5-2018

Downregulation of MicroRNA Eca-Mir-128 in Seminal Exosomes and Enhanced Expression of CXCL16 in the Stallion Reproductive Tract Are Associated with Long-Term Persistence of Equine Arteritis Virus

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Repository Citation
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Notes/Citation Information

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Digital Object Identifier (DOI)
https://doi.org/10.1128/JVI.00015-18
Downregulation of MicroRNA eca-mir-128 in Seminal Exosomes and Enhanced Expression of CXCL16 in the Stallion Reproductive Tract Are Associated with Long-Term Persistence of Equine Arteritis Virus


ABSTRACT Equine arteritis virus (EAV) can establish long-term persistent infection in the reproductive tract of stallions and is shed in the semen. Previous studies showed that long-term persistence is associated with a specific allele of the CXCL16 gene (CXCL16S) and that persistent infection is maintained despite the presence of a local inflammatory and humoral and mucosal antibody responses. In this study, we demonstrated that equine seminal exosomes (SEs) are enriched in a small subset of microRNAs (miRNAs). Most importantly, we demonstrated that long-term EAV persistence is associated with the downregulation of an SE-associated miRNA (eca-mir-128) and with enhanced expression of CXCL16 in the reproductive tract, a putative target of eca-mir-128. The findings presented here suggest that SE eca-mir-128 is implicated in the regulation of the CXCL16/CXCR6 axis in the reproductive tract during long-term persistence.

IMPORTANCE Equine arteritis virus (EAV) has the ability to establish long-term persistent infection in the stallion reproductive tract and to be shed in semen, which jeopardizes its worldwide control. Currently, the molecular mechanisms of viral persistence are being unraveled, and these are essential for the development of effective therapeutics to eliminate persistent infection. Recently, it has been determined that long-term persistence is associated with a specific allele of the CXCL16 gene (CXCL16S) and is maintained despite induction of local inflammatory, humoral, and mucosal antibody responses. This study demonstrated that long-term persistence is associated with the downregulation of seminal exosome miRNA eca-mir-128 and enhanced expression of its putative target, CXCL16, in the reproductive tract. For the first time, this study suggests complex interactions between eca-mir-128 and cellular elements at the site of EAV persistence and implicates this miRNA in the regulation of the CXCL16/CXCR6 axis in the reproductive tract during long-term persistence.
Quine arteritis virus (EVA) is an economically important systemic, reproductive, and respiratory disease of equids (1–7, 12). Its causative agent, equine arteritis virus (EAV), is a positive-sense, single-stranded RNA virus that belongs to the family Arteriviridae, order Nidovirales (8). This virus can be readily transmitted via the respiratory or venereal routes by the acutely infected horse or solely through the venereal route by the carrier stallion (3, 7, 9–11). Viral infection can be either asymptomatic or associated with a wide range of clinical signs (influenza-like syndrome), including dependent edema, conjunctivitis, periorbital or supraorbital edema, respiratory distress, urticaria, and leukopenia (1–3, 7, 12–19). Additionally, EAV infection of pregnant mares can result in abortion or birth of congenitally infected foals that develop a rapidly progressive and ultimately fatal bronchointerstitial pneumonia or pneumoenteric syndrome (20). Very importantly, EAV can establish persistent infection in the reproductive tract of stallions (carrier state), resulting in continuous shedding of infectious virus in their semen (1–3, 26), which guarantees perpetuation of the virus in equine populations (1–3, 6, 7, 9–12, 21–24). The carrier state is testosterone dependent (25) and can last from several weeks or months (i.e., virus shedding in semen ≤1 year following infection [short-term carrier]) to years or even lifelong (i.e., virus shedding in semen >1 year following infection [long-term carrier]) in spite of the development of strong serum neutralizing antibody, mucosal antibody, and local inflammatory responses in the reproductive tract (3, 7, 9, 11, 21, 23, 26–28). Interestingly, EAV carrier stallions do not exhibit clinical signs of disease and exhibit no impairment of fertility (7, 11, 19, 23). To date, the immunopathogenesis of persistent EAV infection in the reproductive tract of the stallion is not completely understood, and it is under investigation in our laboratory. Recently, it has been shown that the outcome of EAV infection in the stallion is dependent on host genetic factors (28–31), precisely associated with a specific allele of the CXCL16 gene (CXCL16S), and that EAV has a specific tropism for a subset of T and B lymphocytes and stromal cells primarily in the ampullae and, to a lesser extent, in the other accessory sex glands (vesicular, prostate, and bulbourethral glands) of the persistently infected stallion (26). Moreover, EAV can persist in the male genital tract despite the presence of strong inflammatory (mediated mainly by CD8+ T lymphocytes) and EAV-specific mucosal antibody responses (26, 27). It is paramount to better understand the mechanisms of EAV persistent infection in the stallion reproductive tract in order to enable development of novel therapies for elimination of the carrier state.

Semen is a complex body fluid composed of cells (mainly spermatozoa and a smaller number of white blood cells) and seminal plasma (32–34). Seminal plasma components in equine semen, as well as in that of other species, have been previously characterized (35–40). Interestingly, in addition to the described soluble constituents, equine seminal plasma contains a diversity of extracellular vesicles, including seminal exosomes (SEs) (34, 41–46). Seminal exosomes encompass a broad group of exosomes, including prostasomes secreted by the prostatic epithelium as well as exosomes derived from other cell types within the reproductive tract (34, 41). Exosomes (including SEs) are derived from multivesicular bodies (MVBs), as opposed to microvesicles, which originate from the process of exocytosis (47). Currently, the properties and function of specific-cell derived exosomes are under extensive investigation. Previous studies have demonstrated that extracellular vesicles are immunosuppressive components in semen and are responsible for mediating the induction of tolerance to paternal antigens (34, 48, 49). Also, they have been implicated in the inhibition of lymphoproliferative responses, phagocytic activity, and natural killer (NK) cell functionality (34, 42). In addition, there is evidence that seminal plasma components may influence viral transmission and replication by either facilitating or blocking it; this property may be dependent on specific viral infections (34, 43, 50). Seminal exosomes participate in cell-cell communication by specific cargo delivery, which may include proteins (e.g.,
cytokines and growth factors), specific lipids, and coding and small noncoding RNAs with potential regulatory functions, including microRNAs (miRNAs) (34, 51). MicroRNAs are noncoding RNA molecules approximately 22 to 24 nucleotides (nt) in length that participate in posttranscriptional regulation of gene expression through specific degradation of mRNA targets or the blockade of their translation (52). Currently, extensive studies have been undertaken to elucidate the role of specific miRNAs in diverse disease processes and their potential use as biomarkers (53–58). Recently, equine miRNAs in normal tissues have been investigated (59, 60), and exosomes and exosomal miRNAs derived from serum and ovarian follicular fluid in mares have been characterized (61–63). Even though SEs have been described for the horse and their role during sperm capacitation has been studied (46, 64), their RNA cargo and function remain to be characterized. Thus, we hypothesized that in light of their potential regulatory functions, SE-associated miRNA species could play a role in the molecular pathogenesis of EAV persistence in the reproductive tract and contribute to viral immune evasion mechanisms. The primary objective of this study was to characterize the equine SE-associated miRNAs and define their potential role during EAV long-term persistent infection in the reproductive tract. We demonstrated that equine SEs are enriched in miRNAs, with potential implications in immune-mediated pathways as observed in humans. Most importantly, we have identified specific downregulation in the relative expression of SE eca-mir-128 during long-term EAV persistent infection and enhanced expression of CXCL16, a chemokine strongly implicated in EAV long-term persistence (21, 29–31), in the ampullae. Interestingly, we have determined that CXCL16 mRNA constitutes a putative target of eca-mir-128 and that consequently, this miRNA may play a critical role in the regulation of the CXCL16/CXCR6 chemokine axis in the reproductive tract of the stallion. This finding is novel and warrants further investigation to identify the specific mechanism whereby it modulates the CXCL16/CXCR6 axis in the reproductive tract of the EAV long-term carrier stallion.

RESULTS

Isolation and size characterization of equine seminal exosomes. The equine exosomes present in two equine seminal plasma samples were precipitated overnight and analyzed by transmission electron microscopy (TEM). TEM analysis demonstrated the presence of membrane-bound particles that were either isolated or in clusters. Size distribution analysis determined a mean vesicle size of 93.43 nm, with 80.8% of the precipitated exosomes having a size ranging from 21 to 120 nm, while <20% comprised larger vesicles (>120 nm) (Fig. 1A and B). The vast majority of the vesicles (67%) had a size ranging from 41 to 100 nm.

Characterization of equine seminal exosomes by Western immunoblotting and immunogold labeling. Exosome-specific markers were evaluated in SEs derived from naive (n = 10), short-term (n = 5), and long-term persistently infected (n = 5) stallions by Western immunoblotting with a panel of specific antibodies (Table 1). Specifically, SEs were characterized by the expression of the tetraspanin protein CD9 and heat shock protein 70 (HSP70) (Fig. 1C) and lack of expression of calreticulin, a calcium-binding protein resident of the endoplasmic reticulum (ER) (Fig. 1C). The expression of CD63 was variable (detected in only 2/20 SE fractions analyzed), while SEs did not express CD81, vimentin, or major histocompatibility complex class II (MHC-II) (Table 2). Furthermore, immunogold staining demonstrated that expression of CD9 is associated with the exosomal membrane (Fig. 1B [inset]). Taken together, these data unequivocally demonstrated successful isolation of SEs from equine seminal plasma and no specific differences in the SE marker expression profiles among naive, short-term, and long-term persistently infected stallions.

Seminal exosomes contain extracellular RNA cargo with a higher abundance of small RNA species. Total RNA was isolated from SEs (derived from semen samples described in Materials and Methods and which were subsequently used for quantitative real-time reverse transcription-PCR [RT-qPCR] analysis as described below [naive stallions, n = 15; short-term carrier stallions, n = 7; and long-term carrier stallions, n = 18]),
and the RNA content was estimated in each individual sample. The average RNA content was 272.5 ng, with a range between 79.8 to 714 ng. Analysis of RNA size distribution using an Agilent Bioanalyzer along with an Agilent RNA 6000 Pico kit (Agilent Technologies, Santa Clara, CA) demonstrated that small RNAs were enriched in this sample type (Fig. 2A). Subsequently, we performed a nuclease protection assay to determine if SE-associated RNA is carried within intact exosomes. The nuclease protection assay demonstrated that 62% of the SE-associated RNA was protected from combined treatment with protease (pronase) and RNase digestion. Exosome lysis by detergent treatment, followed by protease and RNase digestions, resulted in an 80% reduction of total RNA content (Fig. 2B and C).
In order to determine specific SE-associated RNA species and potential differences in their expression levels during EAV persistence, we performed miRNA sequencing on SEs derived from three stallions (L136, L139, and L140) at -30 (prechallenge) and 86 days postinfection (dpi) and from two EAV long-term persistently infected stallions (L136 and L140) at 726 dpi. The average numbers of total reads, mapped reads, mapped fractions, and reads per RNA biotype per sample type are shown in Tables 3 and 4. The large fraction of reads that did not map are largely attributed to rRNA fragments appearing as repeated sequences since the mapping algorithms (miRDeep2 [Max Delbrück Center, Berlin-Buch, Germany]) were set not to retain reads that mapped 5 times to the reference genome for analysis (frequently encountered with rRNA fragments). In the case of rRNAs and tRNAs, these represented a small fraction of the total mapped reads and constituted 3.2% of the total mapped reads in all samples analyzed. MicroRNAs were enriched in SE fractions, with an average number of reads of 2,360,732/1,076,039 (mean ± standard deviation). A group of 23 miRNAs accounted for approximately 90% of the miRNA reads, of which 15 were shared in common between SE fractions collected at prechallenge, 86 dpi, and 726 dpi (Fig. 3; see also Table S1 in the supplemental material). The top five highly enriched miRNAs in SE fractions at those three time points included eca-let-7a-2, eca-let-7a, eca-mir-21, eca-let-7c, and eca-mir-10b. The mean percentages of the total miRNA reads are shown in Table 5. Together, these top five miRNAs accounted for a mean of 63.27% of the total miRNA reads.

**Downregulation of SE-derived eca-mir-128 is associated with EAV long-term persistence in the stallion reproductive tract.** MicroRNA sequencing data analysis identified a subset of 11 miRNAs that were differentially expressed in SEs during the course of EAV infection (Fig. 4A and Table 6). Six out of 11 miRNAs (eca-let-7e, eca-mir-34b, eca-mir-128-1, eca-mir-128-2 [the last two are referred to as eca-mir-128 since they present 100% identity in their mature sequence], eca-mir-132, and eca-mir-191a) showed an overall decrease in their expression levels during the course of EAV infection, while five (eca-mir-10a, eca-mir-28, eca-mir-146a, eca-mir-197, and eca-mir-885) conversely presented an increase in their expression levels (Fig. 4A and Table 6). Relative expression analysis by RT-qPCR of a larger number of samples comprising naive (n = 15), short-term carrier (n = 7), and long-term carrier (n = 18) stallions...
confirmed a significant downregulation of eca-mir-128 (P value = 0.0011 and P value = 0.0106 for the pairwise comparisons between long-term carrier and naive or short-term carrier stallions, respectively) in the last group (Fig. 4B), while no statistically significant differences were observed between short-term carrier and naive stallions (P value = 0.9418). Subsequently, we analyzed the expression dynamics of eca-mir-128 in SEs sequentially collected during the course of EAV infection in two experimentally infected stallions (L136 and L140). While no significant differences were observed in the expres-

**TABLE 3** Mean total reads, mapped reads, and mapped fractions in SEs

<table>
<thead>
<tr>
<th>SE fraction</th>
<th>Mean total reads</th>
<th>Mean mapped reads</th>
<th>Mean mapped fraction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prechallenge</td>
<td>13,207,169 ± 1,780,999</td>
<td>1,780,999</td>
<td>1140,261</td>
</tr>
<tr>
<td>86 dpi</td>
<td>13,431,616 ± 9,565,250</td>
<td>4,948,273 ± 3,345,532</td>
<td>3345,532</td>
</tr>
<tr>
<td>726 dpi</td>
<td>14,791,891 ± 3,331,733</td>
<td>5,924,778 ± 1,664,545</td>
<td>1664,545</td>
</tr>
</tbody>
</table>
sion of eca-mir-128 between prechallenge and 380 dpi, a significant downregulation of this miRNA was observed after 380 dpi (approximately 1 year postinfection; \( P \) values < 0.05). Furthermore, eca-mir-128 reached its lowest expression value by 726 dpi (Fig. 4C). No statistically significant differences were found for the other differentially expressed

<table>
<thead>
<tr>
<th>SE fraction</th>
<th>Mean miRNA reads</th>
<th>Mean tRNA reads</th>
<th>Mean rRNA reads</th>
<th>Mean mRNA reads</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prechallenge</td>
<td>2,744,802 ± 1,273,296</td>
<td>13,156 ± 8,639</td>
<td>26,181 ± 18,681</td>
<td>3,424,562 ± 1,405,618</td>
</tr>
<tr>
<td>86 dpi</td>
<td>2,133,247 ± 1,298,735</td>
<td>13,658 ± 9,556</td>
<td>10,050 ± 13,679</td>
<td>2,997,966 ± 2,043,063</td>
</tr>
<tr>
<td>726 dpi</td>
<td>2,125,853 ± 863,695</td>
<td>52,233 ± 12,358</td>
<td>127,762 ± 47,230</td>
<td>3,239,549 ± 1,077,548</td>
</tr>
</tbody>
</table>

**TABLE 4** Mean total reads per RNA biotype identified in SE fractions

**FIG 3** MicroRNAs that account for >90% of the total miRNA reads in SEs derived from naive (prechallenge) stallions and following equine arteritis virus infection. (A) Prechallenge; (B) 86 days postinfection; (C) 726 days postinfection. Mean percentages of expression ± standard errors of the means along with the cumulative expression are represented. (D) A total of 15/23 miRNAs were commonly shared across time points.
miRNAs identified by miRNA sequencing between naive, short-term carrier, and long-term carrier stallions.

**CXCL16 constitutes a putative target of eca-mir-128.** Computational target prediction yielded a total of 805 putative target genes for eca-mir-128 (Table S2 in the supplemental material). Functional annotation analysis of target genes demonstrated their involvement in several biological processes, with a predominance of genes regulating metabolic and cellular processes but also involved in developmental, immune system, and reproduction processes, locomotion, adhesion, localization, and response to stimuli (Fig. 5A). Specifically, target genes were involved in regulation of gene expression, cell differentiation, development, and morphogenesis, protein phosphorylation and diverse signaling pathways, including the mitogen-activated protein kinase (MAPK) cascade, the phosphoinositide-3 kinase/protein kinase B (PI3K-Akt) signaling pathway, transforming growth factor (TGF) signaling, platelet-derived growth factor (PDGF) and epidermal growth factor (EGF) signaling, Hippo signaling, serine/threonine kinase signaling, DNA damage repair systems, and cell migration/chemotaxis (Table S3 in the supplemental material). Based on their molecular function and protein class, target genes were mainly involved in RNA/DNA/protein/metal binding, catalytic activity, transporter activity, receptor activity, structural activity, and signal transduction (Fig. 5B and C; see also Table S4 in the supplemental material). To analyze the functional relationships between eca-mir-128 target genes, we used PANTHER and DAVID bioinformatics tools. Pathway analysis performed by PANTHER retrieved 20 top pathways (P values < 0.05) potentially under the regulation of mir-128, and these are depicted in Table 7 and Fig. 6A. Among these, six were associated with immune pathways (Table 7). Pathway analysis demonstrated that mir-128 target genes may also be involved in inflammation mediated by chemokine/cytokine signaling, B and T cell activation, interleukin (IL) and TGF-β signaling, and apoptosis.

Interestingly, prediction tools identified CXCL16 as a putative target of mir-128. Consequently, we predictively evaluated the miRNA-mRNA interaction between the mir-128 seed sequence and the 3′ untranslated region (UTR) of its putative mRNA target, CXCL16. For this purpose, we analyzed the interaction between both the human and equine CXCL16 3′ UTR and mir-128, respectively. Analysis of the human CXCL16 3′ UTR by TargetScan predicted that the pairing region is located within nucleotides (nt) 664 and 670 of the 3′ UTR (ACUGUGA) and that the hsa-mir-128-3p seed sequence is located between nt 1 and 7 (Fig. 6B). Prediction of the pairing region is consistent with a 7-mer-A1 site type, as it is a perfect match for the seed sequence of the mature miRNA and it is followed by an adenine. The miRNA-mRNA interaction between the human CXCL16 3′ UTR sequence and mir-128 was determined to have a low free energy (ΔG = −19.2 kcal/mol). In the case of the equine CXCL16 3′ UTR, miRNA-mRNA interaction analysis demonstrated that the mir-128 (nt 1 to 7) predicted pairing region comprises nt 37 to 43 of the 3′ UTR (AUUGUGA) and that interaction occurs with a low free energy as well, estimated at −19.3 kcal/mol. Additionally, a guanine-uracil wobble occurs between nt 6 of mir-128 and nt 38 of the equine CXCL16 3′ UTR (Fig. 6B).

**CXCL16 is upregulated in the ampullae of long-term EAV persistently infected stallions.** In order to establish an association between eca-mir-128 and its putative target, CXCL16, we evaluated the expression of the latter in the main site of EAV infection.

### Table 5: Expression of the top 5 highly enriched miRNAs in SEs

<table>
<thead>
<tr>
<th>miRNA</th>
<th>Mean expression (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>eca-let-7a-2</td>
<td>16.54 ± 3.88</td>
</tr>
<tr>
<td>eca-let-7a</td>
<td>16.51 ± 3.87</td>
</tr>
<tr>
<td>eca-mir-21</td>
<td>11.58 ± 0.17</td>
</tr>
<tr>
<td>eca-let-7c</td>
<td>10.96 ± 0.32</td>
</tr>
<tr>
<td>eca-mir-10b</td>
<td>7.67 ± 3.25</td>
</tr>
</tbody>
</table>

*aThe mean percentages of total miRNA reads ± standard deviation are shown.*
persistence (i.e., ampullae) in the reproductive tract of the stallion (Fig. 7). CXCL16-specific TaqMan qPCR demonstrated a significant upregulation of CXCL16 in the ampullae of both short-term (n = 6) and long-term persistently infected (n = 3) stallions compared to the naive group (n = 4; P values of 0.0279 and 0.0020, respectively; Fig. 7E). RNAscope in situ hybridization (ISH) and immunohistochemistry (IHC) were used to localize the expression of CXCL16 in the ampullae. CXCL16 expression was
significantly abundant in the mucosal epithelial cells of the ampullae, primarily the luminal mucosal epithelium, as well as mucosal and submucosal lymphocyte and plasma cell infiltrates in long-term persistently infected stallions compared to short-term carrier and naive stallions (P values/H110210.05 [Fig. 7A to C and 7E]), in which a lower expression of CXCL16 was observed in scattered epithelial cells (Fig. 7B and C). Furthermore, a significant negative correlation was observed between CXCL16 and SE eca-mir-128 expression levels (H9267/H11005/H110020.73; P value/H110050.0065 [Fig. 7E]).

**DISCUSSION**

The pathogenesis of persistent EAV infection in the male reproductive tract has been the subject of extensive investigation in recent years in our laboratory (26–28, 30, 31). We have demonstrated that long-term persistent infection in the stallion is associated with host genetic factors, specifically with the presence of a specific allele encoding the chemokine CXCL16 (CXCL16S) (30, 31). Furthermore, EAV has the ability to persist despite the induction of a strong systemic immune response and local inflammatory and mucosal antibody responses at the site of persistence (26, 27). Clearly, EAV employs complex strategies to evade host immunity, and the immunopathogenesis of viral persistence remains to be further elucidated (Fig. 8).

Since seminal plasma components have proven to have immunomodulatory effects that are necessary for fertility and may also influence the establishment of sexually transmitted diseases (34, 42, 48, 65–68), their role in enabling EAV persistence in the stallion warrants further investigation. In this study, we have specifically characterized the role of SE-associated miRNAs during EAV persistent infection since specific small RNA cargos within these extracellular vesicles can play a role in modulating local immunity, as recently suggested (34). Thus, we hypothesized that specific equine SE-associated miRNAs may modulate the local inflammatory response and, consequently, contribute to the establishment and maintenance of persistent EAV infection in the reproductive tract of the stallion. In this study, we sought to characterize the mature miRNA profile associated with SEs and to identify the specific miRNAs involved in the maintenance of EAV-persistent infection in the ampullae of persistently infected stallions.

Purified SEs were within the expected size range and were characterized by high expression of CD9, a ubiquitous exosome marker particularly enriched in prostasomes (34, 41, 64). In contrast to human SEs, we observed inconsistent expression of CD63 in equine SE fractions. Since SEs are derived from a number of tissues associated with the male reproductive tract, determining their specific origin constitutes a significant challenge. However, the lack of vimentin expression in SEs may suggest that these

<table>
<thead>
<tr>
<th>miRNA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>eca-let-7e</td>
<td>0.033075 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td>eca-mir-10a</td>
<td>0.03195 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td>eca-mir-28</td>
<td>0.0312 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td><strong>eca-mir-34b</strong></td>
<td><strong>0.0208 (prechallenge vs 86 dpi)</strong></td>
</tr>
<tr>
<td></td>
<td><strong>0.006 (prechallenge vs 726 dpi)</strong></td>
</tr>
<tr>
<td><strong>eca-mir-128-1</strong></td>
<td><strong>0.0045 (prechallenge vs 726 dpi)</strong></td>
</tr>
<tr>
<td></td>
<td><strong>0.0033 (86 dpi vs 726 dpi)</strong></td>
</tr>
<tr>
<td><strong>eca-mir-128-2</strong></td>
<td><strong>0.0051 (prechallenge vs 726 dpi)</strong></td>
</tr>
<tr>
<td></td>
<td><strong>0.00345 (86 dpi vs 726 dpi)</strong></td>
</tr>
<tr>
<td>eca-mir-132</td>
<td>0.02865 (prechallenge vs 86 dpi)</td>
</tr>
<tr>
<td>eca-mir-146a</td>
<td>0.0267 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td>eca-mir-191</td>
<td>0.01575 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td>eca-mir-197</td>
<td>0.0402 (86 dpi vs 726 dpi)</td>
</tr>
<tr>
<td>eca-mir-885</td>
<td>0.02145 (86 dpi vs 726 dpi)</td>
</tr>
</tbody>
</table>

*P values are represented along with the time points that show significant differences, in parentheses. miRNAs that showed a significant downregulation associated with long-term EAV persistence are shown in bold. P values were subjected to a Benjamini-Hochberg correction and a false discovery rate (FDR) set at 0.05.
FIG 5 Functional annotation analysis of mir-128 target genes using PANTHER. A total of 805 putative target genes were classified according to their biological process (A), molecular function (B), and protein class (C).
extracellular vesicles are specifically shed by the lining epithelium of the reproductive tract rather than by cells of mesenchymal origin. Furthermore, Bioanalyzer profiling along with resistance to nuclease activity demonstrated that SEs are enriched in small RNAs and that their RNA repertoire mainly included mature miRNAs as well as other noncoding RNAs. It is noteworthy that when mature miRNA species predominant in equine SEs were analyzed by miRNA sequencing, we found that a small subset \( n = 11005 \) accounted for the vast majority of the miRNAs that were expressed, of which 15 were common to all the analyzed samples independently of infection status or interval after EAV infection. The top five miRNAs expressed in SEs included members of the let-7 family, mir-21 and mir10b, which have been demonstrated to play an active role in the modulation of several immune pathways, including modulation of T cell activation, NK cell function, IL-10 production by CD4\(^+\) T lymphocytes, CD8\(^+\) effector T lymphocyte differentiation, and NF-κB pathway, among others (69–74). From a confirmative viewpoint, these findings are similar to those reported for human SEs (34), suggesting that these miRNAs can be involved in similar modulatory processes in both species. Furthermore, the let-7 family of miRNAs as well as mir-21 are highly abundant in equine serum and plasma and also expressed in diverse tissue types (59, 75–80).

Validation of the miRNA sequencing data by RT-qPCR confirmed specific downregulation of eca-mir-128 in long-term persistently infected stallions, an intronic miRNA encoded by two distinct genes, eca-mir-128-1 and eca-mir-128-2, located on Equus caballus chromosome 16 (chr16:49087163–49087246 [minus strand], corresponding to intron 16 or 18 of the cyclic AMP-regulated phosphoprotein 21-kDa gene [ARPP21], depending on the splice variant) and chromosome 18 (chr18:19492350–19492419 [plus strand], corresponding to intron 22 of the R3H domain containing 1 gene [R3HDM1]), respectively. Similar to their human counterpart (hsa-mir-128), the mature sequences of both eca-mir-128-1 and eca-mir-128-2 are identical and conserved among species (81). While hsa-mir-128 has been extensively implicated in tumorigenesis (81), its role during viral infections and inflammatory processes has been less described (82–85). Recently, downregulation of eca-mir-128 has been implicated in

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**TABLE 7** Top 20 significant pathways associated with eca-mir-128 target genes retrieved by PANTHER following the statistical overrepresentation test

<table>
<thead>
<tr>
<th>PANTHER pathway</th>
<th>No. of targets</th>
<th>Fold enrichment</th>
<th>( P ) value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDGF signaling pathway (P00047)</td>
<td>15</td>
<td>2.63</td>
<td>8.26E-04</td>
</tr>
<tr>
<td>Gonadotropin-releasing hormone receptor pathway (P06664)</td>
<td>19</td>
<td>2.1</td>
<td>2.41E-03</td>
</tr>
<tr>
<td>CCKR signaling map (P06959)</td>
<td>15</td>
<td>2.26</td>
<td>3.40E-03</td>
</tr>
<tr>
<td>PI3 kinase pathway (P00048)</td>
<td>7</td>
<td>3.38</td>
<td>5.36E-03</td>
</tr>
<tr>
<td>Interferon gamma signaling pathway (P00035)</td>
<td>5</td>
<td>4.35</td>
<td>6.47E-03</td>
</tr>
<tr>
<td>Endothelin signaling pathway (P00019)</td>
<td>9</td>
<td>2.73</td>
<td>6.74E-03</td>
</tr>
<tr>
<td>Angiogenesis (P00005)</td>
<td>14</td>
<td>2.1</td>
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equine asthma and hypothetically related to the stimulation of allergic immune responses (82). In this study, we specifically determined that downregulation of SE eca-mir-128 is associated with long-term EAV persistence in the reproductive tract and that this specific miRNA is a putative regulator of CXCL16 as determined by target prediction analysis. CXCL16, which belongs to the CXC chemokine family, acts as one of the cellular receptors of EAV and has been strongly implicated in EAV persistence in the reproductive tract (21, 30, 31, 86). The 3′ UTR and miRNA-mRNA interaction analysis predicted that the binding between the mir-128 seed sequence and its pairing region in the 3′ UTR of CXCL16 mRNA occurs with a low free energy (ΔG of ~19.2 and ~19.3 kcal/mol for the human and equine counterparts, respectively).

![A] Pathway analysis of mir-128 target genes (n = 805) and comparative analysis of CXCL16 3’ UTR as a putative target of mir-128. (A) Pathway analysis by PANTHER retrieved a total of 20 significant pathways, which included pathways in common with the CXCL16/CXCR6 axis, such as PI3K-Akt, G-protein coupled receptor signaling, and some immune pathways. (B) Putative interaction between mir-128 and human and equine CXCL16 3′ UTR. The seed region of mir-128 and binding sites at the 3′ UTRs are in bold. The binding site is predicted as a 7-mer-A1 type, and in the horse sequence, a guanine-uracil wobble (marked in red) is predicted at nt 38 of the 3′ UTR. Both interactions occur with a low free energy (ΔG of ~19.2 and ~19.3 kcal/mol for the human and equine counterparts, respectively).

**FIG 6** Pathway analysis of mir-128 target genes (n = 805) and comparative analysis of CXCL16 3’ UTR as a putative target of mir-128. (A) Pathway analysis by PANTHER retrieved a total of 20 significant pathways, which included pathways in common with the CXCL16/CXCR6 axis, such as PI3K-Akt, G-protein coupled receptor signaling, and some immune pathways. (B) Putative interaction between mir-128 and human and equine CXCL16 3′ UTR. The seed region of mir-128 and binding sites at the 3′ UTRs are in bold. The binding site is predicted as a 7-mer-A1 type, and in the horse sequence, a guanine-uracil wobble (marked in red) is predicted at nt 38 of the 3′ UTR. Both interactions occur with a low free energy (ΔG of ~19.2 and ~19.3 kcal/mol for the human and equine counterparts, respectively).
FIG 7 CXCL16 expression in the ampullae of EAV long-term carrier (n = 3), short-term carrier (n = 6), and naive (n = 4) stallions. (A to C) CXCL16 is significantly upregulated in the ampullae of EAV long-term persistently infected stallions (A) compared to short-term carrier (B) and naive (C) stallions as determined by ISH and IHC (insets). CXCL16 is predominantly expressed in the mucosal epithelium and lymphocytic infiltrates, especially in the luminal area. CXCL16-specific mRNA RNAscope ISH was performed with fast red; images are at a magnification of ×200. Bar = 250 μm. (Insets) An equine CXCL16-specific rabbit anti-peptide antibody was used for IHC. 3,3'-Diaminobenzidine tetrahydrochloride (DAB) stain was used; the image is at a magnification of ×400. (D) High expression of CXCL16 in equine endothelial cell pellets as determined by CXCL16-specific RNAscope ISH and IHC (inset, DAB, ×400). Fast red was used; the image is at a magnification of ×200. Bar = 250 μm. (E) Higher expression of CXCL16 in the ampullae of long-term carrier stallions (n = 3) was quantitatively determined by (top to bottom) TaqMan RT-qPCR, RNAscope ISH, and IHC, and it was demonstrated to be negatively correlated with eca-mir-128 expression. Blue dots represent values for individual stallions, while mean $2^{-\Delta CT}$ values ± standard errors of the means and median scores are represented in red. Letters indicate statistically significant differences between groups (P value < 0.05). For correlation analysis (E, bottom) between CXCL16 and eca-mir-128 expression, mean $2^{-\Delta CT}$ values ± standard errors of the means are depicted.
Differences in CXCL16 expression levels between the three groups along with their negative correlation with eca-mir-128 expression levels demonstrated that long-term viral persistence induces the upregulation of this chemokine in the ampullae, providing further evidence of the key role that CXCL16 plays in the pathogenesis of EAV long-term persistent infection and indirectly suggesting that eca-mir-128 may regulate the expression of this gene in the reproductive tract and allow the establishment of viral persistence (Fig. 8). In this respect, it is possible that the delivery of specific miRNA cargos to epithelial and/or inflammatory cells in the reproductive tract mediated by SEs modulates specific target genes in specific cell populations as shown for other exosomes and, consequently, influences the local immune response (34, 91). It is likely therefore that eca-mir-128 dynamics in SEs influence the outcome of EAV infection in the reproductive tract. However, further *in vitro* experiments are required to confirm that equine CXCL16 is a specific target for eca-mir-128. Additionally, eca-mir-128 is potentially involved in the regulation of several other pathways, some of which are associated with immune mechanisms (82, 83); thus, its role in local immune evasion mechanisms that allow EAV to persist deserves further investigation. The differential expression of the remaining set of miRNAs ($n = 9$) could not be confirmed, and the differences identified by miRNA sequencing could have been overestimated due to the very limited number of samples evaluated (two to three per time point) as well as the inclusion of diverse sequencing biases that can lead to enhancement or reduction of miRNA expression levels as previously reported (92).

Another significant finding from this study is that CXCL16 expression was not limited to inflammatory cells infiltrating the ampullae of long-term carrier stallions but was also abundant in the mucosal epithelium. Even though the mucosal epithelium expresses high levels of CXCL16, recent studies demonstrated that virus is not harbored in the
epithelial cells of the ampullae or other accessory sex glands (26). This observation is supportive of the fact that ongoing studies in our laboratory suggest that EAV requires additional cellular entry factors, among which vimentin seems to play an important role (S. Sarkar and U. B. R. Balasuriya, unpublished data). Thus, the lack of vimentin expression in the mucosal epithelium may be a determinant for the lack of susceptibility of CXCL16+ epithelial cells in the ampullae to EAV infection (26).

In conclusion, the study presented here provides insight into the miRNA species associated with SEs in the stallion and, most importantly, identified the specific downregulation of SE eca-mir-128 during long-term EAV persistent infection in the reproductive tract, a miRNA that predictively targets CXCL16. This study has also unequivocally demonstrated that upregulation of CXCL16 in the reproductive tract is associated with the establishment of long-term viral persistence, providing a strong link between the genetic studies undertaken and the pathogenesis of persistent EAV infection. While the evidence provided by this study indirectly suggests that SE eca-mir-128 modulates the expression of CXCL16 in the reproductive tract, further studies to confirm these observations are warranted (Fig. 8). In addition, other ongoing studies in our laboratory strongly suggest that EAV exploits the CXCL16/CXCR6 chemokine axis in order to modulate local inflammatory and immune responses at the site of persistent infection and that this chemokine axis is probably involved in specific homing of infected lymphocytes into the reproductive tract (M. Carossino and U. B. R. Balasuriya, unpublished data). Interestingly, it is yet to be uncovered how other eca-mir-128 regulated pathways contribute to modulating the CXCL16/CXCR6 axis favoring long-term EAV persistent infection in the stallion reproductive tract. In summary, this study advances our current understanding of EAV persistent infection and demonstrates that diverse and complex host factors are involved in the modulation of viral persistence in the stallion reproductive tract.

MATERIALS AND METHODS

Semen samples. Collection of equine semen samples from stallions that were experimentally infected with EAV (n = 7) has been described elsewhere (19, 26, 27). In addition, archived semen samples from long-term EAV carrier stallions that were naturally infected with EAV were included in the study. These semen samples had been submitted to the OIE Reference Laboratory at the Maxwell H. Gluck Equine Research Center for monitoring EAV carrier state following natural EAV infection in the field. A total of 53 semen samples derived from naive stallions (n = 16), short-term carrier stallions (n = 7), and long-term carrier stallions (n = 30, which include sequential semen samples from two long-term persistently infected stallions [726 dpl to evaluate eca-mir-128 dynamics]) were included in the study. For short-term carrier stallions (duration of viral shedding < 1 year), semen samples collected 1 year following viral clearance were used. In the case of long-term carrier stallions (duration of viral shedding ≥ 1 year), semen samples collected 2 years postinfection were used. All the semen samples were stored in 1-ml aliquots at −80°C.

Animal care and ethics. This study was performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Research Council (93). The Institutional Animal Care and Use Committee (IACUC) at the University of Kentucky, Lexington, KY, approved this protocol (number 2011-0888). Stallions were humanely euthanized by pentobarbital overdose following the American Veterinary Medical Association (AVMA) guidelines for the euthanasia of animals, and all efforts were made to minimize suffering.

Tissue samples. Tissue samples (ampullae) derived from a group of naive stallions (n = 4), EAV experimentally infected stallions (n = 8), and an EAV naturally infected stallion (n = 1) were used in this study (26). These were aseptically collected during postmortem examination and archived at −80°C or formalin fixed and paraffin embedded (FFPE) as previously described (26).

Isolation of seminal exosomes. Before SE isolation, semen samples were thawed on ice and centrifuged at 3,000 × g for 15 min at 4°C to remove cellular debris. The seminal plasma was subsequently filtered through a 0.8-μm filter (EMD Millipore, Billerca, MA). Seminal exosome isolation was carried out using ExoQuick exosome precipitation solution (System Biosciences, Palo Alto, CA). Briefly, 126 μl (63 μl per 250 μl of biofluid as recommended by the manufacturer) of the ExoQuick exosome precipitation solution was added to 500 μl of filtered seminal plasma, mixed, and incubated overnight (minimum of 12 h) at 4°C to allow SE precipitation. Finally, the mixture was centrifuged at 1,500 × g for 30 min at 4°C, the supernatant was removed, and the exosome pellet was either used for downstream applications or stored at −80°C.

Antibodies. A panel of monoclonal and polyclonal antibodies specific for various exosome markers was used for characterization of SEs by TEM/immunogold labeling and Western immunoblotting analysis (Table 1). An equine CXCL16-specific rabbit anti-peptide antibody (86) was used for immunohistochemical (IHC) staining. Secondary antibodies conjugated to horseradish peroxidase (HRP; goat anti-mouse...
IgG [H+L]-HRP and goat anti-rabbit IgG [H+L]-HRP [SouthernBiotech, Birmingham, AL] were used for Western immunoblotting. A secondary antibody conjugated to 12-nm colloidal gold (goat anti-mouse IgG [H+L] [Jackson ImmunoResearch Laboratories, Inc., West Grove, PA]) was used for immunogold labeling. The Bond Polymer Refine Detection kit (Leica Biosystems, Buffalo Grove, IL) was used for IHC.

Transmission electron microscopy and size determination of seminal exosomes. SEs isolated from prechallenged semen samples collected from two naive stallions were used for TEM and immunogold labeling (see below). Briefly, purified SE pellets obtained using the ExoQuick exosome precipitation solution were resuspended in 1/10 of the original volume (50 μl) of phosphate-buffered saline (PBS; pH 7.4; Gibco, Carlsbad, CA) and fixed at a 1:1 ratio in 4% paraformaldehyde for 30 min at room temperature. Subsequently, a Formvar-carbon coated 400-mesh copper grid was floated on a 40-μl drop of the SE fractions for 15 min at room temperature to allow SE adsorption. Excess sample was drained off, and the grids were washed on a drop of Tris-buffered saline (TBS; pH 7.4) for 10 min at room temperature, followed by rinsing once in deionized water. Grids were finally contrasted on a drop of 3% aqueous phosphotungstic acid (pH 7.0) for 2 min, excess stain was removed using filter paper, and grids were observed under a JEOL JEM-1011 transmission electron microscope (University of Georgia Electron Microscopy Services, College of Veterinary Medicine, Athens, GA). Digital images were acquired with an XR80M wide-angle multidiscipline mid-mount charge-coupled-device (CCD) camera (Advanced Microscopy Techniques Corp., Woburn, MA). SE size distribution was assessed by taking measurements using the TEM imaging software.

Immunogold labeling. Purified SE pellets obtained using the ExoQuick exosome precipitation solution were resuspended and fixed as described above. Subsequently, a Formvar-carbon coated 400-mesh nickel grid was floated on a 40-μl drop of the SE fractions for 15 min at room temperature to allow SE adsorption. Excess sample was drained off, and the grids were washed on a drop of TBS (pH 7.4) for 10 min at room temperature and subsequently blocked in TBS containing 2% bovine serum albumin (BSA) for 30 min at room temperature. The grids were incubated with mouse anti-CD9 (1:50 dilution; Table 1) for 1 h at room temperature using a spotting dish in a wet chamber. Excess primary antibody was extensively washed using blocking buffer, followed by secondary antibody incubation (goat anti-mouse IgG conjugated with 12-nm colloidal gold; 1:20 dilution) for 30 min at room temperature using a spotting dish in a wet chamber. Grids were extensively washed in blocking buffer and finally rinsed in deionized water. Grids were contrasted with 3% aqueous phosphotungstic acid (pH 7.0) and visualized as indicated above.

Western immunoblotting. Seminal plasma samples from naive (n = 10), short-term carrier (n = 5), and long-term carrier (n = 5) stallions were evaluated by Western immunoblotting. Purified SE pellets isolated using the precipitation solution were resuspended in 100 μl of radioimmunoprecipitation assay (RIPA) buffer (Santa Cruz Biotechnology, Dallas, TX) containing Halt protease inhibitor cocktail (Thermo Fisher Scientific, Waltham, MA) and vortexed for 15 s, and incubated for 5 min at room temperature for complete lysis. The protein concentration of the lysates was determined using a spectrophotometer (optical density [OD] at 280 nm; NanoDrop 2000 spectrophotometer; Thermo Fisher Scientific). A total of 30 μg of total protein was resuspended in 2× Laemmli sample buffer (Bio-Rad, Hercules, CA) containing 2-mercaptoethanol (Sigma-Aldrich, St. Louis, MO; except for CD9 detection), boiled at 95°C for 5 min, and subjected to denaturing polyacrylamide gel electrophoresis (SDS-PAGE) on a 12% resolving and 4% stacking gel. Subsequently, SE proteins were wet transferred onto a polyvinylidene difluoride (PVDF) membrane (Bio-Rad) for 1 h at 4°C. Transferred PVDF membranes were briefly washed in TBS (pH 7.6) containing 0.05% Tween 20 (TBS-T; Bio-Rad) and blocked with 5% skim milk in TBS-T for 1 h at room temperature. Membranes were incubated with the primary antibody (diluted at a concentration of 1 μg/ml in blocking buffer) overnight at 4°C with gentle rocking and washed 3 times for 10 min each time with TBS-T. The membranes were incubated with the respective secondary antibody (goat anti-mouse or goat anti-rabbit conjugated to HRP; diluted 1:5,000 in blocking buffer) for 1 h at room temperature with gentle rocking. After subsequent washes, membranes were developed using SuperSignal West Pico chemiluminescent substrate (Thermo Fisher Scientific) as indicated by the manufacturer. Developed membranes were imaged using an Azure c300 digital imager system (Azure Biosystems, Dublin, CA). Specific positive controls were run in parallel for each primary antibody.

RNA isolation. Exosomal RNA was isolated from purified SE pellets using the SeraMir exosome RNA isolation kit (System Biosciences) according to the manufacturer’s instructions. RNA yield was determined by fluorometry (Qubit RNA HS assay kit; Thermo Fisher Scientific), and size distribution was assessed using an Agilent RNA 6000 Pico kit (Agilent Technologies, Inc.) according to the manufacturer’s instructions. Additionally, total RNA was isolated from the ampullae for RT-qPCR. Briefly, 50 mg of tissue was dissected, placed in 1 ml of QIAzol lysis reagent (Qiagen, Valencia, CA), and homogenized using MagNa Lyser green beads (Roche, Indianapolis, IN) in a Bead Ruptor 12 (Omni International, Inc., Kennesaw, GA). Lysates were incubated for 5 min at room temperature, followed by addition of 200 μl of chloroform (Sigma-Aldrich). The upper aqueous phase was transferred into a new tube following centrifugation (12,000 × g for 15 min at 4°C), and 1 volume of 70% ethanol was added. RNA isolation was finally performed using the RNeasy minikit (Qiagen) according to the manufacturer’s recommendations.

Nuclease (RNase) protection assay. Purified SE pellets derived from 1 ml of seminal plasma were resuspended in 200 μl of nuclease-free 1× PBS (pH 7.4; Ambion, Thermo Fisher Scientific) and divided into four equal fractions, each receiving one of the following treatments: (i) no treatment, (ii) RNase treatment only, (iii) protease followed by RNase treatment, and (iv) detergent followed by protease and RNase treatment. Detergent treatment was performed with 1% NP-40 (Thermo Fisher Scientific) for 15
MicroRNA library preparation and NGS. To determine the miRNA profile in SEs from naive as well as EAV persistently infected stallions, RNA was isolated from 8 SE samples (L136, prechallenge, 86 dpi, and 726 dpi; L140, prechallenge, 86 dpi, and 726 dpi; and L139, prechallenge and 86 dpi). The exosomal RNA was isolated as described above and submitted to System Biosciences for next-generation sequencing (NGS) (miRNA sequencing). Briefly, the NGS libraries were generated with TailorMix MicroRNA Sample Preparation version 2 (SeqMatic LLC, Fremont, CA). The 3' adapters were ligated to each RNA sample, and excess 3' adapters were subsequently removed. Subsequently, the 5' adapters were then ligated to the 3' adapter-ligated samples, followed by first-strand cDNA synthesis from the ligated products. The cDNA library was amplified and barcoded via enrichment PCR. The final RNA library was size selected on an 8% Tris-borate-EDTA (TBE) polyacrylamide gel. Sequencing was performed on the Illumina NextSeq 500 platform (1 × 75-bp single-end reads).

MicroRNA sequencing data analysis. The raw FASTQ files for each sample were trimmed three times with Trim Galore (Babraham Institute, Cambridge, UK). The trimmed FASTQ files were converted to FASTA format as required for using miRDeep2 (94). miRDeep2 produced an ARF file with the mapping details of all the reads that could be mapped to the Equus caballus reference genome (EquCab2.0) (95). If a read was mapped to >5 different locations in the reference sequence (-r setting in miRDeep2), it was not included in the final data set for analysis. Known miRNAs for Equus caballus were downloaded from miRBase (release 21) (96, 97). Many of the mapped reads could be assigned to either known equine miRNAs or tRNAs as described in the Genomic tRNA database (GtRNAdb) (98). The remaining mapped fragments were compared to the Ensembl annotation for the horse in order to identify miRNA and rRNA fragments. MicroRNA read counts were normalized using the formula log{[(reads × 1,000,000)/total reads] + 1} (79). For further analysis, miRNAs with at least three read counts in every sample were used.

Prediction of miRNA target genes. miRDB, TarBase v7.0 and the miRSearch target prediction tool (Exiqon, Woburn, MA) were used for miRNA target predictions (99, 100). Those genes with a target score of ≥0.80 (miRDB) or ≥0.4 (TarBase) were selected for further analysis. In addition, miRTarBase was used for identification of validated target genes (101). Analysis and identification of miR-128 binding site at the 3'-UTR of human CXCL16 mRNA (GenBank accession numbers NM_022059.3 and XM_001504756.5, respectively) was performed using TargetScanHuman release 7.1 (102) and IntaRNA 2.0 (Freiburg RNA Tools, Universität Freiburg, Germany) (103). The free energy (ΔG) of the predicted miRNA-mRNA interaction was estimated using the RNAhybrid tool (Universität Bielefeld, Germany) (104). A ΔG cutoff value of ≤−15 kcal/mol was used to assess the feasibility of the miRNA-mRNA interaction as previously suggested (105).

Functional annotation and pathway analysis. To investigate the molecular and biological functions of predicted miRNA target genes, DAVID Bioinformatics Resources (version 6.8 [https://david.ncifcrf.gov/]), National Institute of Allergy and Infectious Disease [NIAID], National Institutes of Health [NIH] (106, 107)) along with the PANTHER classification system (http://www.pantherdb.org) (108) was used to functionally annotate genes based on gene ontology (biological process, molecular function, and protein class) and perform pathway analysis.

MicroRNA quantitative real-time PCR. Expression levels of differentially expressed miRNAs identified by microRNA sequencing between any of the sequenced time points (prechallenge, 86 dpi, and 726 dpi) were determined by RT-qPCR. For this purpose, seminal plasma samples from naive (n = 15), short-term carrier (n = 7), and long-term carrier (n = 18) stallions were used. Among the samples derived from long-term carrier stallions (n = 18), two were derived from experimentally infected stallions described elsewhere (L136 and L140), while the remaining samples corresponded to naturally infected stallions which have been monitored for >5 years following infection by the OIE Reference Laboratory (Maxwell H. Gluck Equine Research Center). Additionally, sequential semen samples from the two experimentally infected stallions (n = 12) were used to evaluate eca-mir-128 expression dynamics. Exosomal RNA was reverse transcribed using the miScript II RT kit (Qiagen) following the manufacturer’s recommendations. For reverse transcription of mature miRNA, 10 ng of exosomal RNA was combined with 5× miScript HSpect buffer (4 μL), 10× miScript Nucleics mix (2 μL), and miScript reverse transcriptase mix (2 μL). The final reaction volume was 20 μL. The reaction mixture was incubated at 37°C for 1 h, followed by inactivation of the reverse transcriptase at 95°C for 5 min. cDNA was diluted 1:5 in nuclease-free water, aliquoted, and stored at −20°C until used. For qPCR, the miScript SYBR green PCR kit (Qiagen) was used. Briefly, 1 μL of cDNA was added to a reaction volume (9 μL) containing 2× Quant iTect SYBR green PCR master mix (5 μL), 10× miScript universal primer (1 μL), assay-specific primers (1 μL of a 5 μM stock; final concentration, 0.5 μM), and RNase-free water (2 μL). The cycling conditions included an initial PCR activation step (95°C for 15 min) followed by 40 cycles of denaturation at 94°C for 15 s, annealing at 55°C for 30 s, and extension at 70°C for 30 s. Melt curve analysis was performed to check for nonspecific amplifications along with the inclusion of no template controls. MicroRNA-specific forward primers were designed using miRprimer v2.0 (see Table S5 in the supplemental material) (109, 110). qPCR efficiencies and threshold cycle (CT) values were determined using LinRegPCR v2017.0 (111), with efficiencies in the range of 90.2 to 100.2%. CT values of >37 were not used for analysis. MicroRNA expression levels were normalized to three housekeeping miRNAs (eca-mir-30d, eca-mir-93, and eca-mir-128)
mir-103/eca-mir-107a), which were selected using NormFinder v20 to identify those miRNAs with the most consistent expression pattern across samples (79,112). MicroRNA expression levels were finally expressed as $2^{-\Delta\Delta CT}$.

**CXCL16 quantitative real-time PCR.** CXCL16 expression in the ampuellae was evaluated by TaqMan quantitative real-time PCR. Reverse transcription was performed using a high-capacity cDNA reverse transcription kit (Thermo Fisher Scientific) following the manufacturer’s recommendations. The reaction (20 μl) included 10 μl of total RNA (2 μg), 2 μl of 10X RT buffer, 0.8 μl 25X deoxynucleotide triphosphate (dNTP) mix (100 mM), 2 μl of 10X RT random primers, 1 μl of MultiScribe reverse transcriptase, 1 μl of RNase inhibitor, and 3.2 μl of nuclease-free water. The reverse transcription reaction mixture was incubated for 10 min at 25°C, followed by 120 min of incubation at 37°C and a final step at 85°C for 5 min. cDNA was diluted 1:1 in nuclease-free water and stored at −20°C until used. A custom TaqMan gene expression assay was developed for equine CXCL16 by a commercial company (Thermo Fisher Scientific) using the miRNA sequences derived from GenBank accession number XM_001504756.5. Each 10-μl qPCR mixture consisted of 4.5 μl of cDNA and 5.5 μl of master mix, which included 5 μl of TaqMan gene expression master mix (Thermo Fisher Scientific) and 0.5 μl of the custom TaqMan gene expression primer-probe mix (5’ fluorescein amide [6-FAM] labeled). The cycling conditions included a holding step at 50°C for 2 min and an activation step at 95°C for 10 min, followed by 40 cycles of denaturation (95°C for 15 s) and annealing/extension (60°C for 1 min). RT-qPCR efficiencies and $C_T$ values were determined as indicated above, and expression levels were normalized to those of three housekeeping genes (GAPDH, ACTB, and GUSB) and analyzed as indicated above.

**CXCL16-specific ISH (RNAscope) and immunohistochemistry (IHC) assays.** A CXCL16 mRNA-specific probe based on GenBank accession number XM_001504756.5 was developed by a commercial company (Advanced Cell Diagnostics [ACD], Newark, CA) and contained a total of 16 double Z-branched DNA probe pairs spanning a target region of 730 bp of the mRNA sequence (nt 598 to 1328). The probe was supplied in a ready-to-use format, and its specificity was evaluated using lymphoid tissues (palatine tonsils) and an equine endothelial cell line known to abundantly express CXCL16 (Fig. 7D). Probes specific to dihydrodipicolinate reductase B mRNA of *Bacillus subtilis* and peptidylprolyl isomerase B (PPIB) were used as negative and positive controls to assess assay specificity and RNA integrity in FFPE tissues, respectively. The RNAscope in situ hybridization (ISH) assay was performed using the RNAscope 2.0 HD red detection kit (ACD, Hayward, CA) as previously described, with some modifications (113), which included a second pretreatment for 15 min (100 to 104°C) and a third pretreatment for 20 min (40°C). The signal was finally detected using a fast red solution (red B and red A in a 1:60 ratio) for 10 min at room temperature. Slides were counterstained with 50% Gill hematoxylin I (American MasterTech Scientific, Lodi, CA) for 2 min, bluing was performed with 0.02% ammonium hydroxide in water, and sections were finally dried and mounted using EcoMount (Biocare Medical, Pacheco, CA). The ISH signal in the region of interest was quantified per the manufacturer’s recommendations as follows: 0, no staining or $<1$ dot every 10 cells at a magnification of ×40; 1, 1 to 3 dots every 10 cells at a magnification of ×40; 2, 4 to 10 dots/cell (visible at a magnification of ×20 to ×40); 3, >10 dots/cell, with scattered cells presenting dot clusters (visible at a magnification of ×20); and 4, >10 dots/cell, with frequent cells presenting dot clusters (visible at a magnification of ×20).

For IHC, sections of FFPE tissues (5 μm) were mounted on positively charged Superfrost Plus slides (Fisher Scientific, Pittsburgh, PA) and processed as previously described (26). Heat-induced epitope retrieval was performed using a modified citrate-based ready-to-use solution (pH 6.1; Dako, Carpinteria, CA) as described previously (26), and immunostaining was performed using the Bond Polymer Refine Detection kit (Leica Biosystems, Buffalo Grove, IL). The slides were incubated with 3% hydrogen peroxide (5 min), followed by incubation with CXCL16-specific rabbit antipeptide sera (diluted 1:1,000) in ISH/IHC Super Blocking [Leica Biosystems]) for 1 h at room temperature. This was followed by incubation with a polymer-labeled goat anti-rabbit IgG conjugated to HRP (8 min). 3,3′-Diaminobenzidine tetrahydrochloride (DAB) was used as the substrate, and the slides were incubated for 10 min. Finally, sections were counterstained with hematoxylin and mounted as previously described (26, 27, 113). Immunostaining was quantitatively scored based on the staining intensity within the region of interest (0, no staining; 1, minimal staining; 2, mild staining; 3, moderate staining; 4, high staining).

**Statistical analysis.** Data distribution, descriptive statistics, plots, and statistical tests were generated using JMP11 Pro statistical analysis software and SAS (SAS, Cary, NC). Heat maps were built using package d3heatmap in R. A one-way analysis of variance (ANOVA) was used to identify differentially expressed miRNAs (SAS), followed by a Benjamini-Hochberg correction. RT-qPCR data ($2^{-\Delta\Delta CT}$) values were analyzed using ANOVA (JMP11 Pro), and post hoc comparisons were performed using the Student t test (114). ISH and IHC scores were subjected to a nonparametric test (Kruskal-Wallis) using JMP11 Pro. Correlation analysis between CXCL16 $2^{-\Delta\Delta CT}$ and eca-mir-128 $2^{-\Delta\Delta CT}$ values was performed by the Spearman rank correlation method. Statistical significance was set at a $P$ value of $<0.05$ in all cases.

**Accession number(s).** The RNA sequencing data from this study were deposited in the Gene Expression Omnibus (GEO, NCBI, NIH) database under study GSE108180 (accession numbers GSM2891902 to GSM2891909).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at [https://doi.org/10.1128/JVI.00015-18](https://doi.org/10.1128/JVI.00015-18).

**SUPPLEMENTAL FILE 1**, XLSX file, 0.1 MB.

**SUPPLEMENTAL FILE 2**, XLSX file, 0.1 MB.
REFERENCES


10. Reference deleted.


12. Reference deleted.


SUPPLEMENTAL FILE 3, XLSX file, 0.1 MB.
SUPPLEMENTAL FILE 4, XLSX file, 0.1 MB.
SUPPLEMENTAL FILE 5, XLSX file, 0.1 MB.
ecA218-mir-128 and CXCL16 during EAV Persistence
Journal of Virology

May 2018 Volume 92 Issue 9 e00015-18 jvi.asm.org

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