Virome temporal variation during sea star wasting disease progression in *Pisaster ochraceus* (Asteroidea, Echinodermata)

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**Abstract:** Sea star wasting disease (SSWD) is a condition that has affected asteroids for over 120 years, yet mechanistic understanding of wasting etiology remains elusive. We investigated temporal virome variation in two *Pisaster ochraceus* specimens that wasted in the absence of external stimuli and two specimens that did not experience SSWD for the duration of our study, and compared viromes of wasting lesion margin tissues to both artificial scar margins and grossly normal tissues over time. Global assembly of all SSWD-affected tissue libraries resulted in 45 viral genome fragments represented in >1 library. Genome fragments mostly matched densoviruses and picornaviruses with fewer matching nodaviruses, narnaviruses and sobemoviruses. Picornavirus-like and densovirus-like genome fragments were most similar to viral genomes recovered in metagenomic study of other marine invertebrates. Read recruitment revealed only 2 picornavirus-like genome fragments that recruited from only SSWD-affected specimens, but neither was unique to wasting lesions. Wasting lesion margin reads recruited to a greater number of viral genotypes (i.e. richness) than did either scar tissue and grossly normal tissue reads. Taken together, these data suggest that no single viral genome fragment was associated with SSWD. Rather, wasting lesion margins may generally support viral proliferation.

**Keywords:** Densovirus, Picornavirus, Nodavirus, Sea Star Wasting Disease, Asteroidea

1. Introduction

Sea star wasting disease (SSWD) describes a condition that has affected Asteroidea since at least 1898 [1] and is associated with periodic mass mortality, most recently as 2013-2014 [2]. The disease is pathognomic (i.e. has no distinguishing signs), where grossly abnormal specimens experience loss of turgor, abnormal limb twisting, epidermal lesions, body wall erosion, limb autotomy and in some cases death (reviewed in [3]). The cause of SSWD is currently unknown. Early work suggested association with a densovirus (the Asteroid ambidensovirus-1 [AaV-1]; also referred to as the “Sea Star associated Densovirus” [SSaDV]), and experiments which challenged healthy specimens with filtered tissue homogenates generated some SSWD signs [2, 4]. However, subsequent work found that densoviruses, including AaV-1 / SSaDV, occur in diverse asteroid taxa globally [5, 6], and are highly prevalent within communities inhabiting the northeast Pacific [6] and northwest Atlantic [7] oceans. Recent investigations suggest that wasting response of asteroids to tissue homogenate challenge could generate via non-pathogenic means (i.e. through organic matter enrichment resulting in suboxic conditions through heterotrophic respiration) [8]. Other proposed mechanisms of wasting, including repeated [9] and monotonic [10] temperature excursion, high pCO2 conditions [11], and low precipitation [5] have been hypothesized to influence SSWD. SSWD is not associated with any eukaryotic microorganism nor bacterium [4, 5, 12-14]. Microbiome studies during wasting progression suggest a progressive enrichment of copiotrophic bacteria on surfaces and within tissues [8, 12, 14] but none appear distinct only to affected specimens.
RNA virome studies of SSWD to date have focused on snapshots of viral diversity comparing grossly normal to wasting affected specimens. For example, comparisons of RNA viral composition between disease states in Pycnopodia helianthoides found no RNA viral family consistently associated with SSWD-affected specimens that were absent from grossly normal individuals [2]. Similarly, RNA metavirome surveys of wasting Pisaster ochraceus discovered several candidate RNA viral genotypes [5]. However, subsequent qPCR studies and read recruitment of these genotypes failed to yield significant association with disease [4]. Hence, while RNA viruses occur in grossly normal echinoderm specimens and wasting-affected asteroid specimens, their potential roles in SSWD etiology are poorly resolved.

To further examine the role of viruses in SSWD, we observed time-course progression of wasting in Pisaster ochraceus during a concurrent study in which specimens wasted in the absence of external stimuli [8]. Histopathologic findings of SSWD indicate that affected body wall tissues experienced ulceration, cleft formation and coelomocyte aggregation, along with necrosis and body wall degradation [2, 4, 13, 14]. We focused our study on body wall lesions, since these are the most conspicuous sign of SSWD, were the least subject to observer bias (c.f. deflation, limb curling), and lesions generally precede limb autotomy or body wall erosion. We hypothesized that viral etiologic agents would be would be present in wasting lesion margins, but absent in grossly normal tissues well away from lesions on the same individual, and absent in asteroid specimens that remained grossly normal during our study. Furthermore, we hypothesized that any SSWD-associated viral agent would be absent in physical scar margins in either SSWD-affected or grossly normal specimens, since these may be viruses that replicate during wound healing (c.f. cause pathogenesis). Our results demonstrate that there were no viral genome fragments which recruited reads only from wasting lesion margins in SSWD-affected asteroid specimens (i.e. all wasting lesion genome fragments also recruited reads from scar tissue or control tissues). Rather, we found a progressive enrichment of viral genotypes that recruited sequence reads in lesion margins over time in comparison to artificial scar and grossly normal tissues, suggesting their prominence in prior metaviromic surveys may have been independent of potential pathology.

2. Materials and Methods

Survey Design: Longitudinal survey of sea star wasting microbial ecology was performed in July – August 2018 and reported in separate studies focusing on associated bacteria [8]. We collected six Pisaster ochraceus specimens (mean mass 290 ± 54 g and ray length 11.2 ± 0.9 cm) from the intertidal zone at Davenport, CA (37°1’19”N, 122°12’56”W) on 19 July 2018, which were transported in insulated coolers to the Long Marine Laboratory at UC Santa Cruz and housed in flow-through aquaria indoors in individual containers. After 48 h acclimation, a small scar (~ 5 mm long) was made on a single ray of each specimen using a sterile 4 mm biopsy punch. Individual asteroids were monitored daily for the presence of lesions (which were defined as non-focal loss of epidermal tissues revealing the underlying body wall; see Fig. 1). After 72 h, artificial scar margin tissues (~ 3mm x 2mm) were collected using sterile 4mm biopsy punches. New artificial scars were made on adjacent rays each day, and sampled after each 24 h period. When wasting lesions were observed (the first wasting lesion was observed at 96 h), their margins were sampled following the same approach, and grossly normal tissues well away (> 1cm) from the lesion collected at the same time for comparison. A photographic summary of natural and wasting lesions is provided in Fig. 2. All tissue samples were placed into sterile 1.2 mL cryovials and immediately frozen in liquid N2 or in a -80°C freezer.
Figure 1. Gross examination of SSWD lesion on a *Pisaster ochraceus* specimen retrieved from Davenport, CA at the time of sampling for this survey. A = grossly normal tissue; B = lesion margin; C = lesion (underlying body wall tissues); D = papula and pedicellaria; E = paxilla (spine).
Figure 2. Gross changes in *Pisaster ochraceus* observed in specimens used in viral metagenome analyses over time, and detail (indicated by arrows) of wasting lesion and artificial scars sampled.

**Viral Metagenome Preparation:** We focused viral metagenomic analyses around four specimens -two that developed wasting lesions and died during the experiment (hereafter referred to as “SSWD-affected”), and two that remained grossly normal during the experiment (Table 1). Initial samples (taken at 0 h) were prepared for viral metagenomics from all four specimens. Viral metagenomes from SSWD-affected specimens were prepared from wasting lesion margin, artificial scar margin, and control tissue samples away from artificial scars and wasting lesions at the time of lesion genesis (96 and 288 h for wasting-affected specimens #1 and #2, respectively). In addition, viral metagenomes were prepared from wasting lesion margin samples taken from the SSWD-affected specimens at the time of death. Viral metagenomes were prepared from the two grossly-normal specimens at 0 h, and at 432 h from artificial scar margin tissues and tissue samples collected away from artificial scars.

Table 1. Viral metagenomics library characteristics for *Pisaster ochraceus* wasting temporal assay.
Control = grossly normal tissue; Scar = artificial scar margin; Lesion = wasting lesion margin.

| Library Name | Specimen Name | Date    | Tissue Type | Library Size (Reads) | Matching Viruses |
|--------------|---------------|---------|-------------|---------------------|------------------|
| SC1          | SSWD-affected 1 | 22-Jul-18 | Control     | 3,658,045           | 554,809          |
| SC2          | SSWD-affected 1 | 26-Jul-18 | Scar        | 4,099,676           | 582,091          |
| SC3          | SSWD-affected 1 | 26-Jul-18 | Control     | 3,592,838           | 597,395          |
| SC4          | SSWD-affected 1 | 26-Jul-18 | Lesion      | 3,255,385           | 169,582          |
| SC5          | SSWD-affected 1 | 27-Jul-18 | Lesion      | 3,443,200           | 871,889          |
| SC6          | SSWD-affected 2 | 22-Jul-18 | Control     | 4,948,712           | 1,007,543        |
| SC7          | SSWD-affected 2 | 04-Aug-18 | Control     | 2,703,665           | 624,989          |
| SC8          | SSWD-affected 2 | 04-Aug-18 | Scar        | 2,797,069           | 507,994          |
| SC9          | SSWD-affected 2 | 04-Aug-18 | Lesion      | 3,678,615           | 1,306,746        |
| SC10         | SSWD-affected 2 | 06-Aug-18 | Lesion      | 3,259,328           | 924,368          |
| SC11         | Grossly Normal 1 | 22-Jul-18 | Control     | 2,954,408           | 2,349            |
| SC12         | Grossly Normal 1 | 09-Aug-18 | Control     | 1,919,780           | 2,694            |
| SC13         | Grossly Normal 1 | 09-Aug-18 | Scar        | 1,939,732           | 3,134            |
| SC14         | Grossly Normal 2 | 22-Jul-18 | Control     | 3,273,980           | 3,851            |
| SC15         | Grossly Normal 2 | 09-Aug-18 | Control     | 1,595,206           | 1,112            |
| SC16         | Grossly Normal 2 | 09-Aug-18 | Scar        | 1,196,532           | 882              |

Tissue samples were prepared for viral metagenomics targeting RNA viruses as described previously [15-17]. In addition to RNA viruses, we also examined densovirus (i.e. ssDNA) viruses since they are also captured in libraries prepared using this approach [5]. Briefly, tissue samples were homogenized in 2 mL of 0.02 µm-filtered PBS by bead-beating (Zymo Bead Beaters), briefly centrifuged at 3,000 x g for 30s to remove large particulate matter, and then filtered through 0.2 um filters (Durapore) to remove cell debris. The resulting concentrate was treated with DNAsel 1 (5 U; Thermo Fisher Scientific), RNase One (50 U; Promega) and Benzonase (250 U; Sigma-Aldrich) for 3 h at 37°C to remove co-extracted free nucleic acids, before arresting enzyme activity with 50 µM virus-free EDTA. Viral nucleic acids were extracted using the Zymo Viral RNA kit, before amplification using the TransPlex Whole Transcriptome Amplification kit (Sigma Aldrich). Resulting products were electrophoresed and quantified by Pico Green fluorescence. Samples were then submitted to
Biotechnology Resource Center at Cornell University, where libraries were sequenced on 2 lanes of Illumina MiSeq (2 x 250 bp paired-end) after TruSeq PCR-free library preparation. Sequence libraries are available at NCBI under BioProject PRJNA637333 and SRA accessions SRR11931172- SRR11931187.

**Bioinformatic processing:** Sequence libraries were initially trimmed for adapters and quality (N<0.5). We used an assembly-read mapping approach to examine the presence and absence of viral genome fragments between libraries. First, a global assembly of all 10 samples from SSWD-affected specimens (Table 1) was performed using the CLC Genomics Workbench 4.0 (Qiagen) using a minimum overlap of 0.5 and similarity of 0.8. The resulting contig spectra was aligned against several boutique databases of RNA viruses, encompassing genomes and proteins of invertebrate viral groups, by BLASTx and tBLASTx [18, 19] as described elsewhere [17]. Sequence matches against any of these databases at an E-value<10^{-20} were further aligned against the non-redundant (nr) library at NCBI by BLASTx, and contigs discarded if they matched known bacterial or eukaryote proteins at a higher percentage and lower E-value than viruses. The resulting contig spectra (here termed “viral genome fragments”) were then subject to read recruitment independently across the 10 SSWD-affected specimen libraries and the additional 6 libraries from grossly normal specimens. Viral genome fragments that did not recruit reads from > 2 libraries or which recruited < 2 reads from libraries were discarded from further analysis. Because we did not standardize template nucleic acid quantities nor total viral abundance in amplification reactions, we were unable to gain quantitative insight into representation during wasting progression [20]. Hence, our work focuses only on the presence of viral genome fragments and their presence by read recruitment between viromes. Viral genome fragment sequences are available at NCBI under accessions MW073776-MW073820.

**Statistical analyses:** We analyzed the presence/absence of viral genome fragment recruits between: grossly-normal and SSWD-affected specimens; between initial, wasting lesion, artificial scar, and control tissues away from lesions/scars; and between timepoints on the same individual at the time of lesion genesis and death by performing Fisher’s Exact test to address the hypothesis that the recruitment to viral genome fragments in each specimen condition, tissue types, and sample time were independent. The variation in read recruitment richness between tissue types (initial, grossly normal at time of lesion genesis + conclusion of study, artificial scar at time of lesion genesis + conclusion of study, and wasting lesion at time of lesion genesis + death) was examined by performing pairwise Student’s t-tests with Bonferroni correction to account for Type II error in multiple comparisons. All analyses were performed in XLStat (AddinSoft GmBH).

### 3. Results and Discussion

Our temporal survey of virome composition during SSWD progression further advances understanding that echinoderms, including asymptomatic individuals, harbor rich viral assemblages but that there is little association between specific viral genotypes and wasting signs [5, 6]. Specimens that wasted in our survey experienced loss of turgor, followed by the appearance of epidermal lesions. Microscopic investigation (Fig. 1) showed that these were due to loss of epidermal tissues exposing the underlying body wall. Grossly, lesion margins were unremarkable and were not melanized. Between the time of first lesion appearance and animal death, SSWD-affected specimen #2 autotomized a ray, while SSWD-affected specimen #1 experienced body wall erosions that allowed internal organs (presumably pyloric caeca) to protrude. Wasting lesions were not grossly distinct to artificial scars (Fig. 2).

#### 3.1 Description of viruses recovered in viral metagenomes

Viral metagenomes prepared from 16 samples (Table 1) generated a total of 48,316,171 reads. Assembly of the 10 SSWD-affected specimen libraries (35,436,533 reads) resulted in 96,778 contigs. Of these, only 48 matched RNA or densoviral viral proteins or genomes by BLAST alignment (i.e. viral genome fragments). Next, we recruited these against each of the 10 symptomatic libraries separately, as well as to 6 asymptomatic specimen libraries. Forty-five contigs recruited against >1 library, suggesting that most viruses inhabiting asteroid tissues are cosmopolitan between
individuals within the *Pisaster ochraceus* population at the time of sampling. Viral contigs meeting these criteria were mostly similar to *Picornavirales* (*n* = 19 contigs), with fewer matches to *Piccovirales* (*Parvoviridae; Densovirinae;* *n* = 18 contigs), *Nodavirales* (*Nodaviridae;* *n* = 6 contigs), *Wolframvirales* (*Narnaviridae;* *n* = 1 contig) and *Sobelivirales* (*Solemoviridae;* *n* = 1 contig).

*Picornaviruses* (+ssRNA) feature prominently in most host-associated virome surveys [21-32] and are ubiquitous in marine plankton as free particles [33-39]. Of the 19 picornavirus-like genome fragments recovered in this survey, 9 matched most closely to dicistroviruses (*Dicistroviridae*), 2 to bacillarnaviruses (*Marnaviridae*), and 1 to caliciviruses (*Caliciviridae*) by BLASTx (Fig. 3). Furthermore, contigs clustered with iflaviruses (*Iflaviridae*; contig 91247) and marnaviruses (*Marnaviridae*; contig 12092) by phylogenetic analyses (Fig. 4). Picornavirus-like genome fragments mostly matched viral genomes recovered from transcriptomic surveys of marine invertebrates (mollusks and crustacea) [21, 40] and picoeukaryotes [41]. Picornaviruses have been previously recovered from asteroids [2, 5] and holothurians [17]. Since several genome fragments recovered in this survey matched most closely picornaviruses recovered from protists [41], it is possible that they were associated with protozoa associated with wasting tissues. Previous work has revealed the presence of presumably fungal viruses and a wide richness of protistan rRNAs in metagenomes prepared from material purified from < 0.2 µm filtered tissue homogenates [17]. Picornaviruses have been recovered in stressed marine metazoa [42, 43], cause mortality in protists (e.g. diatom viruses [44, 45]) and disease in marine arthropods (e.g. Taura syndrome virus [46]). However, the wide diversity of picornaviruses recovered from grossly normal specimens in field surveys [21] suggests that their role in disease, especially in mass mortality settings, is unclear for most hosts.

**Figure 3.** Maps of contiguous sequences matching *Picornavirales* recovered in this survey of *Pisaster ochraceus*. Contigs were annotated based on BLASTx (e-value < 1 x 10^-20) against the non-redundant database at NCBI. The color of arrows (open reading frames) indicates the taxonomic identity of their best matches. Numbers above the ORFs indicate the e-value of BLAST results. The total contig lengths are indicated by solid lines running through and between ORFs.
Figure 4. Phylogenetic representations of *Picornavirales*-like genome fragments recovered from *Pisaster ochraceus*. The trees were constructed based on a 98 amino acid (A) and 193 amino acid (D-E) alignment of the RNA dependent RNA polymerase gene, a 187 amino acid alignment of the rhv-like capsid domain (B) and a 160 amino acid alignment of the RNA helicase domain (C) and performed separately for overlapping regions including best matches at NCBI. Trees were constructed by Neighbor Joining and based on Jukes-Cantor distance. Bootstrap values > 50% (based on 1000 iterations) are indicated above nodes. The host identity is indicated by symbols next to branch labels. An additional phylogenetic representations of each tree based on maximum likelihood is presented in Supplemental Fig. 2.

Nodaviruses (+ssRNA) represent significant pathogens of marine vertebrates (reviewed in [47, 48]) and invertebrates [49] and are frequently recovered in grossly normal marine invertebrates [50-53]. Six viral contigs matching most closely nodaviruses by BLASTx (Supplemental Fig. 1) were recovered in our survey. Phylogenetic analyses placed these most similar to alphanodaviruses recovered from cnidarians, nematodes and arthropods as part of transcriptomic viral discovery efforts [21] (Fig. 5). Nodavirus-like genome fragments were identified in the asteroid *Pycnopodia helianthoides* [2] and were present in wasting-affected *Pisaster ochraceus* libraries prepared from the Olympic National Park in 2013 (NCBI BioProject SAMN15704856; tBLASTx e < 1 x 10^{-20}) [5], however
were absent in a RNA viral metagenomic survey of holothurians [17]. Our observation of alphanodavirus genome fragments in asteroids extend the known host range of the Nodamuravirales.

![Figure 5](image_url)

**Figure 5.** Phylogenetic representations of Nodamuravirales-like genome fragments recovered from *Pisaster ochraceus*. The trees were constructed based on: a 234 amino acid alignment of the methyltransferase domain (A); and a 101 amino acid (B-C) and 559 amino acid (D) alignments of the RNA dependent RNA polymerase gene of the nodavirus RNA1 genome fragment including best matches at NCBI. Trees were constructed by Neighbor Joining and based on Jukes-Cantor distance. Bootstrap values > 50% (based on 1000 iterations) are indicated above nodes. The host identity is indicated by symbols next to branch labels. An additional phylogenetic representation of each tree based on maximum likelihood is presented in Supplemental Fig. 3.

Densovirus (Piccovirales; Parvoviridae, ssDNA) have been previously recovered from marine arthropods (reviewed in [54, 55]), mollusks [56, 57], a tunicate [58] and echinoderms [2, 5-7]. Densovirus are associated with mussel [57] and asteroid [2] but see [6]) mass mortality. Contigs matching densoviral proteins and genomes recovered in this survey bore primarily structural ORFs and ORFs bearing the non-structural (NS; i.e. replication-associated protein) 1 regions, with fewer NS2 and NS3 regions (Fig. 6). Genome fragments bearing both structural and non-structural regions (n = 3) bore ambisense genome architecture, suggesting these belonged to the Ambidensovirus genus to which almost all known marine invertebrate densovirus belong [2, 6, 7, 56, 59]. Alignment of structural (coat protein) placed detected densovirus genome fragments within the ambidensovirus genus (Fig. 7). Importantly, we did not recover any genome fragment bearing > 85 % nucleotide identity to Asteroid ambidensovirus 1 (i.e. SSaDV).
Figure 6. Map of densovirus-like genome fragments recovered from *Pisaster ochraceus* from Davenport, CA in July 2018. The length of contig is given by the solid black line running through open reading frames (ORFs; indicated by arrows). The color of arrow indicates the top BLASTx match to the non-redundant database at NCBI, and e-value of the match given above each ORF.

Figure 7. Phylogenetic representations of *Densovirinae*-like genome fragments recovered from *Pisaster ochraceus*. The trees were constructed based on: a 103 amino acid alignment of the structural (coat protein) gene; a 83 amino acid of the non-structural 1 (NS1) gene; a 112 amino acid of the NS2 gene; and a 111 amino acid of the NS3 gene. Phylogenetic representations include best matches by BLASTx against the non-redundant database at NCBI. Trees were constructed by Neighbor Joining and based on Jukes-Cantor distance. Bootstrap values > 50% (based on 1000 iterations) are indicated above nodes. The host identity is indicated by symbols next to branch labels. An additional phylogenetic representation of each tree based on maximum likelihood is presented in Supplemental Figs. 4.
We also recovered two genome fragments matching a narnavirus (Behai narna-like virus 7 NCBI APG77081.1; BLASTx against nr database e-value = $6 \times 10^{-64}$) which was recovered from a razor shell bivalve (presumably Pharidae [21]) and a sobemovirus (Solemoviridae; Rice Yellow Mottle Virus NCBI CAE81305.1; BLASTx against nr database e-value = $1 \times 10^{-35}$) [60]. These likely represent viruses of fungi and plants, respectively. Narnaviruses cause asymptomatic infections in fungi. In Rhizopus microspores, narnaviruses promote sexual reproduction and suppress asexual reproduction in concert with a third endosymbiotic member, the bacterium Mycetohabitans spp. [61]. Sobemoviruses cause a wide range of plant diseases, including mottles and mosaics (reviewed in [62]). Both are unlikely to infect echinoderm cells and instead likely infected microbiome constituents.

3.2 Analysis of virome association with Sea Star Wasting Disease

Of 45 viral genome fragments recovered in this survey, all recruited reads from SSWD-affected asteroids, but only 37 genome fragments recruited reads from grossly normal specimens (Fig. 8). All but 3 of the 45 genome fragments recruited reads from all tissue types. Only two genome fragments (contigs 12092 and 12093), both Picornavirales (putatively assigned to Marnaviridae and Dicistroviridae) recruited reads only from SSWD-affected specimen libraries (Table 2). However, both genome fragments did not uniquely recruit from wasting lesion margin libraries and also recruited reads from artificial scar margin and control tissue libraries. Recruitment to nodavirus-like contig 91444 was significantly associated with wasting lesion margins (p = 0.019; Fisher’s Exact test), but also recruited reads from artificial scar tissues in grossly normal star #1 at the survey conclusion (Tf). Recruitment to contigs 88293 and 2401 (Densovirinae) and 12093 (Picornavirales) were significantly associated with time of sampling (i.e. mostly present in later samples), but also recruited reads from both grossly normal specimens and SSWD-affected specimens. Hence, our data do not support association between any viral genome fragment recovered in this survey and SSWD since none was unique to either wasting lesion margins or to SSWD-affected specimens.

The role of viruses in asteroid wasting etiology has been controversial. Early association between the Sea Star Associated Densovirus (Asteroid ambidensovirus 1; AaV-1 [2]) was not supported by subsequent work [5]. The discrepancy between studies was attributed to inaccurate primer design on investigation outset (providing false positives as a result of a background of ambient densovirus strains), the low numbers of individuals sampled, and presence of AaV-1 in asymptomatic individuals. Further investigation of densoviruses in northwest Atlantic Ocean asteroids revealed the presence of persistent infection by a related strain of AaV-1 (Asteroid ambidensovirus 2) [7], and densoviruses have been recovered from asteroid tissue metagenome surveys elsewhere [5], suggesting that densoviruses may be common constituents of the asteroid microbiome. Of the 19 densovirus genome fragments recovered in the survey presented in this work, all recruited reads from >1 specimen, and 10 were detected in all 4 specimens. These results illustrate that densoviruses may be cosmopolitan within Pisaster ochraceus populations.
Figure 8. Heat map of viral contig read recruitment across all libraries in non-wasted and wasted asteroids. Dark cells = viral contig recruited reads from library, white cells = viral contig did not recruit reads from library. T0 = initial sample, Ti = time of lesion formation, Tf = experiment termination. C = grossly normal tissue, S = artificial scar tissue, L = wasting lesion margin. Phylogeny determined by family-level assignment based on nearest relative match (BLASTx) against non-redundant (nt) database at NCBI.
Interestingly, the number of viral genome fragments recruiting from natural lesion libraries was significantly greater (p<0.008, Student’s t-test, df = 4) than both artificial lesion and control (i.e. away from lesions on SSWD-affected specimens) tissues (Fig. 10). Previous work has noted taxonomic variation in host-associated viral communities in response to stress. For example, Laffy et al [63] examined the impacts of thermal stress on the sponge Rhopaloeides odorabile and observed the proliferation of Calimoviruses and Retroviruses relative to controls. Additionally, Grasis et al [64] noted that the greatest viral diversity was observed in heat-stressed Hydra spp. when compared to controls. Vega Thurber et al. [65] noted an increase in herpesvirus-like sequences in stressed corals. While it is tempting to ascribe this result to enhanced susceptibility to opportunistic pathogens in compromised cells, it is more likely that our observation relates to factors affecting viral replication of normally asymptomatic viral infections. Viral replication in affected tissues is a complex interaction with intracellular properties and environmental cues. All viral groups detected in this study, including the Densovirinae, replicate in actively dividing host cells. Hence, the increase in viral richness in affected tissues, which were not grossly hyperplastic (i.e. did not display gross signs of rapid cell division), is surprising. Wound repair would presumably be associated with an increase in gene transcription in affected tissues, which in turn cause rapid replication of viruses in infected cells.

Table 2. Fisher’s exact test results for viral contigs comparing condition (SSWD-affected vs grossly normal), sampling time (initial, time of lesion formation, time of death or experiment termination), and sample tissue type (control, artificial scar, wasting lesion). Only contigs returning any significant (p< 0.05) result are reported.

| Contig #     | Condition | Time   | Sample Type |
|--------------|-----------|--------|-------------|
| Contig 82293 | ns        | 0.05   | ns          |
| Contig 2401  | ns        | 0.023  | ns          |
| Contig 12092 | 0.001     | ns     | 0.022       |
| Contig 12093 | 0.003     |        | 0.019       |
| Contig 91444 | ns        |        |             |

Figure 9. Richness of viral genome fragments recruiting reads from SSWD-affected and grossly normal tissues (±SE) in viral metagenomes prepared from P. ochraceus during temporal study of wasting. Significance (a,b) determined by Student’s t-test (p<0.008, df=4 with Bonferroni correction for 6 tests). T0 = initial sample, Ti = time of first lesion appearance, Tf = experiment termination.
Recently we observed that SSWD is associated with a proliferation of copiotrophic bacteria near and on their respiratory surfaces, concomitant with and followed by the presence of facultative and strictly anaerobic bacterial taxa [8]. Additionally, SSWD susceptibility correlates with morphological features which influence surface boundary layer extent and respiratory demand compared to theoretical diffusion, which taken together are evidence that wasting is associated with suboxic conditions at the animal-water interface [8]. O₂ tension in cells triggers production of many viruses [66]. For example, flaviviruses and nucleocytoplasmic large DNA viruses use hypoxia inducible factors (HIF; genetic switches and genes that activate under hypoxic conditions), to stimulate production [67-72]. Hence, suboxic stress may influence the proliferation of viruses in SSWD-affected asteroids.

The greater richness of viral genome fragments that recruited reads in SSWD-affected tissues may also relate to the proportion of contaminating co-extracted host nucleic acids in SSWD-affected and grossly normal specimens. Total extracted DNA quantities decrease in affected specimens [20]. Viral nucleic acids, which may be more protected from enzymatic decay within capsids than host RNAs, may become more pronounced when host tissues degrade. Hence, viral nucleic acids may recruit more regularly to viral genome fragments solely because they comprise a greater proportion of virome sequence space.

4. Conclusions

Our results illustrate that both grossly normal and SSWD-affected asteroids are associated with RNA viruses which are similar to those recovered in metagenomic and metatranscriptomic surveys of marine invertebrates performed elsewhere. SSWD in Pisaster ochraceus is not associated with any specific viral genotype detected in this survey. Rather, wasting is associated with an increased richness of viral genome fragments recruiting reads in affected tissues, which may be due to factors influencing their replication or due to the balance between host and viral RNA in tissues. This work provides additional evidence that densoviruses, and particularly the Asteroid ambidensovirus 1 is not consistently associated with sea star wasting, and emphasizes the lack of association more generally between viruses and SSWD. Our work raises interesting questions about the influence of extrinsic factors, e.g. processes driving non-infectious diseases, in influencing the replication and perhaps pathology of viruses in marine diseases.

**Supplementary Materials:** The following are available online at www.mdpi.com/xxx/s1, Figure S1: Nodavirus-like genome fragment contig map, Figure S2: Phylogenetic reconstruction of Picornavirales-like genome fragments by Maximum Likelihood, Figure S3: Phylogenetic reconstruction of Nodamuraivirales-like genome fragments by Maximum Likelihood, Figure S4: Phylogenetic reconstruction of Piccovirales-like genome fragments by Maximum Likelihood.

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