Genetic heterogeneity of dolphin morbilliviruses detected in the Spanish Mediterranean in inter-epizootic period

Consuelo Rubio-Guerri1,2*, M. Ángeles Jiménez3, Mar Melero1, Josué Díaz-Delgado4, Eva Sierra4, Manuel Arbello4, Edwige N. Bellière5, Jose L. Crespo-Picazo2, Daniel García-Párraga2,6, Fernando Esperón5 and Jose M. Sánchez-Vizcaíno1

Abstract

Background: In the last 20 years, Cetacean Morbillivirus (CeMV) has been responsible for many die-offs in marine mammals worldwide, as clearly exemplified by the three dolphin morbillivirus (DMV) epizootics of 1990–1992, 2006–2008 and 2011 that affected Mediterranean striped dolphins (Stenella coeruleoalba). Systemic infection caused by DMV in the Mediterranean has been reported only during these outbreaks.

Results: We report the infection of five striped dolphins (Stenella coeruleoalba) stranded on the Spanish Mediterranean coast of Valencia after the last DMV outbreak that ended in 2011. Animal 1 stranded in late 2011 and Animal 2 in 2012. Systemic infection affecting all tissues was found based on histopathology and positive immunohistochemical and polymerase chain reaction positive results. Animal 3 stranded in 2014; molecular and immunohistochemical detection was positive only in the central nervous system. Animals 4 and 5 stranded in 2015, and DMV antigen was found in several tissues. Partial sequences of the DMV phosphoprotein (P), nucleoprotein (N), and hemagglutinin (H) genes were identical for Animals 2, 3, 4, and 5, and were remarkably different from those in Animal 1. The P sequence from Animal 1 was identical to that of the DMV strain that caused the epizootic of 2011 in the Spanish Mediterranean. The corresponding sequence from Animals 2–5 was identical to that from a striped dolphin stranded in 2011 on the Canary Islands and to six dolphins stranded in northeastern Atlantic of the Iberian Peninsula.

Conclusions: These results suggest the existence of an endemic infection cycle among striped dolphins in the Mediterranean that may lead to occasional systemic disease presentations outside epizootic periods. This cycle involves multiple pathogenic viral strains, one of which may have originated in the Atlantic Ocean.

Keywords: Paramyxoviridae, Endemic occurrence, Dolphin morbillivirus, Stenella coeruleoalba, Mediterranean striped dolphin

Background

Cetacean morbillivirus (CeMV) is an enveloped, negative-strand RNA virus within the genus Morbillivirus and family Paramyxoviridae [1] that may cause serious respiratory, lymphoid and central nervous system (CNS) disease in susceptible cetacean species, leading to strandings and death. Three main lineages of CeMV have been described, dolphin morbillivirus (DMV) [2], porpoise morbillivirus (PMV) [1], and pilot whale morbillivirus (PWMV). Dolphin and porpoise morbilliviruses caused mass mortalities in several cetacean species, and pilot whale morbillivirus has been reported sporadically in pilot whales [3–5], although this species has been reported also to be infected by DMV [6]. In addition to these three lineages of CeMVs, three more recent strains have been identified in stranded cetaceans [7–9].

* Correspondence: crubio@oceanografic.org
1 VISAVET Center and Animal Health Department, Veterinary School, Complutense University of Madrid, Avda. Puerta del Hierro s/n, 28040 Madrid, Spain
2 Fundación Oceanográfic de la Comunitat Valenciana, C/. Eduardo Primo Ylfera (Científic) 1B, 46013 Valencia, Spain
Full list of author information is available at the end of the article

© The Author(s). 2018 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
The first recognized morbilliviral epizootic of cetaceans (actually the first reported for any marine mammal) occurred in 1987–88 on the Atlantic coast of the USA, killing an estimated 50% of the regional bottlenose dolphins (*Tursiops truncatus*) [10]. In 1990 approximately 1000 Spanish Mediterranean striped dolphins (*Stenella coeruleoalba*) were killed by a DMV epizootic [11]. At least another 100 CeMV-caused deaths of bottlenose dolphins occurred in 1993 in the Gulf of Mexico [12, 13]. Epizootic episodes occurred again in the Spanish Mediterranean in 2007 and 2011, with over 200 and 50 deaths of striped dolphins, respectively [14–17], caused by two variants of the same DMV strain (judged from viral phosphoprotein sequences).

Since 2011, CNS-restricted morbilliviral infection (MI) was reported in several animals stranded on the Italian Mediterranean coast in 2012 [18–20] and mass mortality of striped dolphins and of sperm whales due to systemic infection were recorded in Italy in 2013 and 2016, respectively [21, 22]. Although isolated cases of systemic infection (several tissues affected) and CNS-restricted infection have been reported in dolphins in the Atlantic Ocean and in Italy in a fin whale [2, 23–25], no systemic infection has been reported in dolphins in the Spanish Mediterranean outside mass outbreaks. Here we report some cases of systemic DMV infection in striped dolphins in the Spanish Mediterranean in the 2011–2016 period, exploiting sequence comparisons of DMV phosphoprotein (P) sequences to suggest potential Atlantic origins for the cases detected from 2012 till now.

**Methods**

**Animals**

Of 92 dolphins stranded on the coast of the Valencian Community of the Spanish Mediterranean that were necropsied between 2011 and 2016, only five animals, all of them striped dolphins, were DMV-positive. Of these five, animals 2, 4 and 5 stranded dead, whereas the other two animals died shortly after being found. Table 1 and Fig. 1 give the dates and locations of the strandings and relevant features of the stranded animals.

**Necropsy and tissue sampling**

Postmortem examinations were performed within 24 h after stranding. Samples of lung, kidney, pulmonary, pre-scapular and mesenteric lymph nodes, pharyngeal tonsils, urinary bladder, spleen, skin, cerebellum and cerebrum were collected. All these organs, except the mesenteric lymph node and the urinary bladder, were investigated also by RT-PCR for CeMV.

**Histopathology and immunohistochemistry**

Samples from virtually all organs and tissues were collected and fixed in 10% neutral buffered formalin, processed for histopathology using routine methods, and stained with hematoxylin and eosin (HE) according to standard laboratory procedures. Immunohistochemistry to detect morbilliviral antigen was done on cerebrum, lung, kidney, urinary bladder, stomach and intestine of animal 1; lung and cerebrum of animals 2 and 3; lung and pharyngeal tonsil of animal 4; and lung of animal 5. It was carried out on formalin-fixed, paraffin-embedded tissue (FFPE) sections, using IgG2B-isotype monoclonal antibody against Canine Distemper Virus nucleoprotein (CDV-NP; VMRD® Inc., USA), known to cross-react with CeMV [4, 16, 26]. Sections of FFPE canine brain with known CDV infection and immunopositivity were used as positive controls. For negative controls, the primary antibody was replaced with normal mouse serum [27].

**Molecular diagnosis and phylogenetic analysis**

For DMV molecular diagnoses we used RNA isolated from homogeneized tissues (Bullet Blender™ homogeneizer, Next Advance, Averill Park, NY, USA) with the NucleoSpin RNA Table 1 Animals and stranding information

| Animal # | Species             | Date and place of stranding | Age   | Comments                              |
|----------|---------------------|----------------------------|-------|---------------------------------------|
| 1        | *Stenella coeruleoalba* | July 2011 Alicante 38° 11′ 32.26" N O° 33′ 18.75" W | Adult Female | Stranded alive 3 months after outbreak end |
| 2        | *Stenella coeruleoalba* | October 2012 Alcossebre 40° 14′ 44.75" N O° 16′ 34.14" E | Juvenile Male | Dead when found                      |
| 3        | *Stenella coeruleoalba* | July 2014 Nules 39° 50′ 41.69" N O° 5′ 56.50" W | Adult Male | Alive with tremors                    |
| 4        | *Stenella coeruleoalba* | November 2015 Sueca 39° 12′ 12" N O° 18′ 40.78" W | Adult Male | Dead when found                      |
| 5        | *Stenella coeruleoalba* | December 2015 Torrevieja 37° 59′ 5.00" N O° 40′ 51.00" W | Calf Female | Dead when found                      |
II kit (from Macherey-Nagel, Duren, Germany), utilizing our recently described Universal Probe Library (UPL) reverse transcription PCR (RT-PCR) assay, which targets the gene for the viral fusion protein [28]. This real-time PCR assay was performed on the tissue samples routinely used to detect CeMV (described above in the section “tissue sampling”). Afterwards, portions of the viral genes for phosphoprotein (P) [29, 30], nucleoprotein (N) [31] and hemagglutinin (H) [4] were amplified in conventional RT-PCR assays using as templates pure RNA from DMV-positive lung samples from animals 1, 2, 4 and 5, and from a DMV-positive cerebrum sample from animal 3. RNA extracted from the brain of a DMV-positive striped dolphin stranded in 2007 on the Spanish Mediterranean Sea is shown in Fig. 1.
coast served as a positive control. As a negative control, nuclease-free water was used instead of tissue-derived template.

The PCR-amplified regions of the P, N and H genes were purified with the QIAquick PCR purification kit (QIAGEN, Hilden, Germany) and Sanger-sequenced in an ABI Prism 3730 apparatus (Applied Biosystems, Foster City, CA, USA) by a commercial sequencing service (Secugen, CIB-CSIC, Madrid, Spain). These sequences were compared using BLASTN (http://blast.ncbi.nlm.nih.gov) with all the sequences of CeMV and other morbillivirus species deposited in the GenBank. Phylogenetic analysis was carried out using MEGA 4.0 software [32]. P-distance matrices were calculated and tree topology was inferred by the neighbor-joining method based on p-distances. Topology reliability was tested by bootstrapping 2000 replicates generated with a random seed. The partial P gene sequences obtained from the five striped dolphins were aligned using ClustalX to check for differences.

Diagnostic tests for Brucella spp. and Toxoplasma gondii were carried out on DNA extracted from brain homogenates from animals 1 to 4, all of which had histopathological traits of non-suppurative encephalitis because there have been some cases of combination of these infectious agents with CeMV in Italy [33, 34]. All these animals tested negative for Brucella spp. and Toxoplasma gondii. The Brucella spp. assay was based on TaqMan Real time PCR amplification of the brucellar insertion sequence IS711 [35]. For T. gondii, nested PCR was used to target a sequence of the repetitive gene B1 of this microorganism [36].

Results
Necropsy examination
The most relevant gross findings in animal 1 were marked prescapular and pulmonary lymphadenomegaly, pharyngeal tonsil enlargement, and multifocal pulmonary atelectasis. In Animal 2, the lungs were bilaterally expanded with noticeable rib impressions on the dorsolateral surfaces and were diffusely mottled pale gray to dark red. The meninges were diffusely congested and prescapular lymph nodes were enlarged. Animal 3 had focal ventral pulmonary atelectasis and mild caudolateral emphysema. Both prescapular lymph nodes were diffusely congested. The pulmonary-associated lymph nodes were enlarged, edematous and congested. Animal 4 had external cutaneous lesions consistent with interspecific aggressive interaction, throughout the whole body. Multiple organs and cavities were hemorrhagic because of evidence of internal lesions possibly derived from this aggressive interaction. Animal 5 was poorly preserved, and the majority of organs were severely autolysed. However, there was clear evidence that the lungs were atelectatic and that the meninges were diffusely congested, being difficult to dissect.

Histologic and immunohistochemical examination
Animal 1 had diffuse interstitial pneumonia with type II pneumocyte hyperplasia and syncytial cells. The spleen, prescapular, mesenteric and pulmonary-associated lymph nodes were diffusely and moderately depleted of lymphocytes and showed scattered foci of lymphocytolysis. Animal 2 also had severe multifocal bronchointerstitial pneumonia with type II pneumocyte hyperplasia and alveolar septa thickened by infiltrates of lymphocytes, plasma cells, and reactive fibroblasts. Alveoli contained macrophages and syncytial cells mixed with edema fluid. Multiple round, acidophilic intranuclear and cytoplasmic inclusion bodies were observed in syncytial cells, macrophages and type II pneumocytes. There was also evidence of multifocal lymphoplasmacytic encephalitis, with perivascular cuffing, multifocal acute neuronal necrosis, astrocytosis, and gitter cell infiltration. There was marked lymphoid depletion in the spleen and in pulmonary lymph nodes, which also presented sinus histiocytosis and syncytia. In animal 3 there was multifocal lymphoplasmacytic and histiocytic meningoecephalitis with perivascular cuffing. It also presented lymphoplasmacytic and histiocytic bronchointerstitial pneumonia with mild fibrosis, edema and a focal granuloma, and, in the examined lymph nodes, marked diffuse lymphoid depletion with sinus edema and histiocytosis. In animal 4 mixed interstitial bronchopneumonia with intralesional syncitia and both intranuclear and cytoplasmic eosinophilic inclusion bodies were observed. There was also neutrophilic and histiocytic lymphadenitis and pharyngeal tonsillitis, with syncitia and inclusion bodies. Although in animal 5 autolysis masked cellular detail preventing histological evaluation in most tissues, moderate multifocal interstitial bronchopneumonia was evidenced.

The bronchointerstitial pneumonia, syncitia, inclusion bodies and encephalitis in these animals were indicative of morbilliviral infection, which was confirmed by immunohistochemistry for morbilliviral antigen. In Animal 1, the lung was moderately immunopositive (syncytial cells, type II pneumocytes, alveolar and interstitial macrophages, Fig. 2a), and there was intranuclear and cytoplasmic positivity in mononuclear and multinucleated giant and syncytial cells of the prescapular lymph node (Fig. 2b) and in brain oligodendroglia and astrocytes (Fig. 2c). Mild cytoplasmic immunopositivity was observed in the transitional epithelium of the urinary bladder and in kidney's cortical and medullary tubular epithelium. In animal 2 immunopositivity was observed in pulmonary syncytial cells, type II pneumocytes and macrophages (Fig. 2d and e), in cerebral neurons (Fig. 2f) and gitter cells, whereas in animal 3 neuronal cell bodies and dendritic processes were intensely immunoreactive (Fig. 2g and h), with no immunoreactivity detected in lung tissue. In animal 4 pulmonary pneumocytes and syncitia...
and histiocytes of lung (Fig. 2i and j) and pharyngeal tonsils (Fig. 2k and l) were strongly labelled. The lung from animal 5 was clearly immunopositive for morbilliviral antigen.

Molecular and phylogenetic analyses
As expected for systemic morbillivirus infection, all tissues tested from animals 1, 2, 4 and 5 were strongly positive to the fusion protein gene by RT-PCR (assayed by UPL
RT-PCR) (Table 2) and for the P, N and H genes (assayed in conventional PCR assays). In contrast, in animal 3 only CNS samples were strongly positive for mobillivirus by these same criteria, whereas all other tissues proved negative for these four genes (Table 2), indicating CNS-restricted infection in this animal.

Subsequent Sanger sequencing confirmed that all amplicons were mobilliviral sequences. The nucleotide sequences from the P gene, N gene and H gene of animal 1 corresponded to those reported in animals of the 2011 Spanish Mediterranean outbreak (P gene, GenBank JN210891; N gene MG773794; H gene, MG773792), indicating that the DMV responsible for that outbreak caused the systemic infection of animal 1. However, the P, N and H sequences derived from animals 2 to 5 differed from those of animal 1, being identical in these four animals (P gene, GenBank KC572861; N gene, GenBank MG773795; H gene, MG773793) (Figs. 3 and 4).

**Discussion**

The results of this study show that five striped dolphins stranded in the Spanish Mediterranean between 2011 and 2015 had an infectious disease presenting with variable degrees of bronchointerstitial pneumonia, non-suppurative encephalitis and multicentric lymphoid depletion. In animals 1, 2 and 4, which had good tissue preservation, the pathological observations in several organs (Fig. 2) were consistent with acute systemic morbilliviral infection, with immunohistochemical confirmation. In addition, molecular tests proved the presence of DMV in multiple tissues of these animals and of the poorly preserved animal 5 (Table 2). In this last animal, lung preservation was sufficient to also reveal typical morbilliviral pathology and immunohistochemistry (Fig. 2). These animals are the first reported cases in the Spanish Mediterranean Sea of acute systemic DMV infection between epizootic outbreaks. In contrast to the other four dolphins, in animal 3, which presented meningoencephalitis, the immunohistochemistry for the viral antigen and the molecular tests for DMV were only positive in cerebral tissue (Table 2). Although this animal also had bronchointerstitial pneumonia, the signs of chronicity for this arrangement, together with the negativity of the lung for the laboratory tests for DMV did not support active pulmonary disease due to this virus. At stranding, the DMV infection appears limited in this animal to the CNS, corresponding to the chronic encephalitic form of DMV infection [37]. Comparisons of viral sequences suggest that the same DMV strain infected animals 2–5, and that this strain differs from the one in animal 1, as well as from the ones detected in the Spanish Mediterranean outbreaks of 1990, 2007 and 2011 (Fig. 3) [2, 15, 16]. Instead, the strain infecting animals 2–5 appears to be identical (p-distance, 0.000) to a strain identified in the Atlantic Ocean in dolphins stranded in 2011–2013, one in the Canary Islands (GenBank KF695110) and six in the Northwestern coast of the Iberian Peninsula (GenBank, KP836003, KT878656, KT878657, KT878658, KT878660, KT878661). Thus, it appears that the DMV virus infecting animals 2–5 had an Atlantic origin [23, 25]. This conclusion is supported also by the observation that the P gene sequence of animals 2–5 is phylogenetically very close to the P gene sequence of the DMV responsible for the 1990 Spanish Mediterranean outbreak (GenBank AJ608288; p-distance, 0.012), known to have Atlantic origin [11]. This close relation was also found (Fig. 3) for the N gene sequence (p-distance 0.015) (GenBank AJ608288). P-distances for the H gene (1990; GenBank AJ586536) also support the Atlantic origin of the strain found in animals 2–5. Unfortunately, the lack of reporting of these sequences for Atlantic DMV isolates prevented similar comparisons with the

**Table 2** Samples analyzed and results obtained in the molecular and immunohistochemical studies of our animals

| Animal | Conventional RT-PCR for CeMV | UPL RT-PCR for CeMV | Immunohistochemistry for CDV |
|--------|-----------------------------|---------------------|------------------------------|
| 1      | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Lung, cerebrum, kidney, intestine, stomach, urinary bladder |
| 2      | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Cerebrum, lung |
| 3      | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Lung, cerebrum |
| 4      | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Lung, pharyngeal tonsils |
| 5      | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Skin, lung, prescapular lymph node, pulmonary lymph node, kidney, pharyngeal tonsils, cerebrum, cerebellum | Cerebrum, lung |

*The indicated tissues or organs were those analyzed. Italic type highlights tissues that tested negative*
viruses of the 2011–2013 strandings in the Canary Islands and the Northwestern Iberian coast.

Concerning animal 1, since it was found to contain the same DMV strain that caused the 2011 outbreak and it stranded only three months after that outbreak was declared terminated [16], it cannot be excluded that it belongs to the trailing edge of the 2011 outbreak. However, systemic cases were found in Italy in 2011 to 2013 in one fin whale and in numerous sperm whales due to DMV strains closely related to the viruses responsible for the 2007 and 2011 Spanish outbreaks [22, 24], and DMV-positive striped dolphin mass mortality also occurred in Italy in 2013 that could possibly be related to these previous Mediterranean strains [21]. Thus, as already proposed [3, 16] there appears to be endemic viral circulation within the Mediterranean, possibly within the abundant striped dolphin population, although the potential roles of whales in the transmission and maintenance of DMV strains, possible in principle given the reports of pilot whales and a Cuvier’s beaked whale (Ziphius cavirostris) infected with DMV [6, 20, 38], appear unlikely given the low density of these species in the Mediterranean. The fact that the Valencian Community was the epicenter of the previous three DMV epizootics in the Mediterranean makes conceivable that this region could harbor local DMV reservoirs, although this possibility requires further investigation, since other factors like abundance of striped dolphins in the area, and efficient and active surveillance networks may be responsible for the early detection of the epizootics in this region.

Even though most cases of DMV infection outside outbreak periods appear to be of the chronic encephalitic form [30, 39], our findings highlight the need to be
prepared to deal with possible cases of systemic infection within these inter-epizootic periods. In addition, our finding of sequences of apparent Atlantic origin in four of the animals dealt with here stress the importance of sequencing standard fragments of viral genes to make inferences that could suggest paths of virus circulation and provide epidemiological insight, even giving clues for early detection of impending outbreaks.

Comparisons of viral sequences from the three previous Spanish Mediterranean outbreaks suggest that the outbreaks were caused by closely related DMV strains [6, 16]. For example, our results indicate a close relationship between sequences linked to the Spanish outbreaks of 2007 and 2011. Sequence comparisons extending over 9.050 kb [6] suggest that the Spanish 2007 DMV strain is derived from the Spanish 1990 DMV strain. Our finding that the sequences from animal 1 are more closely related to the 2011 sequence (P gene, GenBank JN210891; N gene, GenBank MG773794; H gene, GenBank MG773792) than to the sequences from 1990 or 2007 could be consistent with a sequential lineage (1990 → 2007 → 2011 → DMV in animal 1) or with re-introduction from the Atlantic in 2007. In contrast to the case of animal 1, partial P, N and H gene sequences from animals 2, 3, 4 and 5 were identical to each other and appear to cluster separately from the sequences from the 1990, 2007 and 2011 isolates. Our finding that animal 2, which stranded in 2012, contained an identical P sequence to that of DMV identified in the Atlantic in a dolphin that stranded in 2011 [25], suggests an Atlantic origin for this DMV strain, which was subsequently found also in animals 3, 4 and 5. Consistent with this is the fact that animal 2 had acute disseminated disease, with the virus detected in the lung and the nervous system, which would be expected if the Mediterranean striped dolphin population in 2012 was naive to this Atlantic strain. The fact that we found the same strain in animal 3 that stranded in 2014 with CNS-restricted infection indicates that this strain can induce both systemic and chronic infection.

Conclusions

In summary, our results suggest that multiple DMV strains are circulating in striped dolphins and causing systemic infection in the western Mediterranean during inter-epizootic periods. At least one of these strains can produce both systemic and chronic infection and appears to have originated in the Atlantic Ocean. These findings warrant further studies aiming at clarifying how the virus circulates and causes epidemics in the Mediterranean Sea.

Abbreviations

CDV-NP: Canine Distemper Virus nucleoprotein; CeMV: Cetacean morbillivirus; CNS: Central nervous system; DMV: Dolphin morbillivirus; FFPE: Formalin-fixed, paraffin-embedded; H: Haemaglutinin; HE: Hematoxylin and eosin; MI: Morbillivirus infection; N: Nucleoprotein; P: Phosphoprotein; PMV: Porpoise morbillivirus; PWMV: Pilot whale morbillivirus; RT-PCR: Reverse transcription PCR; UPL: Universal probe library

Acknowledgments

The authors thank Belen Rivera and Rocío Sánchez for technical assistance and Prof. Vicente Rubio (IBV-CSIC, Valencia) for critical reading of the manuscript and for help with writing of the revised versions of this paper. This work was supported by a collaborative agreement involving the Conselleria de Agricultura, Medio ambiente, Cambio Climático y Desarrollo Rural de la Generalitat Valenciana, the Oceanographic Aquarium of the Ciudad de las Artes y las Ciencias of Valencia and the VISAVET Center of the Complutense University of Madrid. This work was cofinanced by Project CGL2009-08125 of the Spanish National Research Plan. Mar Melero is the recipient of a PhD student grant from the University Complutense of Madrid. Consuelo Rubio-Guerri and Josué Diaz-Delgado are recipients of FPU grants from the Spanish Ministry of Education. This research did not receive any other specific grant from funding agencies in the public, commercial, or not-for-profit sectors.
Authors’ contributions
Necropsy and sample taking were performed by CRG, JLC, MM; histopathology and immunohistochemistry were performed by JDD, ES, MA and MAJ; viral study and phylogenetic study were analyzed by CRG, ENB, MM and FE; the manuscript was prepared and critically discussed by CRG, DGP, and JMSV with the contributions of all the remaining authors. All authors have read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1 VISAVET Center and Animal Health Department, Veterinary School, Complutense University of Madrid, Avda. Puerta del Hierro s/n, 28040 Madrid, Spain.  
2 Fundación Oceanográfica de la Comunitat Valenciana, C/ Eduardo Primo Yúfera (Scientific) 1B, 46013 Valencia, Spain.  
3 Medicine and Surgery Department (Anatomic Pathology), Veterinary School, Complutense University of Madrid, 28040 Madrid, Spain.  
4 Unit of Histology and Veterinary Pathology, Institute for Animal Health, Veterinary School, University of Las Palmas de Gran Canaria, Trasmontana, s/n, 35416 Arucas (Las Palmas), Canary Islands, Spain.  
5 National Institute for Agricultural and Food Research and Technology, Ctra. de Algete a El Casar s/n, 28130 Madrid, Spain.  
6 Veterinary Services, Avanqua Oceanogràfic SL, C/ Eduardo Primo Yúfera (Scientific) 1B, 46013 Valencia, Spain.

Received: 20 April 2017 Accepted: 2 August 2018

References
1. Kennedy S. Morbillivirus infections in aquatic mammals. J Comp Pathol. 1998;119:201–25.  
2. Domingo M, Ferrer L, Pumarola M, Marco A, Plana J, Kennedy S, McAliskey M, Rima BK. Morbillivirus in dolphins. Nature. 1990;348:21.  
3. Bellière EN, Esperón F, Fernandez A, Arbelo M, Muñoz MJ, Sanchez-Vázquez JM. Phylogenetic analysis of a new Cetacean morbillivirus from a short-finned pilot whale stranded in the Canary Islands. Res Vet Sci. 2011;90:224–8.  
4. Sierra E, Fernández A, Suárez-Santana C, Xuricu A, Zucca D, Bernaldo de Quiroga Y, García-Alvárez N, de la Fuente J, Sacchini S, Andraza M, Díaz-Delgado J, Arbelo M. Morbillivirus and pilot whale deaths, Canary Islands, Spain. Emerg Infect Dis. 2016;22:740–2.  
5. Taubenberger JK, Tsai MM, Atkin TJ, Fanning TG, Krafft AE, Moeller RB, Kodsi GT, Worthy GA, Taubenberger JK. Morbilliviral epizootic in bottlenose dolphins of the Gulf of Mexico. J Virol Methods. 1996;68:283–90.  
6. Keck N, Kwiatek D, Dhemrain F, Dupraz F, Boulet H, Danes C, Leprie C, Perrin A, Godemar J, Micout L, Lieveau G. Resurgence of morbillivirus infection in Mediterranean dolphins off the French coast. Vet Rec. 2010;166:545–5.  
7. Raga JA, Banyard A, Domingo M, Corteyn M, Van Breeswijk MF, Fernandez M, Aznar FJ, Barrett T. Dolphin morbillivirus epizootic resurgence, Mediterranean Sea. Emerg Infect Dis. 2008;14:471–3.  
8. Rubio-Guerri C, Melero M, Esperon F, Belliere EN, Arbelo M, Crespo JL, Sierra E, García-Parraga D, Sanchez-Vázquez JM. Unusual striped dolphin mass mortality episode related to cetacean morbillivirus in the Spanish Mediterranean Sea. BMC Vet Res. 2013;9:105.

Published online: 24 August 2018

1. Krafft AE, Lichy JH, Lipscomb TP, Klaunberg BA, Kennedy S, Taubenberger JK. Postmortem diagnosis of morbillivirus infection in bottlenose dolphins (Tursiops truncatus) in the Atlantic and Gulf of Mexico epizootics by a polymerase chain reaction-based assay. J Wild Dis. 1995;31:10–5.

Page 9 of 10

Rubio-Guerri et al. BMC Veterinary Research (2018) 14:248
33. Profeta F, Di Francesco CE, Marsilio F, Mignone W, Di Nocera F, De Carlo E, Lucifora G, Pietroluongo G, Baffoni M, Cocumelli C, Eleni C, Terracciano G, Ferri N, Di Francesco G, Casalone C, Pautasso A, Mazzariol S, Centelleghe C, Di Guardo G. Retrospective seroepidemiological investigations against morbillivirus, Toxoplasma gondii and Brucella spp. in cetaceans stranded along the Italian coastline (1998-2014). Res Vet Sci. 2015;101:89–92.

34. Sierra E, Sánchez S, Saliki JT, Blas-Machado U, Arbelo M, Zucca D, Fernández A. Retrospective study of etiologic agents associated with nonsuppurative meningoencephalitis in stranded cetaceans in the Canary Islands. J Clin Microbiol. 2014;52:2390–7.

35. Hinic V, Brodard I, Thomann A, Cvetnic Z, Makaya PV, Frey J, Abril C. Novel identification and differentiation of Brucella melitensis, B. abortus, B. suis, B. ovis, B. canis, and B. neotomae suitable for both conventional and real-time PCR systems. J Microbiol Methods. 2008;75:375–8.

36. Montoya A, Miró G, Blanco MA, Fuentes I. Comparison of nested PCR and real-time PCR for the detection of Toxoplasma gondii in biological samples from naturally infected cats. Res Vet Sci. 2010;89:212–3.

37. Van Bressem MF, Duignan PJ, Banyard A, Barbieri M, Colegrove KM, De Guise S, Di Guardo G, Dobson A, Domingo M, Fauquier D, Fernández A, Goldstein T, Grenfell B, Groch KR, Gulland F, Jensen BA, Jepson PD, Hall A, Kuiken T, Mazzariol S, Morris SE, Nielsen O, Raga JA, Rowles TK, Saliki J, Sierra E, Stephens N, Stone B, Tomo I, Wang J, Waltzek T, Wellehan JF. Cetacean morbillivirus: current knowledge and future directions. Viruses 2014; 6: 5145–5181.

38. Centelleghe C, Beffagna G, Palmisano G, Franco G, Casalone C, Pautasso A, Giorda F, Di Nocera F, Iaccarino D, Santoro M, Di Guardo G, Mazzariol S. Dolphin morbillivirus in a Cuvier’s beaked whale (Ziphius cavirostris), Italy. Front Microbiol. 2017;8:111.

39. Domingo M, Villafranca M, Visa J, Prats N, Trudgett A, Visser I. Evidence for chronic morbillivirus infection in the Mediterranean striped dolphin (Stenella coeruleoalba). Vet Microbiol. 1995;44:229–39.