Development of a Serological Assay for the Sea Lion (Zalophus californianus) Anellovirus, ZcAV

Elizabeth Fahsbender¹, Karyna Rosario¹, John P. Cannon², Frances Gulland³, Larry J. Dishaw² & Mya Breitbart¹,²

¹University of South Florida, College of Marine Science, 140 7th Avenue South, Saint Petersburg, FL 33701 USA, ²University of South Florida, College of Medicine, Department of Pediatrics, 801 6th Street South, Saint Petersburg, FL 33701 USA, ³The Marine Mammal Center, 2000 Bunker Road, Sausalito, CA 94965 USA.

New diseases in marine animals are emerging at an increasing rate, yet methodological limitations hinder characterization of viral infections. Viral metagenomics is an effective method for identifying novel viruses in diseased animals; however, determining virus pathogenesis remains a challenge. A novel anellovirus (Zalophus californianus anellovirus, ZcAV) was recently reported in the lungs of captive California sea lions involved in a mortality event. ZcAV was not detected by PCR in the blood of these animals, creating the inability to assess the prevalence of ZcAV in live sea lions. This study developed an enzyme-linked immunosorbent assay (ELISA) to detect antibodies to ZcAV in sea lion serum. To assess ZcAV prevalence, paired serum and lung samples (n = 96) from wild sea lions that stranded along the California coast were tested through ELISA and PCR, respectively. Over 50% of the samples tested positive for ZcAV by ELISA (34%), PCR (29%), or both (11%) assays. ZcAV is prevalent in stranded wild sea lion populations and results suggest that PCR assays alone may grossly underestimate ZcAV exposure. This ELISA provides a tool for testing live sea lions for ZcAV exposure and is valuable for subsequent studies evaluating the potential pathogenicity of this anellovirus.

The rate of emergence of new diseases in marine animals is increasing¹–⁴, generating a need for surveillance of potential pathogens to protect marine mammals against epidemics. However, it remains difficult to characterize and diagnose viral infections because of methodological limitations⁵. Current detection methods such as degenerate PCR and pan-viral microarrays can detect close relatives of previously described viruses, but are limited for detecting novel viruses. Viral metagenomics (virus particle purification followed by shotgun sequencing) is an effective method for identifying viruses involved in mortality events in marine animals⁶–⁹, yet it remains difficult to establish a connection between a novel virus and disease due to limitations of culturing the viruses as well as difficulties in obtaining fresh diagnostic tissues from wild marine mammals.

Viral metagenomics performed on lung tissue of several necropsied captive California sea lions (Zalophus californianus) with signs of respiratory disease that were involved in a mortality event of unknown cause revealed the presence of a novel anellovirus, Zalophus californianus anellovirus (ZcAV)⁷. ZcAV was found by specific PCR in the lungs of all three of the sea lions that died in the mortality event, but it was not found in sea lions from the same zoo that died of unrelated causes. In addition, ZcAV was found to be actively replicating in lung tissue of a sea lion from the mortality event, further suggesting an association of ZcAV with the death of these animals. In addition to the captive sea lions, 11% of lung samples from wild sea lions stranded off the California coast tested positive for ZcAV by PCR, indicating that this anellovirus is present in wild populations.

The initial ZcAV discovery and prevalence studies have raised many questions about the potential pathogenicity of this virus. Anelloviruses have been extensively studied in humans, where these viruses can be highly prevalent (infecting up to 100% of the population)¹⁰–¹³, yet they have not been linked to human disease. Although anelloviruses have also been found in a wide range of mammals including non-human primates¹⁴, domestic animals¹⁵, Pacific harbor seals⁶, and Risso’s dolphins (E.M. Fahsbender et al., unpublished data), anellovirus-associated pathology remains unknown. Initial evidence suggested that ZcAV may be linked to the mortality event of captive California sea lions; however, studies investigating the pathogenicity of this virus are difficult since ZcAV has only been detected by PCR in tissues of necropsied animals, and cannot be detected by PCR in the blood of infected individuals⁷. The inability to detect ZcAV in blood samples by PCR severely limits further
testing for this virus, since obtaining lung biopsies from live sea lions is not possible. Hence, there is a critical need for an assay to detect ZcAV exposure in blood samples to investigate the epidemiology of this virus, understand its association with disease, and preemptively control this virus, understanding its epidemiology. To overcome these technical limitations of studying the role of ZcAV in disease, here we describe the development of an enzyme-linked immunosorbent assay (ELISA) for ZcAV, and demonstrate that sea lions mount an immune response against ZcAV.

Similar to other anelloviruses, ZcAV contains a small (2140 nucleotide (nt)), negative sense, single-stranded DNA circular genome that encodes three major open reading frames (ORFs). Based on similarities to other anelloviruses, ORF 1 is believed to encode the capsid protein, although this has not been experimentally demonstrated for any anellovirus. For other anelloviruses, the ORF 1 gene product has been predicted to be antigenic due to the presence of major hydrophilic regions, and ORF 1 has been successfully used in sero-prevalence studies in humans and pigs1,17,18. Here we developed an ELISA based on hydrophilic regions of the ORF 1 gene product of the ZcAV genome and demonstrated that this assay is capable of detecting anti-ZcAV antibodies in sea lion serum. This ELISA provides a tool for studying ZcAV epidemiology and identifying seroconversion upon symptom development in captive sea lions, enabling future research investigating the pathogenesis of ZcAV. Finally, the creation of this assay lays the groundwork for bridging the gap between genome discovery via viral metagenomics and assessing the epidemiology of novel viruses and their significance for wild populations.

Methods

Sample collection. Paired serum and lung samples were collected from 100 California sea lions involved in stranding events along the coast of northern California. All samples were collected by The Marine Mammal Center located in Sausalito, California in accordance with Marine Mammal Protection Act permit no 932-1905/MA-009526 to Gulland. Serum samples were collected during routine clinical examinations of live stranded sea lions as described by Bossart et al.21 and archived at 70°C. Lung samples were collected from the same animals upon necropsy, with animals stored at 4°C between time of death and necropsy (within 24 hours of death). All protocols were approved under the Marine Mammal Protection Act. Paired lung and serum samples were shipped to the University of California, Los Angeles for processing using PCR (for lung samples) and ELISA (for serum samples).

PCR testing of sea lion lung samples. Lung samples were tested with a ZcAV-specific PCR assay. For this purpose, DNA was extracted from approximately 25 mg of sea lion lung tissue using the DNeasy Blood and Tissue Kit (Qiagen). DNA was amplified through rolling circle amplification (RCA) using the Illustra TempliPhi Kit (GE Healthcare), which is known to enrich for small circular templates such as the ZcAV genome22,23. RCA products were then used as the DNA templates for the PCR assay using primers ZcAV1056F (5′-AGG CAC TCA GAT CTA TCT AA-3′) and ZcAV1369R (5′-CCC AGG CAT TAC AGG CTT TA-3′) designed to target a 314 nt region from ORF 1 of ZcAV. The PCR (containing 1 μM of each primer, 200 μM of dNTPs, 1 U of Red Taq (Sigma-Aldrich), 1X Red Taq reaction buffer and 2.5 μl of DNA template in a 25 μl reaction) proceeded as follows: 95°C for 5 minutes, 45 cycles of [94°C for 1 minute, 54°C for 1 minute, 72°C for 1 minute], followed by a final extension at 72°C for 10 minutes. PCR products of the correct size were verified by gel electrophoresis. Four samples were cloned using a TOPO TA Cloning Kit (Invitrogen) and commercially sequenced. Sequences were then compared against the GenBank database in order to verify the specificity of the assay. To determine assay sensitivity, a positive control was prepared as previously described. The PCR product was cleaned using the UltraClean PCR Clean-Up Kit (Mo Bio Laboratories) and quantified using a Nanodrop ND-1000 apparatus using ultraviolet light absorption (NanoDrop Technologies). The number of targets per microliter was back-calculated and the positive control was serially diluted to determine the sensitivity of the assay, which was 10 targets.

Optimization of ELISA parameters. An ELISA was developed to screen sea lion serum for antibodies to ZcAV. Since the virus was identified from a mixed community using metagenomics, there were no purified virus particles that could be used as an antigen for the ELISA. Therefore, the ELISA was developed based on regions of ORF 1 that have been used previously in sea lion serological testing17,18. Checkboard titrations were performed to find optimal concentrations of peptide, serum, and proteins. Similar to other anelloviruses16, ZcAV contains a small (2140 nt), negative sense, single-stranded DNA circular genome that encodes three major open reading frames (ORFs). Based on similarities to other anelloviruses, ORF 1 is believed to encode the capsid protein, although this has not been experimentally demonstrated for any anellovirus. For other anelloviruses, the ORF 1 gene product has been predicted to be antigenic due to the presence of major hydrophilic regions, and ORF 1 has been successfully used in sero-prevalence studies in humans and pigs1,17,18. Here we developed an ELISA based on hydrophilic regions of the ORF 1 gene product of the ZcAV genome and demonstrated that this assay is capable of detecting anti-ZcAV antibodies in sea lion serum. This ELISA provides a tool for studying ZcAV epidemiology and identifying seroconversion upon symptom development in captive sea lions, enabling future research investigating the pathogenesis of ZcAV. Finally, the creation of this assay lays the groundwork for bridging the gap between genome discovery via viral metagenomics and assessing the epidemiology of novel viruses and their significance for wild populations.

Results

Development of the ELISA using serum samples from sea lions whose lung samples were PCR-positive for ZcAV demonstrated that sea lions mount an immune response to ZcAV infection. Paired serum and lung samples from 100 stranded California sea lions were then tested for the presence of ZcAV nucleic acids and anti-ZcAV antibodies through PCR and the newly developed ELISA, respectively. Four samples were removed from the data set due to high background with the ELISA. Of the 96 samples tested, 50 (52%) tested positive for ZcAV exposure (ELISA; serum samples; n = 33 (34%) and/or DNA presence (PCR; lung samples; n = 28 (29%)) (Table 1). Eleven percent of the sea lions tested positive for ZcAV by both ELISA and PCR, and 48% of sea lions tested negative by both methods. Of the 28 sea lions with PCR-positive lung samples, 39% were also serum-positive by ELISA. However, 32% of the PCR-negative sea lions (n = 68) also tested positive for ZcAV exposure via ELISA. The Kappa value of 0.066, calculated using GraphPad, indicates poor correlation between the PCR and ELISA test results.

Discussion

The ELISA developed in this study provides the first and only method to test for ZcAV exposure in live sea lions. Here, this assay was used in combination with PCR to assess the prevalence of ZcAV in the wild, stranded sea lion population along the northern coast of California. Protein-L-based detection of ZcAV-specific antibodies suggests the exposed sea lions are able to recognize and mount immune responses to the virus. These data, which are the first obtained for marine mammal anelloviruses, support prior studies showing that human anelloviruses are immunogenic, with both IgG and IgM class antibodies produced upon exposure to ORF 17,18. The combined ELISA and PCR results demonstrate that ZcAV infections are prevalent and persistent in wild California sea lions.

The development of this ELISA was unconventional due to the many difficulties of working with marine mammals. Gaining access to 100 paired serum and lung samples from a protected species is both arduous and time consuming. Furthermore, the sea lion...
Anelloviruses are known to demonstrate a high degree of genetic variability. The ZcAV infection occurred in another sea lion organ instead of the lungs, we could expect a PCR-negative but ELISA-positive result. Conversely, sea lion samples may have tested positive by PCR but negative by ELISA (18%) if the sea lions were actively infected upon standing and died before a detectable antibody response was mounted. Although the timing of the response of the sea lion immune system to ZcAV is unknown, development of the human antibody response to TTV is slow and known to take up to 21 weeks after virus inoculation, although once the antibodies are produced they persist in serum for long periods. Finally, slight sequence mutations or antigenic diversity of anelloviruses may also contribute to data incongruency. Anelloviruses are known to demonstrate a high degree of genetic variability which can pose problems for both DNA detection by PCR primers and antibody detection. For certain TTV genotypes, it has been shown that multiple serotypes may exist in circulation.

Future work should compare the diversity of anelloviruses that can be detected by the PCR and ELISA utilized in this study, as the discrepancy between the two assay types may reflect differences in the diversity of genotypes and serotypes recovered.

Through a combination of PCR- and ELISA-based testing, this study demonstrated that sea lions mount an immune response to the anellovirus ZcAV and revealed the high prevalence of ZcAV in a stranded wild sea lion population. The ELISA created here will enable future research on the epidemiology of ZcAV in live sea lions and allow further study of ZcAV pathogenesis by measuring serocorrelation in infected captive sea lions. Finally, this assay has direct implications for protecting sea lion health since it is the only method available for the detection of ZcAV in live sea lions. The ZcAV ELISA enables the screening of rehabilitation animals before admitting them into facilities with healthy animals, which can prevent the spread of this virus to other captive sea lions and provide a valuable management step if further work links ZcAV with disease.

Table 1 | Prevalence of ZcAV in paired serum and lung samples from a stranded wild population of California sea lions (Zalophus californianus; n=96) assessed by ELISA and PCR, respectively

| Lung PCR positive n (%) | ELISA positive n (%) | Serum ELISA positive n (%) | Serum ELISA negative n (%) | PCR totals n (%) |
|------------------------|----------------------|---------------------------|---------------------------|-----------------|
| 11 (11%)               | 17 (18%)             | 46 (48%)                  | 46 (48%)                  | 63 (66%)        |
| 22 (23%)               |                      |                           |                           |                 |
| 33 (34%)               |                      |                           |                           |                 |

1. Harvell, C. D. et al. Emerging marine diseases—climate links and anthropogenic factors. *Science* **285**, 1505–1510 (1999).
2. Gulland, F. D. & Hall, A. Is marine mammal health deteriorating? Trends in the global reporting of marine mammal disease. *EcoHealth* **4**, 135–150 (2007).
3. Jones, K. E. et al. Global trends in emerging infectious diseases. *Nature* **451**, 990–993 (2008).
4. Keeling, F. et al. Impacts of biodiversity on the emergence and transmission of infectious diseases. *Nature* **468**, 647–652 (2010).
5. Delwart, E. L. Viral metagenomics. *Rev Med Virol* **17**, 115–131 (2007).
6. Ng, T. F. F. et al. Discovery of a novel single-stranded DNA virus from a turtle fibropapilloma by using viral metagenomics. *J Virol* **83**, 2500–2509 (2009).
7. Ng, T. F. F., Suedmeyer, W. K., Wheeler, E., Gulland, F. & Breitbart, M. Novel anellovirus discovered from a mortality event of captive California sea lions. *J Gen Virol* **90**, 1256–1261 (2009).
8. Li, L. et al. The fecal viral flora of California sea lions. *J Virol* **85**, 9909–9917 (2011).
9. Ng, T. F. F. et al. Metagenomic identification of a novel anellovirus in Pacific harbor seal (*Phoca vitulina richardsii*) lung samples and its detection in samples from multiple years. *J Gen Virol* **92**, 1318–1323 (2011).
10. Prescott, L. E., Simmonds, P. & Collaborators International. Global distribution of transfusion-transmitted virus. *New Engl J Med* **339**, 776–777 (1998).
11. Handa, A., Dickstein, B., Young, N. S. & Brown, K. E. Prevalence of the newly described human circovirus, TTV, in United States blood donors. *Transfusion* **40**, 245–251 (2000).
12. Biagini, P. et al. Distribution and genetic analysis of TTV and TTMV major phylogenetic groups in French blood donors. *J Med Virol* **78**, 298–304 (2006).
13. Okamoto, H. in *TT Viruses Vol. 331 Current Top Microbiol* (eds Ethel-Michele de Villiers & Haraldzur Hausen) Ch. 1, 1–20 (Springer Berlin Heidelberg, 2009).
14. Romeo, R. et al. High prevalence of TT virus (TTV) in naive chimpanzees and in hepatitis C virus-infected humans: frequent mixed infections and identification of new TTV genotypes in chimpanzees. *J Gen Virol* **81**, 1001–1007 (2000).
15. Biagini, P. et al. Circular genomes related to anelloviruses identified in human and animal samples by using a combined rolling circle amplification/sequence-independent single primer amplification approach. *J Gen Virol* **88**, 2696–2701 (2007).
16. Biagini, P. in *TT Viruses Vol. 331 Current Top in Microbiol* (eds Ethel-Michele de Villiers & Haraldzur Hausen) Ch. 2, 21–33 (Springer Berlin Heidelberg, 2009).
17. Huang, Y. W. et al. Expression of the putative ORF1 capsid protein of Torque teno sus virus 2 (TTSuV2) and development of western blot and ELISA serodiagnostic assays: Correlation between TTSuV2 viral load and IgG antibody level in pigs. *Virus Res* **158**, 79–88 (2011).
18. Huang, Y.-W. et al. Serological profile of torque teno sus virus species 1 (TTSuV1) in pigs and antigenic relationships between two TTSuV1 genotypes (1a and 1b), between two species (TTSuV1 and -2), and between porcine and human anelloviruses. *J Virol* **86**, 10628–10639 (2012).
19. Okamoto, H. et al. The entire nucleotide sequence of a TTV isolate from the United States (TUS01): Comparison with reported isolates and phylogenetic analysis. *Virology* **259**, 437–448 (1999).
20. Ott, C. et al. Use of a TTV virus ORF1 recombinant protein to detect anti-TTV virus antibodies in human sera. *J Gen Virol* **81**, 2949–2958 (2000).
21. Bossart, G. D., Reidarson, T. H., Dierau, L. A. & Duffield, D. A. in CRC Handbook of Marine Mammal Medicine 2nd edn (eds Dierau, L.A. and Gulland, F.M.D.) (CRC Press) 2001.
22. Kim, K.-H. et al. Amplification of uncultured single-stranded DNA viruses from rice paddy soil. *Appl Environ Microbiol* **74**, 5975–5985 (2008).
23. Kim, K.-H. & Bae, J.-W. Amplification methods bias metagenomic libraries of uncultured single-stranded and double-stranded DNA viruses. *Appl Environ Microbiol* **77**, 7663–7668 (2011).
24. Kessler, S. W. Rapid isolation of antigens from cells with a staphylococcal protein A-antibody adsorbent: Parameters of the interaction of antibody-antigen complexes with protein A. *J Immunol* **115**, 1617–1624 (1975).
25. Nilsson, B. H., Solomon, A., Björck, L. & Akerström, B. Protein L from *Pectrostreptococcus magnus* binds to the kappa light chain variable domain. *J Biol Chem* **267**, 2234–2239 (1992).
26. De Chateau, M. et al. On the interaction between protein L and immunoglobulins of various mammalian species. *Scand J Immunol* **37**, 399–405 (1993).
27. Tsuda, F. et al. Determination of antibodies to TT virus (TTV) and application to blood donors and patients with post-transfusion non-A to G hepatitis in Japan. *J Virol Methods* **77**, 199–206 (1999).
28. Tsuda, F. et al. IgM-class antibodies to TT virus (TTV) in patients with acute TTV infection. *Hepatol Res* **19**, 1–11 (2001).
29. Tawara, A. et al. Transmission of human TT virus of genotype 1a to chimpanzees with fecal supernatant or serum from patients with acute TTV infection. *Biochem Biophys Res Commun* **278**, 470–476 (2000).
30. Maggi, F. & Bendinelli, M. in TT Viruses Vol. 331 Current Top Microbiol (eds Ethel-Michele de Villiers & Harald zur Hausen) Ch. 5, 65–90 (Springer Berlin Heidelberg, 2009).
31. Okamoto, H. et al. Molecular cloning and characterization of a novel DNA virus (TTV) associated with posttransfusion hepatitis of unknown etiology. Hepatol Res 10, 1–16 (1998).
32. Hijikata, M., Takahashi, K. & Mishiro, S. Complete circular DNA genome of a TT virus variant (isolate name SANBAN) and 44 partial ORF2 sequences implicating a great degree of diversity beyond genotypes. Virology 260, 17–22 (1999).

Acknowledgments
This project was funded through an Interdisciplinary Research Development Grant from the University of South Florida’s College of Marine Science. The authors thank Dr. Michael Shamblott, Marci O’Driscoll, and Elizabeth Wheeler for guidance with experiments, use of equipment, and feedback on the manuscript, and the staff and volunteers of The Marine Mammal Center for care and sampling of the stranded sea lions.

Author contributions
E.F. developed the ELISA, tested samples, and wrote the manuscript. F.G. collected the sea lion samples. K.R., M.B., J.C. and L.D. advised and aided with the development of the ELISA and data analysis. All authors reviewed the manuscript.

Additional information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Fahsbender, E. et al. Development of a Serological Assay for the Sea Lion (Zalophus californianus) Anellovirus, ZcAV. Sci. Rep. 5, 9637; DOI:10.1038/srep09637 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder in order to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/