African swine fever in backyard pigs of Sabah state, East Malaysia, 2021

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INTRODUCTION

African swine fever (ASF) is a transboundary haemorrhagic viral disease that affected domestic and wild pigs of all ages. The disease is caused by African swine fever virus (ASFV) and was introduced to China in 2018 before spreading rapidly to neighbouring Asian countries. As such, putting countries free from ASF like Malaysia at risk. ASF is highly lethal with no vaccine or treatment available. In February 2021, we confirmed backyard pigs from various locations in Sabah were infected with ASF using real time polymerase chain reaction (real-time PCR). Further characterization of the Sabah ASFVs indicated that they were of p72 genotype II with intergenic region (IGR) variant II that displayed an addition tandem repeat sequence (TRS) insertion, similar to ASFV from Indonesia, Vietnam and China. These results indicate and support the transboundary expansion of a homogenotypic ASFV (p72 genotype II and IGR variant II) in the Europe and Asia-Pacific, emphasizing the need for a holistic international collaboration in control and preventing further spreading of the current ASF pandemic. Importantly, our results informed the first detection and characterization of ASF, a disease previously not detected in Malaysia. This information is crucial for further mitigation and preventive measures.

Keywords: African swine fever; Malaysia; p72; TRS; backyard pig.
analysis of few different genetic regions (Sanna et al., 2017). The B646L gene (encoding p72 major capsid protein) is a key fragment for ASFV genotyping (Bastos et al., 2003) with 24 geographically related genotypes reported so far (Ge et al., 2018). However, the abundant genotype of ASFV reflected the complexity of ASF epidemiology especially in discriminate closely related isolates. Therefore, variable tandem repeat sequences (TRS) between I73R and I329L gene was used as a tool for subtyping closely related ASFV from Eastern Europe, the current circulating strain in Europe and Asia region (Gallardo et al., 2014; Mazur-Panasiuk et al., 2020).

The ongoing uncontrolled spreads of ASF in the region have impacted the relevant industries hard and put countries free from ASF like Malaysia at risk. Unexpectedly, high sudden mortality in backyard pigs was reported in February 2021 at Sabah state, suspected for ASF. Here, we described the first detection along with the initial genetic characterization (partial p72 and TRS) of ASFV for the first outbreak in Malaysia.

MATERIALS AND METHODS

Samples
A total of 14 samples were received from the Sabah State Department of Veterinary Services (Sabah DVS) in February 2021 for the diagnosis of swine viral diseases including ASF. The samples (bone marrow, liver, kidney, spleen, lung, intestine, heart and brain) were from cases of high sudden mortality involving backyard pigs in three neighboring districts (Pitas, Kota Marudu and Beluran) within Kudat and Sandakan division of Sabah, East Malaysia (Figure 1). The pigs were reported depressed and loss of appetite prior sudden death. The samples from different districts were received in different date and were processed separately according to standard operating protocol to avoid contamination.

Deoxyribonucleic acid (DNA) extraction and real time polymerase chain reaction (real-time PCR)

The specimens were homogenized with mortar and pestle in the presence of sterile sand. The homogenate was centrifuged at 3 000 rpm for 10 minutes before the supernatant was filtered. Viral DNA extraction was done from the supernatant with the IndiSpin Pathogen Kit (Indical Bioscience, Germany) according to the manufacturer’s instructions. The extracted DNA was stored at -20°C until further analysis. The ASFV detection was carried out by following the manufacturer’s instructions of the real-time virotype ASFV PCR Kit (Indical Bioscience, Germany) which the endogenous control is included. The amplification was performed on CFX Connect Real-Time PCR System (Bio-Rad, USA). Besides that, primer-probe from King et al. (2003) as recommended by the

Figure 1. Map of Sabah State, East Malaysia shows the locations of ASF positive cases. The Sabah base map was used to map out the distribution of ASF cases according to World Geodetic System (WGS) 1984 projection with ArcGIS 10.8 by ESRI.
OIE was also used to detect the ASFV (OIE, 2021a). For the King et al. (2003) probe, BHQ1 was used as quencher instead of TAMRA. The 15 uL real-time PCR mixes contained final concentration of 0.5 pmol of both sense and antisense primer, 0.5 pmol of probe and 1x LightCycler® Taqman® Master mix (Roche, Switzerland), before 5 uL of DNA was added. The reaction was subjected to 95°C for 10 minutes for initial denaturation, followed by 45 cycles of denaturation at 95°C for 10 sec, annealing at 60°C for 35 sec and extension at 72°C for 1 sec with QuantStudio 3 System (Applied Biosystems, USA).

Gene sequencing and phylogenetic analysis
For each district, one positive ASF sample was amplified and further characterized on the partial B646L gene (encoding the p72 major capsid protein) (Bastos et al., 2003) and intergenic region (IGR) variant between I73R and I329L genes (Gallardo et al., 2014). Both of these molecular markers are used in the standard genotyping procedures (Ge et al., 2018; Le et al., 2019). The IGR between I73R and I329L genes contain tandem repeat sequence (TRS) that is important in distinguishing between closely related ASFVs. The conventional PCR was carried out with MyCycler™ (Bio-Rad, USA) using GoTaq® Green Master Mixes (Promega, USA) in a total volume of 25 uL reaction. The reactions were subjected to 95°C for 2 minutes for initial denaturation, followed by 35 cycles of denaturation at 95°C for 30 sec, annealing at 59°C for 30 sec, extension at 72°C for 30 sec and a final extension at 72°C for 5 minutes. The amplicons were subjected to gel electrophoresis and visualized on 1.5% agarose gel, stained with SYBR® Safe DNA gel stain (Invitrogen, USA). The 100 bp DNA ladder (Promega, USA) was used as the molecular weight marker. The amplicons were then excised and sent for Sanger sequencing (Apical Scientifics [M] Sdn Bhd). Raw nucleotide sequences were processed using BioEdit version 7.1.9 (Hall, 1999). The nucleotide sequences were deposited to GenBank and their accession numbers are MW788576-MW788578 (p72) and MW788579-MW788581 (TRS), respectively. Clustal W was used to align the partial p72 sequenced for Sabah ASFVs along with representative sequences for all 24 ASFV genotypes retrieved from GenBank. Phylogenetic analysis for partial p72 using neighbor-joining method with Kimura 2 parameter was run in 1 000 bootstrap replicates with Molecular Evolutionary Genetics Analysis (MEGA) version 7 (Kumar et al., 2016).

RESULTS
All 14 samples (bone marrow, liver, kidney, spleen, lung, intestine, heart and brain) of backyard pigs from three different locations in Sabah were detected positive for ASFV by real-time PCR. In contrast, these samples were negative for other swine viral diseases including classical swine fever (CSF) and porcine reproductive and respiratory syndrome (PRRS). In addition, both virotype ASFV PCR Kit and King et al. (2003) assays detected consistent threshold cycle (Ct) for all the samples. The Ct value range from 18 to 27 with ± 0.5 Ct value difference between assays.

Further amplification and sequencing of one amplified product each from Pitas, Kota Marudu and Beluran with both p72 and TRS, confirmed and characterized the first occurrence and the outbreak of ASF in Malaysia. Analysis on the partial p72 shown all three Sabah strains were 100% identical to each other and to other ASFV strains from a wide geographical location, includes Indonesia, Vietnam and China. Phylogenetic analysis using partial p72 indicated the Sabah strain belonged to p72 genotype II (Figure 2). Analysis on the TRS revealed that all three Sabah ASFVs were IGR variant II with an additional 10 bp TRS (S’-GGAAATATATA-3’) insertion between I73R and I329L gene, compared to the reference sequence of the Georgia 2007 strain (Figure 3). These results were similar to ASFV strains from Indonesia, Vietnam and China.

DISCUSSION
Clinically, ASF is indistinguishable from other porcine diseases including CSF and PRRS, making differential diagnosis challenging. Moreover, the confirmation of ASF depends on laboratory testing (OIE, 2021a). In this study, both virotype ASFV PCR and King et al. (2003) assay showed specific detection and consistent performance among each other’s. Together with the characterization markers, rapid reliable detection and confirmation of ASF are crucial for disease differential diagnosis and timely implementation of control measures in preventing disease spreading.

The p72 and IGR molecular markers have been used extensively in ASF phylogenetic and molecular epidemiological purposes. So far, there are 24 geographical related ASFV genotypes based on the p72 characterization (Ge et al., 2018). Notably, all three Sabah ASFVs were of p72 genotype II. Grouped in a monophyletic with ASFV strains from Indonesia, Vietnam, China, Estonia and Georgia, this ASFV strain is in accordance with the genotype that is responsible for the current ASF epizootic in Europe and Asia since its introduction into Georgia in 2007 (Ge et al., 2018; Le et al., 2019; Dharmayanti et al., 2021). In combination with the Asia and the Pacific ASF epidemiology, our result suggests potential spillover or introduction of this TAD from the neighboring countries concordance with the outbreak timeline.

Comparisons of TRS of Sabah strains with Indonesia, Vietnam, Korea, China, few other Eastern Europe countries and Georgia are also being carried out to explore the potential epidemiology and introduction of ASFV p72 genotype II into East Malaysia. Likewise, the Sabah strains show a homogenous nucleotide pattern with an additional 10 bp TRS (S’-GGAAATATATA-3’) insertion between I73R and I329L gene (Figure 3), which categorized them as an IGR variant II (Mazur-Panasiuk et al., 2020). This additional motif is absent in domestic pig isolate from Georgia, the older p72 genotype II isolate. Conversely, this motif is present in all recent ASFV strains isolated from domestic pigs from Asia and both domestic and wild pigs from Eastern Europe. Taken together, this information implied the current circulating strain may originate from the recent evolved strain compared to the original Georgia strain which was introduced from Africa in 2007 (Rowlands et al., 2008). Importantl, this information is in agreement with the current epidemiological events and reaffirmed the expansion of a homogenotypic ASFV from Europe to Asia-Pacific region.

All the Sabah strains were identical to each other albeit been sampled from distant locations. The three districts (Pitas, Kota Marudu and Beluran) in this study were separated by the Crocker Range with the shortest road access connecting Kota Marudu to Beluran via Pitas. The distance between each location ranges from approximately 66 km to over 150 km through rough terrain. This emphasized the Sabah ASFVs are from the same introduction and indicated the disease had spread. To date, a total of 33 outbreaks have been officially confirmed across Sabah, involving both domestic pigs and wild boars with over 1 100 domestic pigs have died or killed (OIE, 2021b).
Figure 2. Phylogenetic analysis of the partial p72 gene of Sabah ASFV strains. The evolutionary history was inferred with the Neighbor-Joining method using Kimura 2 parameter model in MEGA 7 at 1 000 bootstrap replicates. Only bootstrap values >70% are shown along the branches. Black triangle indicates causative virus from this study. Roman numerals to the right indicate p72 genotypes. GenBank accession numbers are provided for all sequences. Scale bar indicates nucleotide substitutions per site.
There are needs for further characterization of the Sabah ASFV especially on the wild boars sample which is lacking, to establishing the source of disease introduction and spread (Vergne et al., 2020). Notably, whole genome sequencing of ASFV could shed light on better understanding of the epidemiology, evolution and useful in distinguishing the circulating strains on the genome scale. Collectively, this study provides evidence and supports the common origin and transboundary expansion of a homogenotypic (p72 genotype II and IGR variant II) circulating ASFV strain in Asia-Pacific region.

The rapid transboundary transmission of this homogenotypic ASFV strain could be from a variety ways (Dixon et al., 2020; Vergne et al., 2020), particularly through infected pork products, swill feeding and loose border vigilance. However, the introduction of ASF into Sabah, East Malaysia remains to be elucidated. At present, there is no vaccine available for ASF. As such, the best strategies for infected countries like Malaysia are the implementation of stamping out, cleaning and disinfection, zoning and movement control, strengthen awareness, strict biosecurity, surveillance and rapid disease detection to contain and control the disease spreading.

The rapid transboundary expansion of ASF in this region is unprecedented. As such, holistic international collaborations encompass One Health paradigms are needed for timely response, effective implementation of control measures to mitigate the further spreading of ASF pandemic in this region and beyond.

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Conflict of interest

The authors declare that they have no conflict of interest.

REFERENCES


