton in the estuary. Year hydrologic classification was based on the
- California Department of Water Resources indices for the San Joaquin Valley at
- 242 (https://cdec.water.ca.gov/reportapp/javareports?name=WSIHIST) (Figure 5).

243 Abundance and distribution analysis were limited to stations in the lower estuary,

- the southern Delta, and the San Joaquin River.
- 245 We selected five of the most abundant taxa in the estuary to focus analysis on:
- 246 Pseudodiaptomus forbesi, Limnoithona tetraspina, Diaphanosoma spp.,
- 247 Synchaeta spp., and Hypercanthomysis longirostris. Abundance and distribution
- 248 analysis were limited to the dry years (2012-2016) and the wet years (2011,
- 249 2017, and 2019) and the summer months when abundance is highest for most
- taxa. Due to nonparametric distributions of abundance estimates a Kruskal-
- Wallis test was performed on the monthly mean CPUE for the summer months to
- compare the CPUE and the centers of distribution between drought and flood
- years (Figure 5). The center of distribution for each month was plotted for each
- year and taxa so that:

$$D_{t,i} = \frac{\sum CPUE_{t,s} * km_s}{\sum CPUE_{t,s}}$$

- where $D_{t,i}$ is the center of distribution for taxa t for month i and km_s is the
- 257 distance of station s in km from the Golden Gate Bridge.
- 258 When examining the variation in abundances between dry and wet years both *P.*
- 259 forbesi and Diaphanosoma spp. saw significant increases in their abundances
- during wet years, while *L. tetraspina* saw a significant decrease (Figure 6).
- 261 Interestingly only *Diaphanosoma* spp. and *H. longirostris* had significant shifts of
- their distribution seaward in wet years compared to dry years. These trends
- witnessed over the past decade of the study suggest that outflow can have an
- influence on the abundance and distribution of different zooplankton taxa in the
- estuary. This supports prior research showing the correlation between summer
- outflows and zooplankton abundances or distribution amongst mysid species
- before the invasion of *P. amurensis* (Siegfried et al. 1979; Cloern et al. 1983).
- 268 Analyzing how outflow affects zooplankton abundance and location in the estuary

will require more research but will be important to understanding the spatial and
 temporal relationships between zooplankton and their fish predators.

Conclusion 271 272 In 2019 the Zooplankton Study recorded the abundances of calanoids, cladocera, rotifers, and mysids at lower densities comparable to other recent 273 years and consistent with the downward historic trends in the estuary. Calanoid 274 and cyclopoid copepod abundance peaked in fall, whereas cladocera, rotifers, 275 and mysids peaked in summer. The small, abundant *Limnoithona tetraspina* was 276 again the most abundant copepod in the upper estuary. This multi-decade 277 zooplankton study has enabled researchers and managers to track the shifts in 278 zooplankton abundances and community composition across the estuary for 279 nearly 5 decades. The Zooplankton Study has documented the introduction and 280 281 dominance of Pseudodiaptomus forbesi, Limnoithona tetraspina, and Hypercanthomysis longirostris, as well as the community's response to the 282 invasive clam *Potamocorbula amurensis*. Understanding these dynamics and 283 how they have fundamentally changed trophic interactions is critical to assessing 284 food resources for fish and conservation strategies in the San Francisco Estuary. 285 References 286

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370 Figures

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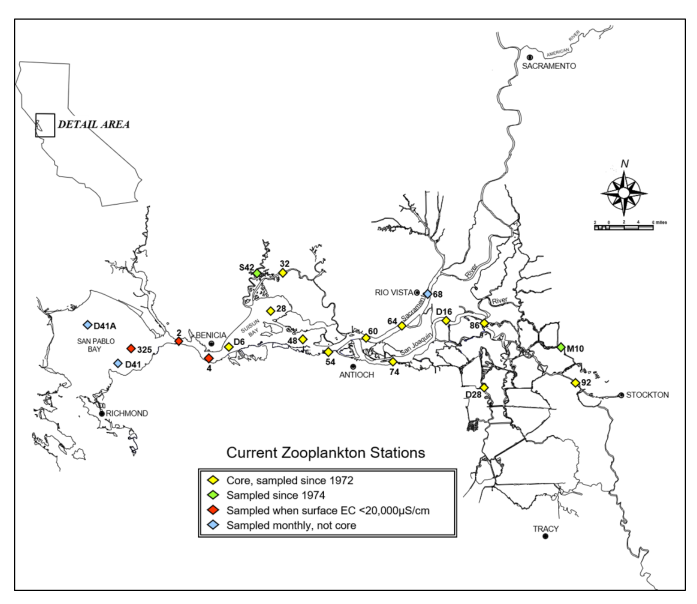


Figure 1. Map of fixed Zooplankton Study stations in the San Francisco Estuary.

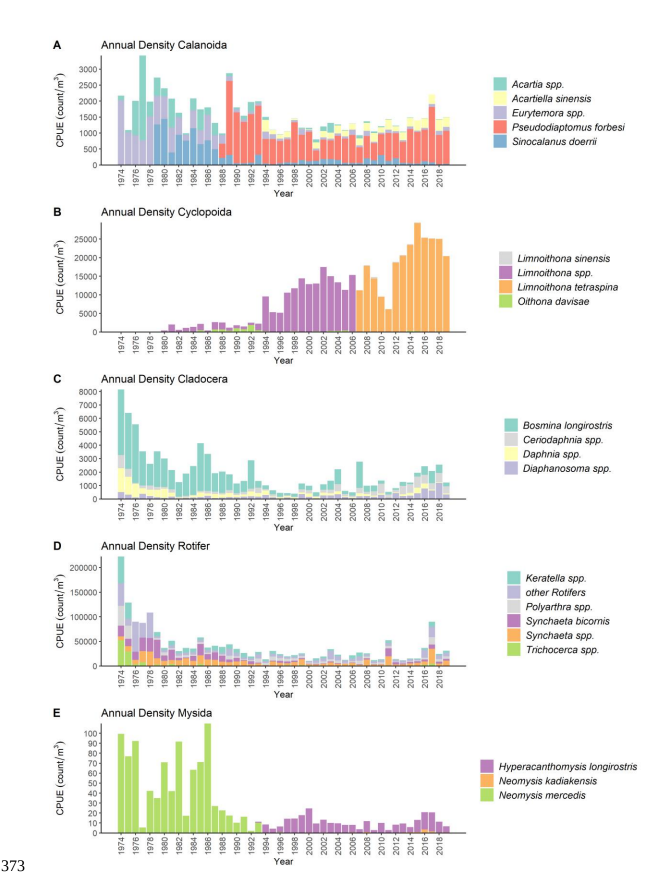


Figure 2. Annual (Mar-Nov) mean zooplankton CPUE for A) Calanoid CPUE in the CB net, B) Cyclopoida CPUE in pump samples, C) Cladocera CPUE in the CB net, D) Rotifer CPUE in pump samples, and E) Mysid CPUE in the mysid net.

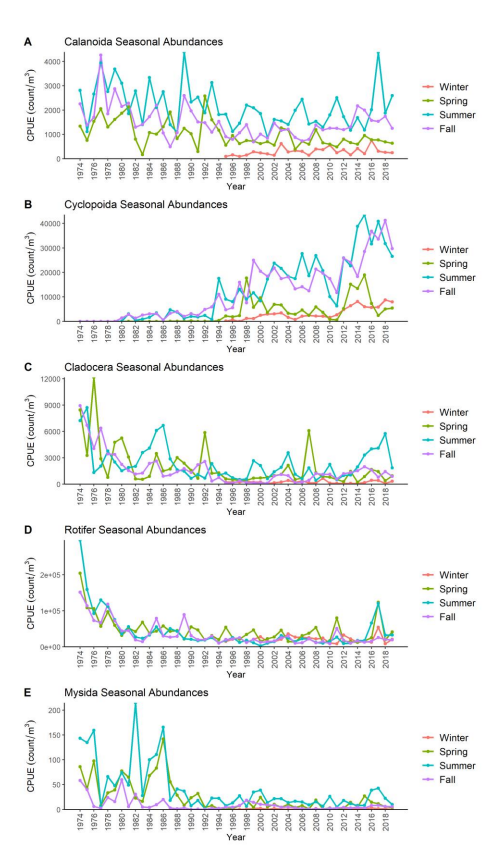


Figure 3. Seasonal mean zooplankton CPUE. Spring, summer, and fall are reported for 1974-2018, winter is reported for 1995-2019. A) Calanoid CPUE in the CB net. B) Cyclopoida CPUE in pump samples. C) Cladocera CPUE in the CB net. D) Rotifer CPUE in pump samples.

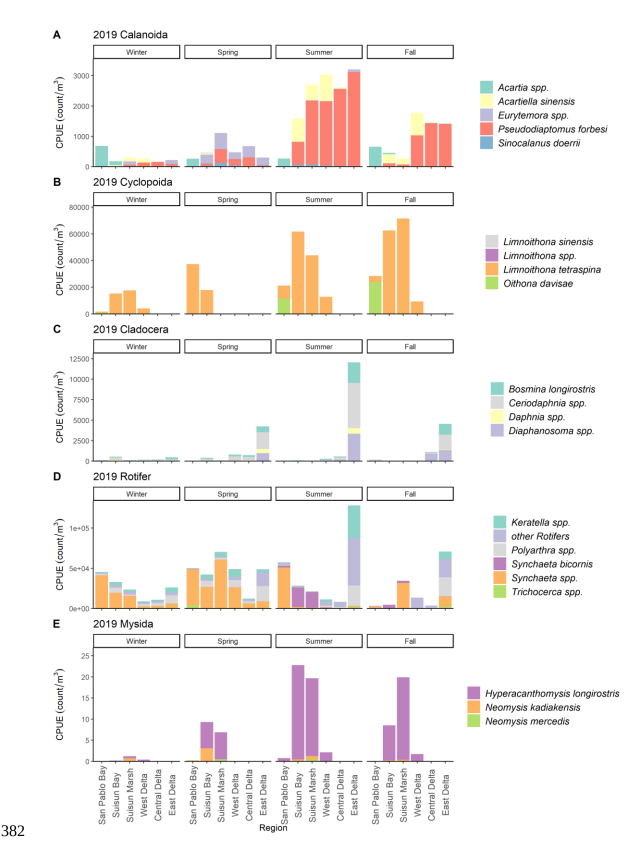


Figure 4. Seasonal mean zooplankton CPUE for 2019 by region for A) Calanoid CPUE in the CB net, B) Cyclopoida CPUE in pump samples, C) Cladocera CPUE in the CB net, D) Rotifer CPUE in pump samples, and E) Mysid CPUE in the mysid net.

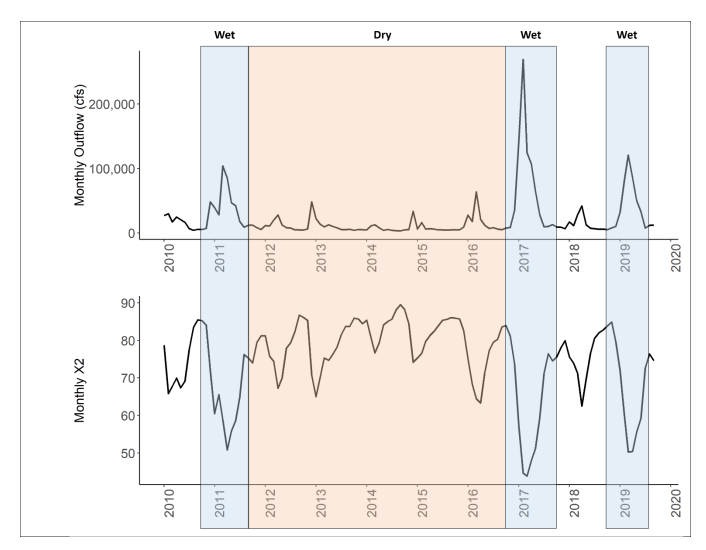


Figure 5. Average monthly outflow (cfs) and average monthly X2 position for 2010 – 2019. Dayflow data from Department of Water Resources https://water.ca.gov/Programs/Environmental-Services/Compliance-Monitoring-And-Assessment/Dayflow-Data.

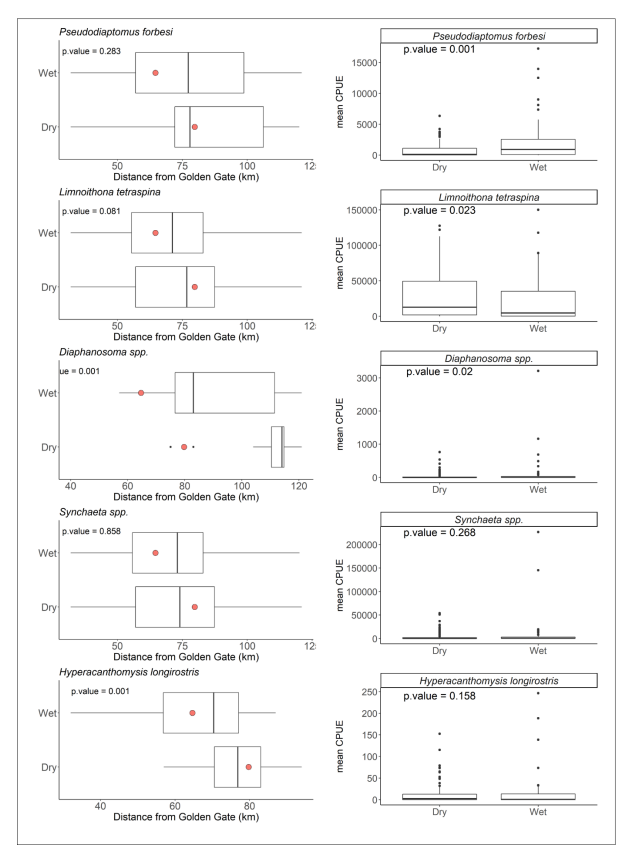


Figure 6. Average summer month CPUE and center of distribution for select taxa in drought (2012-2016) and flood years (2011, 2017, and 2019). Red points in distribution charts represent mean X2.