Densovirus associated with sea-star wasting disease and mass mortality

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Populations of at least 20 asteroid species on the Northeast Pacific Coast have recently experienced an extensive outbreak of sea-star (asteroid) wasting disease (SSWD). The disease leads to behavioral changes, lesions, loss of turgor, limb autotomy, and death characterized by rapid degradation (“melting”). Here, we present evidence from experimental challenge studies and field observations that link the mass mortalities to a densovirus (Parvoviridae). Virus-sized material (i.e., <0.2 μm) from symptomatic tissues that was inoculated into asymptomatic asteroids consistently resulted in SSWD signs whereas animals receiving heat-killed (i.e., control) virus-sized inoculum remained asymptomatic. Viral metagenomic investigations revealed the sea star–associated densovirus (SSaDV) as the most likely candidate virus associated with tissues from symptomatic asteroids. Quantification of SSaDV during transmission trials indicated that progression of SSWD paralleled increased SSaDV load. In field surveys, SSaDV loads were more abundant in symptomatic than in asymptomatic asteroids. SSaDV could be detected in plankton, sediments and in nonasteroid echinoderms, providing a possible mechanism for viral spread. SSaDV was detected in museum specimens of asteroids from 1942, suggesting that it has been present on the North American Pacific Coast for at least 72 y. SSaDV is therefore the most promising candidate disease agent responsible for asteroid mass mortality.

Significance

The cause of SSWD remains a mystery. Scientific hypotheses given for other asteroid mortality events include storms (6–11), temperature anomalies (1, 3, 12), starvation (13), and infection by unidentified pathogens (5). For instance, pathogens in the bacterial genus Vibrio (12, 14, 15) and an unidentified eukaryotic parasite (4) were seen in die-offs of the tropical asteroid Astropyga muricata (4) and the Mediterranean asteroid Actinaster jacksoni (4). However, it is difficult to distinguish the cause of an infectious disease from the associated microbial community that can flourish in a sick or injured animal.

Some early patterns from the field supported the hypothesis that SSWD is contagious. Within a region, SSWD has sometimes moved from site to site similar to an infectious disease. For example, the disease spread north to south in Southern California. All of the major aquaria on the North American Pacific Coast

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Since June 2013, millions of sea stars (asteroids) of the west coast of North America have wasted away into slime and ossicle piles, due to a disease known as sea-star wasting disease (SSWD). SSWD has been used to collectively describe die-offs of sea stars in the Northeast Pacific since at least 1979; however, this SSWD event differs from other asteroid mass mortalities (1–5) due to its broad geographic extent (from Baja California, Mexico to Southern Alaska; pacificrockyintertidal.org) and many (n = 20) species affected, representing several major lineages of Asteroidea (Fig. 1, Table S1, and SI Text). The extensive geographic range and number of species infected might make SSWD the largest known marine wildlife epizootic to date. Outward signs of SSWD vary slightly among species but generally start with behavioral changes, including lethargy and limb curling, followed by lesions, ray autotomy, turgor loss (deflation), and end with animal death (Fig. 1). Histology of dead and dying asteroids from geographically widespread natural habitats and aquaria, showed epidermal necrosis and ulceration, and dermal inflammation and edema in the body wall. Clinically affected (i.e., symptomatic) individuals rarely recover in the laboratory and only occasionally in the field.


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were affected by SSWD in Fall 2013, with mass mortality of captive asterozoids. However, SSWD did not spread into aquaria that sterilized inflowing seawater with UV light but did spread into nontreated aquaria. Furthermore, the incursion of SSWD in aquaria with sand-filtered intake suggests a microscopic infectious stage, rather than a disease that spread only via contact with an infected host or vector. Overall, these patterns suggest a microscopic, water-borne, infectious disease agent, rather than an infected host or vector. Overall, these patterns suggest that the disease is transmissible from symptomatic to asymptomatic individuals and that the pathogenic agent is virus-sized (i.e., <200 nm in diameter).

**Investigation of Viral Agents in SSWD-Affected Tissues**

We sought to identify candidate viruses in inoculum by conducting metagenomic surveys to compare viral consortia inhabiting asymptomatic and SSWD-affected tissues. Tissue samples (~1-cm cross-section of a ray including body wall, epidermis, pyloric caeca, and gonad) were excised from 28 specimens (Tables S2–S5) with sterilized dissecting scissors and placed into sterile 50-mL plastic tubes. Viral metagenomes (i.e., metaviromes) were then prepared and analyzed following Gudenkauf et al. (16) (SI Text). All sequences have been deposited in GenBank under BioProject accession no. PRJNA253121.

Metaviromic analysis indicated a greater prevalence of parvovirus-like sequences in symptomatic relative to asymptomatic asteroid homogenates (in 2 of 13 asymptomatic libraries and 7 of 15 symptomatic libraries). Global assembly of all viral metagenomes (n = 28 libraries, representing 5 \times 10^7 sequences and 1 \times 10^{10} nt of information) resulted in assembly of a near-complete densovirus (*Parvoviridae*) genome, which we have named SSaDV (Table S5).

**Inoculation with Virus-Sized Homogenates Induces Wasting-Disease Signs**

We turned to viruses as potential infectious agents in SSWD because histologic evaluation of hundreds of slides indicated no evidence of prokaryotic or eukaryotic microbial infection within tissues of asteroids affected by SSWD. Asymptomatic adult *Pycnopodia helianthoides* were collected from several sites in Puget Sound, Washington, in January–April 2014 before disease occurrence at the collection sites. Animals were placed in separate 10-gallon glass aquaria with sand-filtered and UV-treated flow-through seawater on separate draw lines (i.e., seawater was not recirculated between aquaria; US Geological Survey Marrowstone Laboratory) and held at ambient temperature (8–9 °C). Viral challenges were performed with homogenates prepared from symptomatic tissues. Tissues from symptomatic asterozoids were homogenized in 10 mL of seawater from the aquarium housing the symptomatic animal. The tissue homogenate was initially centrifuged at 1,000 \times g for 5 min at 4 °C to remove large debris before being split into two 5-mL aliquots. One aliquot was boiled for 7 min to serve as a control before both heat-treated (control) and non-heat-treated (virus exposed) homogenates were serially syringe-filtered through 1-μm pore size polycarbonate and 0.22-μm pore size polyethersulfone filters, resulting in a virus-sized inoculum. Each asterozoid was injected with 0.5 mL of either control (i.e., heat-treated) or viral inocula (0.5 mL) into the coelom cavity. After injection, animals were observed every morning and evening for SSWD signs.

The animals inoculated with virus-sized material showed disease signs after 10–17 d whereas control (heat-treated inoculum) individuals did not develop SSWD signs (Fig. 2). From one of the virus-exposed animals showing SSWD signs in the first challenge, we again isolated virus-sized material to challenge a new set of asymptomatic animals. This second set of inoculated individuals had disease signs after 8–17 d (and again no control animals that received heat-treated inoculum developed SSWD signs). These experiments demonstrate that the disease is transmissible from symptomatic to asymptomatic individuals and that the pathogenic agent is virus-sized (i.e., <200 nm in diameter).

**Fig. 1.** Photographs of SSWD-affected stars (A) asymptomatic *P. helianthoides*, (B) symptomatic *P. helianthoides*, and (C) symptomatic *P. ochraceus*. Disease symptoms are consistent with loss of turgor, loss of rays, formation of lesions, and animal decomposition. (D) Map showing occurrence of SSWD based on first reported observation. (E) Transmission electron micrograph of negatively stained (uranyl acetate) viruses extracted from an affected wild *E. troshelii* from Vancouver. The sample contained 20–25-nm diameter nonenveloped icosahedral viral particles on a background of cellular debris (primarily ribosomal subunits) and degraded viral particles of similar morphology. (Scale bar: 100 nm.)

**Fig. 2.** (A) Proportion of stars remaining asymptomatic after inoculation with control (heat-killed) or virus-sized fraction (VSF) of asteroid homogenates in first (Expt 1) and second (Expt 2) challenge. Survival analysis (20) indicates that the time to lesion development differs among treatment and control groups (log-rank test: \(\chi^2 = 18.6, df = 1, P = <0.0001\)), but there is no difference in time to development of lesions in the two experiments (log-rank test: \(\chi^2 = 0.2, df = 1, P = 0.698\)). (B) Change in SSaDV load between initiation of viral challenge and termination of experiment (i.e., animal expiry in live challenge or euthanasia of control animals). Note difference in scale. Viral load was determined by quantitative PCR (qPCR) targeting the NS1 gene of the SSaDV genome. Different letters above bars indicate that the mean viral abundance change is significantly different (\(P < 0.05\), two-tailed Wilcoxon signed-rank test to account for heteroskedasticity; Fisher’s F-test \(P > 0.05\)). Error bars = SE.
SSaDV Is Linked to Wasting Disease in Field Surveys

Due to the association of SSaDV with asteroid tissues, we examined the incidence of SSaDV among symptomatic (n = 286 individuals) and asymptomatic (n = 49 individuals) asteroids of 14 species. Viral load (number of SSaDV copies detected per mg of tissue) and prevalence (i.e., percentage of samples where SSaDV was detected) were higher in symptomatic than in the asymptomatic animals in all three species where both symptomatic and asymptomatic animals were obtained (Fig. 5). However, the virus was present in both asymptomatic and symptomatic individuals in species where animals in both health states were sampled, including P. ochraceus, P. helianthoides, and E. troschelii. Because SSaDV detection varied by tissue type and location on each animal (Fig. S2), the single tissue sample taken from each individual likely led to some false negatives (in repeated sampling of body-wall tissues from symptomatic P. ochraceus, SSaDV was detected in 11–38% of samples). Due to the potential for these false negatives, it was not surprising that we observed SSaDV in some asymptomatic asteroids. Conversely, SSaDV in asymptomatic animals almost certainly represents viral presence before disease signs develop because we know from our inoculation experiment that signs can take 2 wk to progress after inoculation (or could represent viruses present on animal surfaces that had not yet gained entry to animal tissues).

Despite these procedural challenges, asteroids were more likely to be diseased if they had a high viral load (Fig. 6). In our statistical models involving viral load, we started with all factors and their first-order interaction terms. To focus our interpretation of model effects and increase power, we sequentially removed interaction terms that were not statistically significant (in order of their associated P value) and then did the same for main effects (this removal generally followed Akeike information criterion model selection). For the relationship between SSaDV load and disease, we used a logistic model of symptomatic vs. asymptomatic to evaluate the potential independent effects of SSaDV load, asteroid species, geography (San Diegan province or south of Point Conception vs. Oregonian province or north of Point Conception), and asteroid size (measured as arm circumference) in a sample of 107 symptomatic and asymptomatic P. ochraceus, P. helianthoides, and E. troschelii for which we had size measurements. The main significant predictive variable for being symptomatic was the SSaDV load [logistic regression, square root-transformed load of SSaDV, estimate = 0.00013 (0.00008 SE) chance of being symptomatic increasing with viral load, P = 0.006]. In addition, for a given viral load, asteroids from southern sites were more likely to be symptomatic [logistic regression, estimate = 0.95 (0.53 SE), P = 0.03] (Fig. 6). This result was consistent with analyses limited to P. ochraceus, which was the only species sampled in the North and South. However, neither asteroid size nor species had significant independent or interactive effects with the other factors. Given that viral load was the main predictor of disease, we then asked...
P. helianthoides found a significant interaction between species and size (likelihood ratio test: $\chi^2 = 19.7$, $df = 2; P < 0.0001$) and disease status (likelihood ratio test: $\chi^2 = 7.4$, $df = 1; P = 0.0065$) with additive effects of SSaDV abundance and disease in wild asteroids. Logistic regression (generalized linear model with binomial distribution and logit link) comparing models with species and disease status and their interaction indicated that a model including species (likelihood ratio test: $\chi^2 = 25.0$, $df = 1; P = 0.029$) had the greatest support. This result indicates that viral prevalence differs among species and disease status, but the difference among disease status does not differ significantly among species (likelihood ratio test: $\chi^2 = 5.4$, $df = 1; P = 0.020$). The odds ratios suggest that symptomatic stars are 3.2 times more likely to be virus-positive than asymptomatic stars. Error bars = SE.

Fig. 5. (A) Mean viral load and (B) prevalence (i.e., proportion of SSaDV-positive individuals) (B) as determined by qPCR targeting the VP4 gene of the SSaDV genome. qPCR was applied to P. helianthoides ($n = 10$ asymptomatic and 79 symptomatic), P. ochraceus ($n = 26$ asymptomatic and 72 symptomatic), and E. troschelii ($n = 6$ asymptomatic and 31 symptomatic) whole-tissue DNA extracts. (C) SSaDV load in sympatric asteroid species. All stars except Dermasterias imbricata, Orthasterias sp., and AstreopLeo polycanthus were symptomatic. The number of individuals tested is indicated for each species. Different letters within bars represent significant difference in the percentage of viral reads between asymptomatic and symptomatic asteroids ($P < 0.001$; $df = 113$ for P. helianthoides, $df = 42$ for E. troschelii, and $df = 117$ for P. ochraceus; data log-transformed and corrected for heteroskedasticity by $\sqrt{n}/\sqrt{c}$; t test). The probability of being infected with the virus was higher in symptomatic asteroids. Logistic regression (generalized linear model with binomial distribution and logit link) comparing models with species and disease status and their interaction indicated that a model including species (likelihood ratio test: $\chi^2 = 19.7$, $df = 2; P < 0.0001$) and disease status (likelihood ratio test: $\chi^2 = 7.4$, $df = 1; P = 0.0065$) had the greatest support. This result indicates that viral prevalence differs among species and disease status, but the difference among disease status does not differ significantly among species (likelihood ratio test: $\chi^2 = 5.4$, $df = 1; P = 0.020$). The odds ratios suggest that symptomatic stars are 3.2 times more likely to be virus-positive than asymptomatic stars. Error bars = SE.

Fig. 6. Square root-transformed viral load in asymptomatic and symptomatic asteroids in San Diegan (i.e., south of Point Conception) and Oregonian (i.e., north of Point Conception) biogeographical provinces. For the relationship between SSaDV abundance and disease, we used a logistic model of symptomatic vs. asymptomatic to evaluate the potential independent effects of SSaDV abundance, sea star species, geography (San Diegan vs. Oregonian province), and sea-star size (measured as arm circumference) in the 107 P. ochraceus, P. helianthoides, and E. troschelii for which we had both size measurements and a mix of asymptomatic and symptomatic stars. The main significant predictive variable for being symptomatic was the abundance of SSaDV (logistic regression, square root-transformed count of SSaDV, estimate = 0.0013 (0.0008 SE) chance of being symptomatic increasing with viral count, $P = 0.006$). Error bars = SE.

Fig. 7. Transcription of the SSaDV VP4 as assessed by qRT-PCR comparing asymptomatic and symptomatic tissues. qRT-PCR was performed on whole-tissue RNA extracts from E. troschelii ($n = 5$ asymptomatic and 5 symptomatic), P. ochraceus ($n = 5$ asymptomatic and 6 symptomatic), and P. helianthoides ($n = 10$ asymptomatic and 10 symptomatic), and normalized to quantities of SSaDV assessed by qPCR in cDNA extracts. Transcript levels were significantly higher in asymptomatic P. ochraceus than symptomatic individuals (Mann-Whitney U test; $P = 0.039$, $df = 11$). Error bars = SE.
SSaDV Present in Plankton, Sediments, and Nonasteroid Echinoderms

Given evidence for viral transmission between asteroids in the laboratory, we then sought to understand how SSaDV might move between wild host individuals and populations. To investigate whether uninfected asteroids could contact viruses free in the water, associated with suspended particles, or in sediments, we tested different environmental samples for SSaDV. SSaDV had its highest abundance in the virioplankton size fraction of the water and was present in the suspended particulate material size fraction > 0.2 μm (Fig. 8). SSaDV presence in particulate material is congruent with other observations of paroviruses (16) and might represent viruses adsorbed to abiotic material, viruses in detrital particles from decayed animals, or viruses within larval asteroids. SSaDV was also found in sediments collected from public aquaria that had experienced SSWD several months earlier, and SSaDV was concentrated in sand filters used to treat incoming water and between aquaria (Fig. 9). Therefore, SSaDV might transmit between asteroids and among populations by mechanisms other than direct contact between diseased and healthy individuals, consistent with the observation that SSaDV-infected asteroids shed virus into the water column (Fig. S3). Water-column SSaDV transport helps explain how SSWD spreads among disjunct asteroid populations.

The SSWD epizootic has hit an alarming number of asteroid species in all shallow water habitats. Eight of 11 asteroid species sampled from SSWD areas contained SSaDV (Fig. 4). The broad range of species in which SSaDV was detected is unexpected because most viruses infect a narrow range of host species. However, paroviruses are known to infect across families, and variations in capsid protein secondary structure can result in variable host range (18). Paroviruses gain entry to host cells via transferrin receptors, which are among the most highly expressed proteins in coelomic fluid (where, notably, it is found in coelomocytes, which are a major defense mechanism against cellular microbial infection) (19). It is also possible that receptors are shared between closely related sea-star species. This phenomenon is especially a possibility here because six of the eight species known to be affected by SSWD in which SSaDV was detected are within a single asteroid family (Asteriidae; the remaining two were within the Asterinidae) that may share common cell-surface features through which viruses may infect.

This broad host range inspired us to look for SSaDV in other Northeastern Pacific echinoderms. We observed SSaDV in sympatric, nonasteroid echinoderms, including echinoids (Strongylocentrotus purpuratus and Dendraster excentricus) and ophiuroids (Fig. S4). The impact of SSaDV on these taxa is unknown; however, the presence of the virus suggests that they could form a reservoir of SSaDV. If some echinoderms are
tolerant reservoirs of infection, it might help keep SSaDV in the system long after it extirpates less tolerant hosts, helping explain how SSWD can extirpate some host species while still persisting in an ecosystem. However, we cannot eliminate the possibility that other non-commercial invertebrates are reservoirs for SSaDV.

SSaDV Present in Asteroids 72 Years Ago

Its broad host range suggested that SSaDV could be associated with asteroid mortalities in other times and places. To better understand the geographic distribution of SSaDV, we studied Northwest Atlantic Coast asteroids with SSWD-like signs in 2012 and 2013. We used qPCR targeting two loci (VP4 and NS1) on the SSaDV genome to detect SSaDV from diseased *Asterias forbesii* from the Mystic Aquarium (Connecticut). Although we did not detect SSaDV gene transcription, the SSaDV DNA detected in 9 of 14 samples suggests that this virus might be present in other oceanic basins. Additionally, to investigate whether SSaDV was present in Northeast Pacific Coast asteroids before 2013, we tested for SSaDV in ethanol-preserved museum specimens collected between 1923 and 2010 at sites in British Columbia, Washington, Oregon, and California (Table S6).

qPCR amplification detected SSaDV DNA (NS1 and VP4 loci) in asteroids that had been field-collected and preserved in July 1942, October 1980, September 1987, and July 1991 (Fig. 10). We also detected one locus in at least nine other individual asteroids, suggesting that viruses with homologous NS1 or VP4 were present in these populations, too. Therefore, SSaDV and related viruses might have infected asteroids on the North American West Coast decades before the current SSWD event.

If SSaDV is the cause of the current SSWD event, it is unclear why the virus did not elicit wide disease outbreaks in the past during periods in which it was detected; however, there are several possible reasons why the current SSWD event is broader and more intense than previous occurrences. SSaDV may have been present at lower prevalence for decades and only became an epidemic recently due to unmeasured environmental factors not present in previous years that affect animal susceptibility or enhance transmission. There are anecdotal reports from fishers and scuba divers that by 2012, the Salish Sea was severely overpopulated with adult *P. helianthoides*. Our finding of a strong relationship between size and SSaDV load, and anecdotal observations of SSaDV commonly isolated but less so in juveniles in the field, suggest that the current event may be exacerbated by a large number of adult sea stars present in small bays and inlets. Because of its wide host range in the current event, we also speculate that variation of SSaDV (possibly by modification of capsid structure, as seen in other parvoviruses) (18) may have led to greater virulence. There remains much to be learned about the interactive effects of environmental transport, virulence, and environment on finding virus and disease transmission to healthy asteroids after exposure in two trials with virus-sized particles from diseased asteroids, finding replicating densovirus in diseased tissue, and an association between viral load and disease. Furthermore, our observation of SSaDV from 72 y ago suggests that, like many marine pathogens, SSaDV was already present in the environment before the outbreak. The detection of SSaDV in diverse echinoderm species and some sediments suggests a high potential for persistence in nonasteroid reservoirs. SSaDV is present in environmental samples, suggesting that it can spread outside of a host. However, it remains to be seen how infection with SSaDV kills asteroids, what the role is for other microbial agents associated with dying asteroids, what triggers outbreaks, and how asteroid mass mortalities will alter near-shore communities throughout the North American Pacific Coast. More generally, viral pathogens are poorly known for all noncommercial invertebrates yet may play an unrecognized, yet major role, in marine ecosystems.

Conclusions

In summary, SSWD has caused widespread and, until now, unexplained mass mortality in asteroids. SSWD spread has been most consistent with an infectious agent, which we suggest is a virus. Based on our observations, the densovirus, SSaDV, is the most likely virus involved in this disease. We base this statement on finding virus and disease transmission to healthy asteroids after exposure in two trials with virus-sized particles from diseased asteroids, finding replicating densovirus in diseased tissue, and an association between viral load and disease. Furthermore, our observation of SSaDV from 72 y ago suggests that, like many marine pathogens, SSaDV was already present in the environment before the outbreak. The detection of SSaDV in diverse echinoderm species and some sediments suggests a high potential for persistence in nonasteroid reservoirs. SSaDV is present in environmental samples, suggesting that it can spread outside of a host. However, it remains to be seen how infection with SSaDV kills asteroids, what the role is for other microbial agents associated with dying asteroids, what triggers outbreaks, and how asteroid mass mortalities will alter near-shore communities throughout the North American Pacific Coast. More generally, viral pathogens are poorly known for all noncommercial invertebrates yet may play an unrecognized, yet major role, in marine ecosystems.

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