Pathogenicity of novel goose-origin astrovirus causing gout in goslings
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Abstract

Background: A novel goose-origin astrovirus (GoAstV) has broken out across China in recent years, causing gout in goslings with a mortality rate of around 50%. However, our understanding of the dynamic distribution, tissue tropism and pathogenesis of GoAstV is incomplete. In order to assess its pathogenicity, one-day-old goslings were inoculated separately with GoAstV via oral and subcutaneous injection routes.

Results: Clinical symptoms, gross and microscopic lesions, blood biochemical parameters and viral loads were detected and recorded for 20 days after infection. Typical gout was observed in experimental goslings. GoAstV can be replicated in tissues and cause pathological damage, especially in the kidney, liver, heart and spleen. Virus-specific genomic RNA was detected in blood, cloacal swabs and all representative tissues, and virus shedding was detected up to 20 days after inoculation, suggesting that GoAstV has a wide tissue tropism and spread systematically after inoculation. The viral copy numbers examined in kidney were the highest, followed by spleen and liver.

Conclusion: This experiment determined the accurate value of viral loads and biochemical indicators of GoAstV-induced goslings. These findings increase our understanding of the pathogenicity of GoAstV in goslings and provide more reference for future research.

Keywords: Novel goose-origin astrovirus, Goslings, Gout, Histopathology, Biochemical parameters, Viral loads

Background

Astroviruses are non-enveloped positive-sense single-stranded RNA viruses belonging to the Astroviridae family [1]. Currently, the family was classified into two genera, Mamastroviruses and Avastroviruses [2]. Astroviruses are characterized with a size of approximately 30 nm in diameter [3]. The length of genomes ranges from 6.1 to 7.9 kb, consisting of a 5'-untranslated region (UTR), three open reading frames (ORFs), 3'-UTR and a poly (A) tail [4]. ORF 1a and ORF 1b encode non-structural protein for genomic replication, and ORF 2 encodes capsid proteins associated with the virion formation [5]. The astrovirus polymerase has no proofreading function during the replication process, so the replication of the virus is prone to error, resulting in an increase in the mutation rate of the virus [6]. The high genetic diversity and recombination potential of astroviruses indicate that cross-species transmission may be frequent [7, 8].

Since the initial discovery in faeces of infants in 1975, astroviruses have been detected in many species, including poultry [9–12]. Infection with astroviruses mainly causes acute gastroenteritis in human and animals, occasionally with encephalitis [8, 13]. In poultry, astroviruses infection has been found to be associated with a variety of diseases, such as nephritis in chicken, visceral gout in broilers and fatal hepatitis in ducklings, leading to severe economic losses [14–16]. The typical characteristics of...
astroviruses infection in poultry are high morbidity and mortality, low feed conversion rate and growth retardation [4].

Since November 2016, a new disease in goslings, of which the specific symptoms are gout, hemorrhages and renomegaly, has spread to many provinces of China, including Shandong, Jiangsu and Anhui, as well as other regions. It is estimated that the economic losses caused by the epidemic are between 1.2–1.5 billion yuan [17]. The infection rate of the disease was usually up to 80%, and the mortality rate even up to 50% [18]. The infection can be identified from the goslings aged from 5 to 20 days. Autopsies showed that severe urate deposition in the internal organs, especially the heart, liver and kidney. A novel goose-origin astrovirus (GoAstV) has been identified in infected goslings, which is different from previously reported avian astroviruses in terms of genotypes and has a nucleotide homology of less than 70% of other avian astroviruses [17, 19–21]. In the present study, we detect the dynamic distribution of GoAstV in goslings infected by oral and subcutaneous injection routes respectively, which revealed virus shedding and virus distribution, helping to further understand the pathogenicity and transmission mechanism. It provided theoretical support for the prevention and control of the disease.

Results
Clinical signs, body weight and gross lesions
The goslings in oral infection group began to present loss of appetite and depression at 5 dpi. Compared with oral infection group, those clinical signs were observed as early as 4 dpi in subcutaneous infection group. Survival curves of goslings were performed using Prism 7 program (GraphPad) (Fig. 1). Goslings in oral infection group began to die at 7 dpi and presented a mortality 30% at 13 dpi. While goslings in subcutaneous infection group began to die at 6 dpi and exhibited a mortality of 38% at 14 dpi. The survival rate showed extremely significant difference between the infected groups and the control group at the end of observation period ($P < 0.01$). Compared with the control group, goslings in the infected groups grew slowly and gained less weight. In addition, the body weights of goslings in the subcutaneous infection group were lower than those in the oral infection group (Fig. 2).

The lesions of all dead goslings in experimental groups had a high similarity. At necropsy, gout with urate deposits in visceral organs and articular cavity, severe hemorrhages and renal enlargement, and white substance-filled ureters can be observed in dead goslings (Fig. 3). No mortality or any clinical symptoms were found in control goslings in this experiment.

Histopathology in different tissues
Histopathological analysis showed that there were obvious and almost identical histopathological changes in both experimental groups at 7 dpi. And the most obvious lesions of this disease were in liver, spleen and kidney (Fig. 4). Colorless to basophilic radiating, sharp, acicular, crystalline deposits (urate tophi) are often surrounded by low to moderate numbers of viable and degenerate heterophils, small amounts of eosinophilic cellular and karyorrhectic debris (necrosis), which were frequently observed in the livers (Fig. 4a, d). The spleen showed diffuse hemorrhage (Fig. 4b, e). As showed in Fig. 4c and f, renal tubular epithelial cell degeneration, necrosis and exfoliation were prominent features in infected goslings in our experiment. In summary, no lesions were observed in the corresponding tissues of goslings in the control group, while similar lesions were observed in various tissues of goslings in both infected groups.

Detection of biochemical parameters
The detection results of three groups were presented in Fig. 5. The analysis suggested that the ALT, AST, UA and UN of infected groups were higher than that of control group. Briefly, ALT, AST and UA of subcutaneous infection group all reached the peak at 6 dpi, UN at 8

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**Fig. 1** Survival rates of goslings after inoculation with SDPY strain. The percentage of goslings that survived in the infected groups were extremely significantly lower than in the control group. **p < 0.01. **Means that the difference between infection groups and control group was extremely significant.
ALT, AST and UN of oral infection group all reached the peak at 8 dpi, UA at 6 dpi. After reaching peak levels, the above biochemical parameters declined with time.

Detection of viral load in the blood
The detection results of GoAstV copy number in blood were shown in Fig. 6. Note that the viral RNA was not detected in the control group. Viral copy numbers of both infected groups could be detected in the blood as early as 2 dpi and reached double replication peaks at 6 dpi and 10 dpi. Moreover, viral copy numbers of subcutaneous infection group were higher than that of oral infection group.

Detection of viral load in cloacal swabs
The dynamic changes of GoAstV in different tissues of goslings were detected by qRT-PCR (Fig. 8). The virus RNA can be detected throughout the experiment and attained high levels in all investigated tissues at 6 and 10 dpi. Maximum virus copy numbers were achieved at 6 dpi. Notably, more viral copy numbers were found in the kidneys than in other tissues. Furthermore, virus copy numbers in all investigated tissues showed similar trend in both infected groups. No positive viral RNA was recorded in the control goslings in this study.

Discussion
In recent years, increasing clinical cases of GoAstV infections have been concerned and are reported in different provinces of China, inducing huge economic losses for goose industry [18, 19]. Although many studies have been conducted on virus isolation and characterization of GoAstV, less is known about the pathogenicity and the replicator dynamics in vivo, which is critical for
enclosing the pathogenesis of the emerged GoAstV. So, in this study, a systematic trial of the pathogenicity of GoAstV in goslings was carried out.

As expected, this study revealed that goslings infected with GoAstV showed clear symptoms of gout. Goslings in subcutaneous infection group could have been symptomatic much more or earlier than those in oral infection group. Severe and identical lesions were observed in both infection groups, suggesting that GoAstV could infect goslings horizontally through the digestive tract.

Fig. 4 Histopathology lesions of tissues from goslings infected with GoAstV at 7 dpi. HE-stained liver section showed urate crystals and necrosis (A, D). HE-stained spleen section showed diffuse haemorrhage (B, E). HE-stained kidney section, renal tubular epithelial cell degeneration, necrosis and exfoliation (C, F). G H and I are negative control (histology changes in healthy goslings). Magnification, × 100

Fig. 5 Dynamics of ALT, AST, UA and UN in serum of the goslings post artificially infected with GoAstV. *Means that the difference between infection groups and control group was significant. **Means that the difference between infection groups and control group was extremely significant
The results laid the foundation for us to better understand the route of GoAstV transmission. GoAstV-infected goslings had severe urate deposits on the internal organs and ureters. These histopathological observations of GoAstV-induced renal tubular epithelial cell degeneration and necrosis in consistency with the previous study that the Moser reported in 2007, which provided further evidence for the impaired renal function [22]. In experimental groups, severe urate deposits may have blocked the renal tubules and caused renal damage [23], thereby reducing uric acid excretion and increasing serum uric acid. Uric acid accumulates in the blood and can be transferred to any of the organs of the body through the blood circulation. Therefore, organs with larger blood flow, such as heart, liver and kidney, have more obvious gout lesions. Furthermore, the clinical symptoms and histological lesions in goslings were largely consistent with the other latest reports [1, 19]. These results indicated that GoAstV could cause kidney damage, which maybe the main cause of gosling gout in China. Apart from gout, GoAstV infection can decrease feed intake and lower down feed conversion efficiency, which leads to growth repression. The weight loss of infected goslings is an important problem, because they lost 37% of their weight at 14 dpi, which caused considerable economic losses.

AST mainly exists in cardiomyocytes, followed by hepatocytes [24]. ALT is present in high concentration in hepatocytes, it’s the specific marker of hepatocellular injury because it occurs exclusively in the liver [25]. If hepatic cells or myocardial cells are damaged, above enzymes will leak into the blood and levels of these will increase rapidly. It is reported that markedly elevated serum UA is clearly associated with gouty arthritis and nephrolithiasis [26]. UA and UN levels can reflect tissue damages, especially for renal impairment [27]. The detection results of this study showed that almost all blood biochemical parameters of experimental groups were higher than those of control group and then decreased with time. One such factor may be tissue and cells damage caused by the invasion of virus, resulting in the leakage of enzymes into the circulation. And we know that tissue damage is able to self-repair. Therefore, the subsequent reduction in the level of enzyme is likely caused by the cellular self-repairing capability. This also supports changes in the above histopathological damage.

Related to the qRT-PCR data, viral loads could be detected in the blood, cloacal swabs and various organs collected from infected goslings at 2 dpi, indicating the primary viremia can be developed less than 48 h post infection. Viral RNA was found in all investigated tissues sampled from the infected goslings suggesting that GoAstV has a wide tissue tropism and spread systematically after inoculation, which was in accord with those in other studies [1]. GoAstV was continuously detected in infected goslings throughout the experiment indicating that the virus can replicate efficiently in vivo. The kidney may be one of the main target organs of GoAstV due to the highest virus copy number, which further indicated that GoAstV can cause gout owning to the impaired kidney. In addition, target organs may also include spleen and liver, and they could be used as options for detection of GoAstV due to high viral copy numbers. In this study, the changes of viral loads in cloacal swabs, blood and various organs were basically the same. Reasonable immunization procedures can be formulated according to the shedding virus in production practice. During the peak period of viral loads, the disinfection of goose house should be strengthened to prevent the wide spread of the disease.
Conclusions
The commercial vaccines against GoAstV have not yet been developed in China due to a lack of knowledge of the relevant diseases. Therefore, further pathogenic analysis of GoAstV is necessary and would provide useful information for the development of the efficient vaccine against gout. In conclusion, this experiment determined the accurate value of viral loads and biochemical indicators of GoAstV-induced goslings. It fills in the gaps in this aspect of data and provides more reference for future research.

Methods
Animals
One hundred fifty one-day-old healthy goslings were purchased from a commercial hatchery in Shandong Province. Three goslings from each group were randomly selected every 2 days after inoculation for the test. We only needed 30 goslings for each group, and in the actual trial we prepared 50 goslings much more than we planned. Those goslings were fed ad libitum in specific pathogen-free (SPF) isolators without any immunization. Cloacal and tracheal swabs were collected from goslings before inoculation to ensure that goslings were GoAstV-negative by TaqMan-based one-step real-time quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR) assay.

Virus
The SDPY strain (accession number: MH052598) used as the challenge virus in this study was isolated in our laboratory from the kidney of goslings with gout [17, 18]. The titer of challenge virus was determined as $10^{5.25}$ median embryo lethal dose (ELD$_{50}$)/0.2 mL by infection of goose embryos and calculation of the titer by the Reed and Muench method [28].

Pathogenicity assessment of GoAstV in goslings
To determine the pathogenicity of GoAstV, 150 one-day-old goslings were randomly divided into three groups of 50 and housed in separate SPF isolators. The goslings in the experimental groups were inoculated with 0.2 mL of GoAstV (SDPY strain; $10^{5.25}$ ELD$_{50}$/0.2 mL) either orally or by subcutaneous injection in the neck. The control group was inoculated with equal doses of phosphate buffer saline (PBS) at the same injection site. Water and food were autoclaved before feeding and automatically refilled. Clinical symptoms, gross and microscopic lesions were recorded after infection. On the 2nd, 4th, 6th, 8th, 10th, 12th, 14th, 16th, 18th, and 20th days post-inoculation (dpi), three goslings in each group were randomly selected for weighing. At the same time, serum samples were collected from all selected goslings and stored at $-20^\circ$C, which were used to detect and record the changes of biochemical parameters including alanine aminotransferase (ALT), aspartate aminotransferase (AST), urea nitrogen (UN) and uric acid (UA). The biochemical parameters were detected by Taian City Central Hospital (Taian, China). Blood specimens, cloacal swabs and tissue samples of heart, liver, spleen, lung, kidney, bursa, thymus, pancreas, brain, proventriculus and intestine were also collected at each dpi and stored at $-80^\circ$C until were using for detecting and recording the changes of viral load by qRT-PCR. At the end of this study, the remaining geese were euthanized by intravenous injection of sodium pentobarbital (100 mg/kg body weight). The preferred method of euthanasia is intravenous administration of excessive barbiturates, which can suppress the central nervous reflex and render the animal unconscious. Excessive doses of barbiturates can cause the animal to stop breathing, followed by cardiac arrest.
Histopathology
The goslings that died after inoculation were autopsied and tissue samples (liver, spleen and kidney) were collected for histopathological analysis. Tissue samples were fixed in 10% formalin for 48 h at room temperature, and then routinely processed, embedded in paraffin wax, and cut into 5-μm sections. The sections were stained with hematoxylin and eosin (HE) and examined using light microscopy.

RNA extraction and qRT-PCR
RNA was extracted by MiniBEST Universal RNA Extraction Kit (TaKaRa, Dalian, China) following the manufacturer’s instructions. The concentration of each RNA sample was measured using the DeNovix DS-11 Spectrophotometer. To determine the viral load in each sample, GoAstV was detected by qRT-PCR method previously established in our laboratory. A pair of specific primers and probe used in this study are as follows, forward: GGTGGGCTAAATAACGGAACCTCAG, reverse: GACC TATTTTCCTTGGGATCAC and probe: TCGGCTCA ACATCGGCTGA TGGG. The qRT-PCR assay was developed and validated using the LightCycler (Roche Diagnostics) and TaKaRa One Step PrimeScript™ RT-PCR Kit (TaKaRa, Dalian, China). The optimized qRT-PCR reaction volume was 20 μL containing 10 μL 2 × One Step RT-PCR Buffer III, 0.4 μL TaKaRa Ex Taq HS, 0.4 μL PrimeScript RT Enzyme MixII, 0.4 μL forward primer, 0.4 μL reverse primer, 0.4 μL probe, 6.0 μL RNase Free ddH2O, and 2.0 μL RNA template. An initial reverse transcription at 45 °C for 5 min, reverse transcriptase inactivation at 95 °C for 10 s, followed by 45 cycles at 95 °C for 5 s and at 60 °C for 20 s, and ending at 4 °C. Fluorescence signals for each sample were harvested at the end of each step at 60 °C.

Statistical analysis
Each experiment was repeated three times independently and average values were taken. No data exclusion in the manuscript. All data were presented at means ± standard deviation and analyzed using a one-way analysis of variance (ANOVA) procedure of GraphPad Prism 7.0 (GraphPad Software Inc., San Diego, CA, United States). Statistical significance was set at P < 0.05.

Abbreviations
GoAstV: Goose-origin astrovirus; SPF: Specific pathogen-free; qRT-PCR: Real-time quantitative reverse transcriptase-polymerase chain reaction; ELID50: Median embryo lethal dose; PBS: Phosphate buffer saline; dpi: Days post-inoculation; ALT: Alanine aminotransferase; AST: Aspartate aminotransferase; UN: Urea nitrogen; UA: Uric acid; HE: Hematoxylin and eosin; ANOVA: Analysis of variance

Acknowledgements
Not applicable.

Authors’ contributions
YD and YT made substantial contributions to conception and design, and had the initial idea, was partially guiding the experiments and contributed substantially to the manuscript. DY and JT performed the experiment and statistical analysis. JY made substantial contributions to conception and design, was actively involved in the planning and conduction of the experiment, and revised the manuscript critically. DY drafted the manuscript. All authors have read and approved the final version of the manuscript.

Funding
This work was supported by the National Key Research and Development Program of China (2017YFD0500803); the China Agriculture Research System (CARS-42-19); and Funds of Shandong “Double Tops” Program. These programs supported entirely this project starting from the design of study, sample collection, analysis and interpretation of data.

Availability of data and materials
The datasets used and/or analyzed in the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
All the animal infection experiments were approved by the Animal Care and Use Committee of Shandong Agricultural University and conducted in accordance with the “Guidelines for Experimental Animals” of the Ministry of Science and Technology (Beijing, China). All goslings used in the experiment were euthanized by intravenous injection of sodium pentobarbital (100 mg/kg body weight).

Consent for publication
Not applicable.

Competing interests
All authors declare that they have no conflict of interest with any organization.

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Received: 22 June 2020 Accepted: 29 December 2020

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