



Extracellular Vesicle-Mediated RNA Release in Histoplasma capsulatum

🔟 Lysangela R. Alves,ª Roberta Peres da Silva,¹ David A. Sanchez,^c Daniel Zamith-Miranda,^c 🕑 Marcio L. Rodrigues,^{a,d} Samuel Goldenberg,^a Rosana Puccia,^b Joshua D. Nosanchuk^c

^aInstituto Carlos Chagas, Fiocruz, Curitiba, Cidade Industrial de Curitiba, Brazil

^bDepartamento de Microbiologia, Imunologia e Parasitologia da Escola Paulista de Medicina, Universidade Federal de São Paulo—UNIFESP, São Paulo, Brazil Departments of Medicine (Division of Infectious Diseases) and Microbiology and Immunology, Albert Einstein College of Medicine, Bronx, New York, USA ^dInstituto de Microbiologia, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil

ABSTRACT Eukaryotic cells, including fungi, release extracellular vesicles (EVs). These lipid bilayered compartments play essential roles in cellular communication and pathogenesis. EV composition is complex and includes proteins, glycans, pigments, and RNA. RNAs with putative roles in pathogenesis have been described in EVs produced by fungi. Here we describe the RNA content in EVs produced by the G186AR and G217B strains of Histoplasma capsulatum, an important human-pathogenic fungal pathogen. A total of 124 mRNAs were identified in both strains. In this set of RNA classes, 93 transcripts were enriched in EVs from the G217B strain, whereas 31 were enriched in EVs produced by the G186AR strain. This result suggests that there are important strain-specific properties in the mRNA composition of fungal EVs. We also identified short fragments (25 to 40 nucleotides in length) that were strain specific, with a greater number identified in EVs produced by the G217B strain. Remarkably, the most highly enriched processes were stress responses and translation. Half of these fragments aligned to the reverse strand of the transcript, suggesting the occurrence of microRNA (miRNA)-like molecules in fungal EVs. We also compared the transcriptome profiles of *H. capsulatum* with the RNA composition of EVs, and no correlation was observed. Taking the results together, our study provided information about the RNA molecules present in H. capsulatum EVs and about the differences in composition between the strains. In addition, we found no correlation between the most highly expressed transcripts in the cell and their presence in the EVs, reinforcing the idea that the RNAs were directed to the EVs by a regulated mechanism.

IMPORTANCE Extracellular vesicles (EVs) play important roles in cellular communication and pathogenesis. The RNA molecules in EVs have been implicated in a variety of processes. EV-associated RNA classes have recently been described in pathogenic fungi; however, only a few reports of studies describing the RNAs in fungal EVs are available. Improved knowledge of EV-associated RNA will contribute to the understanding of their role during infection. In this study, we described the RNA content in EVs produced by two isolates of Histoplasma capsulatum. Our results add this important pathogen to the current short list of fungal species with the ability to use EVs for the extracellular release of RNA.

KEYWORDS Histoplasma capsulatum, RNA, extracellular vesicles

istoplasma capsulatum is a major human fungal pathogen on the global stage that causes disease in both immunocompetent and immunocompromised individuals, albeit the risk for severe disease increases with compromised immunity (e.g., in patients with HIV infection or cancer as well as in individuals receiving steroids or tumor necrosis

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Address correspondence to Lysangela R. Alves, lys.alves@gmail.com, or Joshua D. Nosanchuk, josh.nosanchuk@einstein.yu.edu.

We describe the RNA content in EVs produced by two isolates of Histoplasma capsulatum. Our results add this important pathogen to the current short list of fungal species with the ability to use EVs for the extracellular release of RNA. @LyseBia

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factor alpha [TNF- α] blockers). In the United States, it is the most common cause of fungal pneumonia (1). *H. capsulatum* is of particular concern in certain developing regions (2), especially in Latin American countries, including Brazil (3, 4), Guatemala (5), and French Guiana, where it is considered the "first cause of AIDS-related death" (6). Despite its clear importance, enormous gaps exist in our understanding of the pathogenesis of histoplasmosis, the disease caused by *H. capsulatum*. An interesting facet of the biology of *H. capsulatum* is its ability to release extracellular vesicles (EVs) (7, 8).

EVs are bilayered lipid structures released by remarkably diverse cells across all kingdoms (9). We have demonstrated that EVs are present in both ascomycetes and basidiomycetes (7, 10–14). This observation implies that mechanisms for EV production and release are truly ancient, as they appear to predate the divergence of these branches 0.5–1.0 billion years ago. Fungal EVs can carry biologically active proteins, carbohydrates, lipids, pigments and nucleic acids (15, 16), many of which are constituents of the fungal cell wall and diverse others are associated with stress response and pathogenesis.

EV-mediated transport of fungal RNA was recently shown in both commensal and opportunistic fungi. EV RNA molecules, mostly smaller than 250 nucleotides (nt), were identified in *Cryptococcus neoformans, Paracoccidioides brasiliensis, Candida albicans, Saccharomyces cerevisiae*, and *Malassezia sympodialis* (17, 18). Since *H. capsulatum* packages diverse compounds within EVs, we postulated that it too would use these compartments to export RNA. In this study, the EV-associated RNA components were characterized in two different isolates of *H. capsulatum*. As described in other fungi, *H. capsulatum* EVs carry both mRNAs and noncoding RNAs (ncRNAs). In addition, proteomic data allowed the identification of 139 RNA-binding proteins (RBPs) in the EVs, suggesting that proteins involved in RNA metabolism might play an important role in cell communication through the EVs. Our results add this important pathogen to the list of fungal species with the ability to use EVs for the extracellular release of RNA.

RESULTS

Histoplasma capsulatum EVs contain RNA. We characterized the RNA molecules contained in EVs isolated from culture supernatant samples of *H. capsulatum* strains G186AR and G217B. These strains belong to distinct clades, and G217B has been shown to be more virulent than G186AR in experimental models (19, 20). The best-known difference between these two strains is that G217B lacks alpha-1,3-glucan on the yeast form cell wall (19, 20).

The reads obtained from the mRNA libraries (reads of >200 nt) were aligned with each strain-specific genome available at the NCBI (G186AR ABBS02 and G217B ABBT01). For data validation, we considered only sequences with expression values of transcripts per million (TPM) of \geq 100 in all biological replicates and transcripts with reads covering at least 50% of the coding DNA sequence (CDS). The small RNA (sRNA) fraction was analyzed for the presence of different species of noncoding RNAs (ncRNAs) by aligning the sRNA fraction (reads of <200 nt) with the *H. capsulatum* G186AR strain. These RNA molecules were compared between the strains in order to gain insights into the role of the EV RNA in this fungus and also to determine if there were differences with respect to composition between the two strains with distinct phenotypes.

Strain-specific content of EV RNA in *H. capsulatum.* We identified a total of 124 mRNA sequences in EV samples from the two strains and carried out paired comparisons between the G186AR and G217B samples. We applied the statistical negative binomial test with filters corresponding to TPM values of \geq 100, log2 values of \geq 2, and false-discovery-rate (FDR) values of \leq 0.05. We observed 93 transcripts enriched in EVs derived from the G217B strain, while 31 transcripts were enriched in the G186AR strain (see Table S1 in the supplemental material). In the G217B-associated transport (18%), oxidation-reduction mechanisms (12%), transmembrane transport (11%), and translation (8%) (Fig. 1). In the G186AR strain, the mRNA sequences were enriched only in general cellular and metabolic processes (59%). These results suggest that there are





FIG 1 Gene ontology analysis. The pie charts present the gene ontology of mRNA sequences enriched in EVs isolated from (A) *H. capsulatum* G217B (n = 93) and (B) *H. capsulatum* G186AR (n = 31).

important differences with respect to the mRNA composition of EVs derived from these two strains of *H. capsulatum*.

H. capsulatum EVs contain mRNA fragments and microRNA (miRNA)-like molecules. In addition to the identification of full-length transcripts in EVs, we also detected short reads of averages of 25 to 40 nt in length that aligned consistently in the CDS but at specific positions of the mRNAs (3' end, 5' end, or middle sequence); about 50% of these short fragments aligned to the reverse strand, including 172 (G217B) and 80 (G186AR) sequences of this type (Table 1). A total of 172 fragments were represented in the G217B sample compared to only 80 in the G186AR EVs (Table 1). About 47% of the reference mRNA translate proteins of unknown biological processes; this could be explained by the fact that around 33% of the genes annotated in H. capsulatum genome code hypothetical proteins and/or do not present a conserved domain, which impedes our current ability to determine specific biological activities. Those associated with DNA metabolism/biogenesis were the second most abundant for both EV samples (22 for G217B versus 16 for G186AR), followed by transport for G217B and by protein modification for both strain EVs. Other processes related to short RNAs identified in both strain EVs were oxidation-reduction, signaling, and carbohydrate and lipid metabolism (Table 1). RNA fragments associated with translation were highly enriched in G217B (n = 11) but not in G186AR (n = 2) EVs, while those related to response to stress were found exclusively in the G217B sample. The corresponding proteins are stress response protein whi2, DNA repair protein rad5, and a thermotolerance protein (Table 1). Analysis of translation-related sequences allowed identification of mRNA fragments associated with distinct steps of the translation process, such as ribosome biogenesis and processing. Other metabolic pathways identified in both strains were protein modification, carbohydrate, and lipid metabolism, signaling, oxidation-reduction, and transmembrane transport, among others (Table 1).



TABLE 1 Fragments of mRNAs identified in the EVs isolated from the G217B and G186AR strains^a

	G217B	G186AR		
Feature ID	alignment	alignment	Sequence description	GO
Protein modification				
HCBG_03026	5′R	5′R	Tetratricopeptide-like helical	Amino acid metabolic process
HCBG_05660	MR		CMGC SRPK protein kinase	Protein modification process
HCBG_05782	MF		Dihydrofolate synthetase fol3	Cofactor metabolic process
HCBG_06582	5′F		Aspartyl aminopeptidase	Peptidase activity
HCBG_07777	MF		Mitochondrial processing peptidase alpha	Peptidase activity
HCBG 08965	MF	MF	Tyrosine phosphatase	Protein modification process
HCBG 09127	3'R / 3'F		Proteasome component C5	Peptidase activity
HCBG 09175	5'F	5'F	Aspartic-type endopeptidase	Peptidase activity
HCBG 09182	MR		Protein kinase	Protein modification process
HCBG 01228	5'E		Oxidative stress-induced growth inhibitor 2	Pentidase activity
HCBG 01665	MF	MF	pH domain-containing protein	Protein modification process
HCBG_03811	MR	3'R	Heat shock protein Hsp98 Hsp104	ATPase activity peptidase activity
HCBG_00544	MF	5	Ubiquitin conjugating enzyme	Ligase activity
HCBG 02715	3'F	3'F	Ubiquitin family protein	2.9000 00000
HCBG 05116	3'F	5.	Protein	Protein modification process
HCBG 07497	51	3′F	Protein	Peptidase activity
		5.		
Carbohydrate metabolism				
HCBG_00058	5′R		Mannosyl-oligosaccharide alpha-mannosidase	Catabolic process
HCBG_00633	3'R / 3'NS		Class V chitinase	Catabolic process
HCBG_06620	3′R	3′R	Transaldolase	Carbohydrate metabolic process
Lipid metabolism				
HCBG_02433	MF	5'F	Acyl carrier protein	Biosynthetic process
HCBG_01540	MF	MF	Predicted protein	Lipid metabolic process
HCBG_04372		3′R	GPI anchor biosynthesis protein (Pig-f)	Lipid metabolic process
Description to attract				
	2/5		Concerned attended to the section White	
HCBG_02224	3 F		General stress response protein whiz	
HCBG_01643	3'K			Response to stress
HCBG_00190	3 K		Thermotolerance protein	
Translation				
HCBG 00808	ME	ME	60S ribosomal protein L15	
HCBG_00853	3'F		Small nucleolar ribonucleoprotein complex	
HCBG_01544	5'R / F	5'R	Ribosome biogenesis protein	
HCBG_02168	5'F / MF	5 11	60S ribosomal protein 125	Translation
HCBG_02499	5'R		rRNA processing protein Utp6	Oxidoreductase activity
HCBG_02762	3'F		60S ribosomal protein 31	Translation
HCBG_02702	MR		Prenyl cysteine carboxyl methyltransferase Ste14	mRNA processing
HCBG_04500	5'R		LeucyLtRNA synthetase	Translation
HCBG_03084	5'P		Transcription initiation protein Spt5	Translation
HCBG_04703	5'P		115 small nuclear ribonucleoprotein component	Chromosome organization
HCBG_06802	5'R		Ribosome biogenesis protein Ssf2	chromosome organization
11606_00002	511		hibbsome biogenesis protein 552	
Signaling process				
HCBG_00598	5'F / 5'NS		MinD kinetochore complex component Nnf1	Signal transduction
HCBG_03086*	5′R / F		Ste Ste20 paka protein kinase	Reproduction
HCBG_04646*		3′R	Protein Ras-2	Signal transduction
Uxidation-reduction	2/0	2/0 / 2/10	Developed a developed and the developed at 450	
HCBG_00763	3 K 2/D / 2 F	3 K / 3 NS	Benzoate 4-monooxygenase cytochrome p450	
HCBG_03251	3'R/3F		Tim-barrei enzyme family protein	Oxidoreductase activity
HCBG_04436	5'R/3'R	0/5	Flavin-containing monooxygenase	Oxidoreductase activity
HCBG_05481	3'F	3'F	Like subfamily b member 4	Protein folding
HCBG_05591	3'F	3 F	Emn-binding split-barrel-like protein	Oxidoreductase activity
HCBG_06890	5'F		Glutaredoxin	Homeostatic process
HCBG_08366	3'F		Conserved hypothetical protein	Oxidoreductase activity
HCBG_01233	5'R / 5'F		Galactose oxidase beta-propeller	
HCBG_00232		5'F	Tyrosinase	Oxidoreductase activity
HCBG_03159		MR	Ste Ste7 Mek1 protein kinase	Reproduction
Transport				
HCBG 00485	3′R		Vacuolar ABC heavy-metal transporter	Transmembrane transport
			· I	· · · · · · · · · · · · · · · · · · ·

TABLE 1 (Continued)



	G217B	G186AR		
Feature ID	alignment	alignment	Sequence description	GO
HCBG_00680	3′F		Arsenine resistance protein	Transmembrane transport
HCBG_00850	MR		MFS monocarboxylate	Transmembrane transport
HCBG_01089	5'F / 5'NS	5'R / 5'NS	Mitochondrial carrier	Iransport
HCBG_02374	5'K	E'D	Endosomal cargo receptor	Vesicle-mediated transport
HCBG_02985	5'D	5'D	Witechondrial dicarboxylate carrier	Transmombrane transport
HCBG_03738	211	ME	Execuse complex component Sec10	Vesicle-mediated transport
HCBG_04312	3'F	5'R / 3'F	Nonrepetitive nucleoporin	Nucleocytoplasmic transport
HCBG_04317	5'F	5 117 5 1	mRNA transport regulator	Transport
HCBG 04719	5'F		Nucleoporin	
HCBG_04608	3′R		MFS transporter	Transmembrane transport
HCBG_05671	MR		Actin-associated protein	Vesicle-mediated transport
HCBG_05941	5′F	5′R	Potassium uptake protein	Transmembrane transport
HCBG_05942	MR		Potassium uptake protein	Transmembrane transport
HCBG_06437	MF	MF	Oligopeptide transporter	Transport
HCBG_06658	MR		PX domain-containing protein	Transmembrane transport
HCBG_07112	MF		Ap-2 adaptor complex subunit	Vesicle-mediated transport
HCBG_07566	3'K	3 K / MK	Pan1	vesicie-mediated transport
HCBG_08252*	5'F		MFS multidrug transporter	Transmembrane transport
HCBG_09093	5′R		Kinetoplast-associated protein Kap	Transmembrane transport
HCBG_09150	5'R / 3'R		Cap binding protein	Transport
HCBG_04513	5'F		3-Oxoacyl-acyl-carrier-protein synthase	
DNA metabolism or				
biogenesis				
HCBG_00397	E/E		PHD finger domain	Chromosome organization
HCBG_00799		5 F 5'D / 2'E	C6 zinc finger domain containing protein	Pepudase activity
HCBG_07996	3'F	JN/JF	Recombination bot spot-binding protein	DNA metabolic process
HCBG_01721	3'F		Nitrogen assimilation transcription factor nira	Chromosome organization
HCBG 03125	5.	MF	White collar	Signal transduction
HCBG 03879	MR	MR	DNA-directed RNA polymerase I subunit	Biosynthetic process
HCBG_04485		3′F	Centromere protein Cenp-o	Chromosome organization
HCBG_04625	MR		C6 finger domain	Biosynthetic process
HCBG_04221	3′R		Chromatin remodeling complex subunit	Helicase activity
HCBG_05411	3′R	3′R	Transcription factor SteA	Reproduction
HCBG_05417	MF		Elongator complex protein 3	Biosynthetic process
HCBG_05986	5'F	a/5	G _{1/S} regulator	DNA metabolic process
HCBG_05814	3′R	3′R	Histone H2a	Chromosome organization
HCBG_06244		MF	double-strand-break repair protein	reproduction
HCBG_07395	MR		CP2 transcription factor	Biosynthetic process
HCBG_07428	3'F		Caf1 family ribonuclease	
HCBG_09164	MF	MF	C2H2 finger domain transcription factor	Biosynthetic process
	5'F 2'P	2'D	Formamidopyrimiding DNA alycosylase	DNA motobolic process
HCBG_04340	S R ME	S K ME	Telomere length regulation protein Elg1	lon hinding lipid hinding
HCBG_06146	5'R	5'R	Telomerase-binding protein Est1a	ion binding, lipid binding
HCBG_07560	5'R / 5'F	5'R / 5'F	DNA repair protein protein	
HCBG_05625	3′R	3′R	p60-like cell wall	
HCBG_09024	MR		Hlh transcription factor	
HCBG_06915	5'F	5'F	Proline-rich protein-15	Chromosome segregation
Other/unknown function				
HCBG_00048	5′R	5′R	Hypothetical protein HCBG_00048	Less less die e
HCBG_00453	5'K		NIL zinc finger protein	ion binding
HCBG_00075	3'F 5'P	5'P	ATPase AAA-5 protein	lon binding
HCBG 01015	э к MF	э к MF	Arrase AAA-3 protein Predicted protein	
HCBG_01082	3'R / 3'F	3'R	Zinc knuckle domain protein	
HCBG_01086	5'R	51	Predicted protein	
HCBG 01127	5'R / 3'R		Predicted protein	
HCBG_01146	MF		Predicted protein	
HCBG_01161	MF		Predicted protein	

TABLE 1 (Continued)



Feature ID	G217B alignment	G186AR alignment	Sequence description	GO
HCBG 01256	3′R		Conserved hypothetical protein	
HCBG_01258	MR		Predicted protein	
HCBG_01500	MR		Predicted protein	
HCBG_01656	MF		Predicted protein	
HCBG_01888	3'R	3'R	Conserved hypothetical protein	
HCBG_01052	3'E	511	Conserved hypothetical protein	
	51		Drotoin	
	5 K 5/F		Protein Dradistad protain	
HCBG_02107	5 F	2/5		
HCBG_02158		3'F	Conserved hypothetical protein	
HCBG_02464	3'R / 3'F	3'F / 3'R / 3'NS	Carbohydrate-binding module family 48 protein	
HCBG_02569	MR / MF	MF	Predicted protein	
HCBG_02659	MR / MF	MR	Predicted protein	
HCBG_02697	3′R	3′R	Predicted protein	
HCBG_02981	MF		Phosphotransferase enzyme family protein	
HCBG_02986	MF	5'F	Predicted protein	
HCBG 03093	MR		PH domain protein	
HCBG_03374	MF	MF	Glutathione transferase	
HCBG_03658	3'R / 3F		Conserved hypothetical protein	Helicase activity
HCBG_03692	3'R / 3F		Predicted protein	Theneuse derivity
			Dredicted protein	
			Predicted protein	
HCBG_03805	MF		MD was not an establish	
HCBG_03899	MR	MR/3R	wD repeat protein	
HCBG_03911	3′R	3′R	Protein	
HCBG_03913	MR		Hypothetical protein HCBG_03913	
HCBG_03980	MR		Phosphatidylserine decarboxylase	
HCBG_04009	MR		Hypothetical protein HCBG_04009	
HCBG_04186	MR		Conserved hypothetical protein	
HCBG_04193	3′R	3′R	Conserved hypothetical protein	
HCBG 04201	3′F		Hypothetical protein HCBG 04201	
HCBG_04208	3′F	3′F	Conserved hypothetical protein	
HCBG_04365	MF		Hypothetical protein HCBG 04365	
HCBG_04371	5'R / 5'F		Bifunctional uridylyltransferase uridylyl-removing	
HCBG 04380	3'D	3'P	Predicted protein	
	2/0	211	Predicted protein	
	ס <i>ה</i> סיס	2/0	Producted protein	
HCBG_04432		5 N	Predicted protein	
HCBG_04780	5 K	5 R	Bromodomain-containing protein	
HCBG_04887		MR	Predicted protein	
HCBG_05336	5'R		UPF0160 domain protein	
HCBG_05404	3′R / 3′F		Predicted protein	
HCBG_05580	3′R		Methyltransferase domain-containing protein	
HCBG_05638	5′R		Predicted protein	
HCBG_05703	5′R		Conserved hypothetical protein	
HCBG_05744	5'F		T-complex protein 1 subunit beta	
HCBG 05763	3′R	3′F	Conserved hypothetical protein	
HCBG_05878	3′F		Hypothetical protein HCBG 05878	
HCBG_06018	5'F		Cytomegalovirus GH-receptor family	
HCBG_06054	MR		Phosphotransferase family protein	Ion hinding kinase activity
HCBG_06071	ME	ME	Protein	ion binang, kindse detivity
	MD	1411	Conserved hypothetical protein	
HCBG_06114	3 F		Protein	
HCBG_06176	3'F	= 10	KH domain protein	RNA binding
HCBG_06239		5'R	Nonsense-mediated mRNA decay protein	
HCBG_06270	MR		Predicted protein	
HCBG_06364	MR		F-box domain-containing protein	
HCBG_06436	MF		Predicted protein	
HCBG_06661		5′NS	Predicted protein	
HCBG_06677	3′F		Predicted protein	
HCBG_06927	3′R / 3′F		Predicted protein	
HCBG_07002	5'R / 5'F	5′R / 5′F	Ketoreductase	
HCBG 07065	5'F		Predicted protein	
HCBG 07214	5'R	5′R	Predicted protein	
HCBG 072/7	MR	5 11	Acultransferase 3	Transferring acyl groups
		MD	Acylitatistetase 3	mansiening acyl groups
NCBG 0/296	MIK	MK	hypothetical protein HCBG 0/296	

TABLE 1 (Continued)

Feature ID	G217B alignment	G186AR alignment	Sequence description	GO
HCBG_07377	MF	MR	Predicted protein	
HCBG_07484	3′F		Rhomboid family membrane protein	Peptidase activity
HCBG_07611	MR / MF	MR / MF /	Protein	
		MNS		
HCBG_07676	3′R / 3′F		Lyr family protein	
HCBG_07802	3′R / 3′F	3′R / 3′F	Predicted protein	
HCBG_07811	3′F	3′F	Predicted protein	
HCBG_08059	MR	MF	DUF833 domain protein	Protein complex assembly
HCBG_08505	3′F		Sucrase ferredoxin domain-containing protein	
HCBG_08661	MF	MF	Predicted protein	
HCBG_08693	3′R		Set domain protein	
HCBG_08838	5′R		WW domain	
HCBG_08850	5′R		Integral membrane protein	
HCBG_09013	5′F	5'F	Predicted protein	
HCBG_09099	5′R	5′R	Conserved hypothetical protein	
HCBG 09144	MF		Predicted protein	

^aFor some transcripts, there was an alignment in specific positions of the mRNA, not covering the entire sequence. 5', 3', or M (middle of the mRNA) followed by an "F" or an "R" represents forward (F) or reverse (R) orientation. GO, gene ontology; GPI, glycosylphosphatidylinositol; ID, identifier; mtDNA, mitochondrial DNA.

To gain further insight into the role of EV RNAs, to determine if they could be derived from a miRNA-like pathway, and to assess if they could play a biological role in the recipient cell, we searched for RNA secondary structures, since they are fundamental for gene expression regulation (21). A broad study of RNA structures in distinct cells revealed regulatory effects of the RNA structure throughout mRNA life cycle such as polyadenylation, splicing, translation, and turnover (22, 23). Using the entire range of EV RNA sequencing (RNA-seq) data, a total of 33 RNAs with putative structures were generated by a probability distribution, using a free energy (Δ G) value of less than or equal to -7.0 (Table S2). On the basis of this parameter, we identified transcripts for U3 small nucleolar RNA-associated protein, L-isoaspartate O-methyltransferase, serine/threonine-protein kinase, proteasome component C5, pre-rRNA processing protein Utp22, C-x8-C-x5-C-x3-H zinc finger protein, fungus-specific transcriptional repressor RPH1 (Fig. 2; see also Table S2).

Comparison of EV ncRNA classes in *H. capsulatum* **EVs.** We used the ncRNA database from *H. capsulatum* to identify the classes of ncRNA present in EV RNAs. The data analysis revealed 73 different sequences of ncRNA in *H. capsulatum* EVs from the G186AR strain and 38 from the G217B isolate. A total of 33 molecular species were common to both strains, 40 were exclusively identified in the G186AR strain, and the most abundant class of ncRNA found in *H. capsulatum* EVs consisted of tRNAs (Table 2).

Analysis of proteins putatively associated with RNA metabolism in the EVs. As a rule, cellular RNAs are covered with proteins and exist as ribonucleoprotein (RNP) complexes. The proteins associated with RNAs are named RNA-binding proteins (RBPs). These proteins participate in several biological processes, ranging from transcription to RNA decay (24). In this context, we investigated the presence of RBPs in the H. capsulatum EVs. We analyzed the proteomic EV data available for the G217B strain (25), and we identified 139 proteins related to RNA metabolism (8) (Table 3; see also Table S3). We found many RBPs, such as poly(A) binding protein (PABP), Nrd1, Prp24, and Snd1; splicing factors, exosome complex components, and ribosomal proteins (Table 3; see also Table S3) were identified. In addition, we also found quelling-deficient protein 2 (QDE2), an Argonaute protein important in the RNA machinery in fungi. Because we identified the QDE2 in EVs, we searched for the components of the RNA interference (RNAi) machinery in H. capsulatum and compared them with the proteins from Neurospora crassa and Schizosaccharomyces pombe, which are the fungal species for which the RNAi machinery was best described previously (26, 27). H. capsulatum EVs contained one Argonaute protein (QDE2), two Dicer-like proteins, the QIP (quelling interaction protein), and the RNA-dependent RNA polymerase (QDE1) (Table 4).







FIG 2 RNA secondary structure. We used ppFold software to predict the secondary structure from the putative miRNAs extracted from the obtained reads. The numbers in parentheses represent the alignment E values. The colors indicated for the nucleotides represent the reliability percentage for each position of the RNA molecule (bottom panel). The stability value corresponding to each structure is given in kilocalories/mole.

Comparisons of cellular RNA versus EV RNA showed a distinct enrichment of molecules in the vesicles. We next assessed the composition of cellular RNA from *H. capsulatum* yeast cells (28) and compared this information to that obtained from analyses of EV-associated RNA composition under the same conditions. There was no correlation between the transcripts with highest expression levels and their presence in the EVs (Table S4). Examples of highly expressed cellular transcripts included histones 4, 2B, and 2A, allergen Aspf4, chaperones, and translation factors, among others (Table S4). In contrast, zinc knuckle domain-containing protein, vacuolar ATP synthase subunit C, $G_{1/5}$ regulator, thermotolerance protein, histone variant H2A.Z, and proteasome component C5 had an enrichment value of greater than 7,000 in the EVs, while they showed low expression values in the cell (Table S4). The differences in composition between cells and EVs were also evaluated by grouping the transcripts into biological

TABLE 2 Classes of ncRNA sequences identified in EV preparations from *H. capsulatum* strains G186AR and G217B^a

RNA category and ncRNA	G186AR	G217B
rRNA		
15S_rRNA	_	Х
NTS1-2	Х	_
RDN18-1	Х	Х
RDN18-2	Х	Х
RDN25-1	Х	—
RDN25-2	Х	Х
RDN37-1	Х	_
RDN37-2	Х	—
RDN5-1	Х	Х
RDN5-2	Х	Х
RDN5-3	Х	Х
RDN5-4	Х	Х
RDN5-5	Х	Х
RDN5-6	Х	Х
RDN58-1	Х	Х
RDN58-2	Х	Х
ncRNA		
RUF21	Х	Х
snoRNA	X	V
SNR54	X	X
tRNA		
tRNA-Ser	—	Х
tRNA-Met	—	Х
tRNA-Gln	—	Х
tRNA-Cys	—	Х
tRNA-Ser	Х	Х
tRNA-Pro	Х	Х
tRNA-Ala	Х	Х
tRNA-Thr	Х	Х
tRNA-Ala	Х	Х
tRNA-Phe	Х	Х
tRNA-Ala	Х	Х
tRNA-Asn	Х	Х
tRNA-Met	Х	Х
tRNA-Arg	Х	Х
tRNA-Trp	Х	Х
tRNA-Gly	X	Х
tRNA-Asp	X	Х
tRNA-Pro	X	Х
tRNA-Thr	X	Х
tRNA-His	X	Х
tRNA-Glu	X	X
tRNA-GIn	X	X
tRNA-Tyr	X	X
tRNA-GIn	X	Х
tRNA-GIy	X	
tRINA-Lys	X	_
tRNA-lie	X	—
tRNA-Leu	X	_
tRNA-Met	X	_
	X	—
INNA-IIE	×	_
	^ V	_
INIA-LYS	X	—
	A V	_
tDNA Dha	^ V	_
tRNA-FIE	A X	
tRNA-Sec	x	_
tRNA-Asp	X	_
tRNA-Thr	Х	_



TABLE 2 (Continued)

RNA category and ncRNA	G186AR	G217B
tRNA-Ile	Х	_
tRNA-Ser	Х	_
tRNA-Ser	Х	_
tRNA-Arg	Х	_
tRNA-Lys	Х	_
tRNA-Leu	Х	_
tRNA-Ser	Х	_
tRNA-Leu	Х	_
tRNA-Ala	Х	_
tRNA-Cys	Х	_
tRNA-Thr	Х	_
tRNA-His	Х	_
tRNA-Tyr	Х	_
tRNA-Ser	Х	_
tRNA-Leu	Х	_
tRNA-Lys	Х	_
tRNA-Ala	Х	_
tRNA-Pro	Х	_
tRNA-Arg	Х	_
tRNA-Glu	Х	_

^aX, present; —, absent.

processes (Fig. 3). For the yeast cells, the main pathways were associated with transport, translation, and general metabolic processes (Fig. 3). For the EVs, the enriched pathways were transmembrane transport, protein phosphorylation, and transcription regulation (Fig. 3). This result demonstrates the low levels of correlation between the most highly expressed cellular mRNAs and EV cargo, providing evidence that there might be a mechanism directing the RNA molecules to the EVs.

DISCUSSION

As previously described (17, 18), RNA molecules associated with fungal EVs are remarkably diverse. For instance, mRNAs, tRNA fragments, snoRNAs, small nucleolar RNAs (snRNAs), and miRNA-like molecules were characterized in EVs from C. albicans, C. neoformans, P. brasiliensis, and S. cerevisiae (17). We observed similar distributions of RNA molecules in H. capsulatum EVs. The comparison between the G186AR and G217B EVs revealed important differences in the variety of mRNAs identified. When the mRNA composition was compared to what was described for other fungi, important similarities were observed. For example, the most abundant biological process identified in G217B EVs was vesicle-mediated transport, which was also the most abundant process in C. albicans EVs (17). Molecules required for ribosome biogenesis, which were observed in G217B EVs, belonged to the most highly enriched process in S. cerevisiae EVs (17). However, in the comparisons of the ncRNA molecules, different profiles were observed. Most of the ncRNAs in H. capsulatum strains derived from tRNAs; a similar profile was obtained with C. albicans (17). In addition, almost no snoRNAs were identified in H. capsulatum, but this class of ncRNAs was one of the most abundant in the EVs of other fungi (17). Differences in EV composition were observed previously in C. neoformans; the EV-associated RNA produced by mutant cells with defective unconventional secretion differed considerably from similar samples produced by wild-type cells (29).

In our study, we identified short reads that aligned specifically to exons; however, these sequences did not correspond to complete mRNAs in the EVs. They instead corresponded to 25-nt-long fragments that were enriched in specific exons of the transcript. These fragments of mRNAs were previously described in human cells (30), where most of the transcripts identified in the EVs corresponded to a fraction of the mRNA with an enrichment of the 3' UTR of the transcript (30). The results of that human study led to the hypothesis that the mRNA fragments had a role in gene expression regulation in the recipient cells as the secreted mRNA could act as competitors to





TABLE 3 Proteins related to RNA metabolism identified in EV preparations from H. capsulatum strain G217B

Majority protein ID	Protein name	Gene name
C0NMG7	QDE2 protein	HCBG_03944
C0P170	Cap binding protein	HCBG_09150
C0NJ23	Exosome complex exonuclease RRP4	HCBG_03153
CONM03	Exosome complex exonuclease RRP45	HCBG_04533
CONCT3	KH domain RNA-binding protein	HCBG_00929
CONUHO	KH domain RNA-binding protein	HCBG_07001
C0NIU5	KH domain-containing protein	HCBG_02352
CONUS5	mRNA 3'-end-processing protein RNA14	HCBG_06689
CONNW0	mRNA cleavage and polyadenylation factor CLP1	CLP1 HCBG_04840
CONP91	mRNA decapping enzyme	HCBG_04971
CONC87	mRNA export factor Mex67	HCBG_00733
C0NJ33	Nuclear and cytoplasmic polyadenylated RNA-binding protein Pub1	HCBG_03163
C0NQQ9	Poly(A) ⁺ RNA export protein	HCBG_05339
CONSS5	Polyadenylate-binding protein (PABP)	HCBG_06205
CONKR4	Ribonucleoprotein	HCBG_03744
C0NSY4	RNA binding domain-containing protein	HCBG_06264
CONWH9	RNA-binding protein	HCBG_07509
CONB22	RNA-binding protein	HCBG_00318
CONPA1	RNA-binding protein Nrd1	HCBG_04981
CONZI9	RNA-binding protein Prp24	HCBG_08569
CONTZ5	RNA-binding protein Snd1	HCBG_06625
CONMQO	RNP domain-containing protein	HCBG_04027
C0NLQ4	RRM domain-containing protein	HCBG_04434
C0NJ27	Transcription elongation factor Spt6	HCBG_03157
C0NTQ1	Transcription initiation factor TFIID complex 60-kDa subunit	HCBG_06531
CONRU6	U1 snRNP-associated protein Usp106	HCBG_05876
C0NZZ2	U1 snRNP-associated protein Usp107	HCBG_08722
CONBS3	U2 snRNP auxiliary factor large subunit	HCBG_00569
C0NAD4	U3 small nucleolar RNA-associated protein	HCBG_00080
C0NZA3	U3 small nucleolar RNA-associated protein 22	HCBG_08483
CONLW4	U3 snoRNP-associated protein Rrp5	HCBG_04494
COPORO	U6 snRNA-associated Sm-like protein LSm2	HCBG_08990
C0P041	30S ribosomal protein S10	HCBG_08883
CONFV8	40S ribosomal protein S15	HCBG_01774
CONX47	40S ribosomal protein S18	HCBG_08039
C0NZD2	40S ribosomal protein S20	HCBG_08512
CONBDO	40S ribosomal protein S21	HCBG_00426
CONUDO	40S ribosomal protein S3	HCBG_06961
CONLP3	405 ribosomal protein 54	HCBG_04423
CONF40	405 ribosomal protein 55A	HCBG_01506
CUNLRS	40S ribosomal protein S9	HCBG_04445
CONTH6	5'-3' exoribonuclease 1 (EC 3.1.13)	HCBG_06456
CONKI2	605 ribosomal protein L1	HCBG_03662
CONNL2	605 ribosomai protein L3	HCBG_04742
CUNCP3	605 ribosomal protein L30	HCBG_00889
CUNRDO	COS ribosomal protein LO	HCBG_05366
CUNQRO	And DNA seven has extra the	HCBG_05346
	Acyl-RINA-Complex Subunit	
	Aldnine-trive evidese (EC 0.1.1.7) (aldnyi-trive synthetase) (Aldrs)	ALAT HCBG_03098
CONDEE	Arcinul +DNA curthotoco	
	Arginyi-triva synthetase	
	Asparaging tPNA synthetase	
CONCV7	Asparagingi-trink synthetase	
	ATP dependent bolicase NAM7	
	ATP dependent PNA belicase DOP1	
	ATP-dependent NNA helicase DOBT	HCBG_02344
CONECZ	Coll cycle control protoin	
	Cleavage and polyadenylation specific factor 5	HCBG_01393
	Clustered mitochandria protein homolog (protein TIC21 homolog)	
	Custoinul_tPNA_synthetase	
	Cystemyrthina synthetase D Aminoacul tDNA doaculaso (EC 2111) (EC 21106)	
	D-Anninuacyr-inina uedcyldse (EC 5.1.1.7) (EC 5.1.1.90)	
	DNA directed RNA polymerase il polypeptide	
	Diva-unecteu Riva polymerase subunit deta (EC 2.7.7.6)	
	Elicitor protein Eukanyatia paptida chain ralaaca factar CTD hinding subunit	
	Eukaryouc peptice chain release factor GTP-binding subunit	LCRG_02A10

TABLE 3 (Continued)



Majority protein ID	Protein name	Gene name
COPO imes 7	Eukaryotic translation initiation factor 3 subunit D (EIF3D)	HCBG_09057
CONEV9	Fibrillarin	HCBG_01425
CONZT8	Glutaminyl-tRNA synthetase	HCBG_08668
CONKS5	Glutamyl-tRNA synthetase	HCBG_03755
CONE28	Glycyl-tRNA synthetase	HCBG_02121
CONN35	Histidyl-tRNA synthetase	HCBG_04162
CUNL66	Isoleucyl-tRNA synthetase, cytoplasmic	HCBG_03896
CUNZR4	LeucyI-tRNA synthetase	HCBG_08644
CONICO	Leucyi-tRNA synthetase	
CONMS8	Mitotic control protein dis3	HCBG_03034
CONBIS	mRNA splicing protein PRP8	HCBG_00494
CONV83	NAM9 ⁺ protein	HCBG_07877
CONG69	Nucleic acid-binding protein	HCBG_01885
CONUD1	Phenylalanyl-tRNA synthetase subunit beta	HCBG_06962
CONBD1	Phenylalanyl-tRNA synthetase subunit beta cytoplasmic	HCBG_00427
CONUP1	Polymerase II polypeptide D	HCBG 06655
C0NNC4	Pre-mRNA-processing factor 39	HCBG 04251
C0NJB4	Pre-mRNA-processing protein prp40	HCBG_03244
C0NXM8	Pre-mRNA-splicing factor	HCBG_08220
C0NLW7	Prolyl-tRNA synthetase	HCBG_04497
C0NW72	Ribonuclease T2-like protein	HCBG_07402
C0NEF9	Ribonuclease Z	HCBG_01275
C0NIJ3	Ribosomal biogenesis protein Gar2	HCBG_02250
C0NHN4	Ribosomal protein L14	HCBG_02856
C0NI43	Ribosomal protein L6	HCBG_03015
C0NVX9	Ribosomal protein S5	HCBG_07309
C0NN82	RNA helicase (EC 3.6.4.13)	HCBG_04209
C0NEY2	RNA polymerase II largest subunit	HCBG_01448
C0NL28	RNA polymerase subunit	HCBG_03858
CONYA7	RNase H domain-containing protein	HCBG_07901
CONH14	RNP domain-containing protein	HCBG_02636
CONDP9	RNP domain-containing protein	HCBG_01992
CONC99	SAM domain-containing protein	HCBG_00/45
CONEPT	Seryi-tRNA synthetase	HCBG_02184
CUNSR2	Signal recognition particle subunit SRP68 (SRP68)	HCBG_06192
	Silicing factor 2A subunit 2	
CONTRO	Splicing factor 3B	HCBG_06950
CONBR2	Splicing factor 3B subunit 1	HCBG_00558
CONG79	Threonyl-tRNA synthetase	HCBG_02621
CONSBO	Transfer RNA-Trn synthetase	HCBG_06040
CONL23	tRNA (cytosine-5-)-methyltransferase NCI 1	HCBG_03853
CONUP2	tRNA [guanine(37)-N1]-methyltransferase (EC 2.1.1.228)	TRM5 HCBG 06656
CONEYO	tRNA guanylyltransferase	HCBG 01446
C0NJJ2	tRNA ligase (EC 6.5.1.3)	HCBG_03322
C0NM44	tRNA pseudouridine synthase	HCBG_04574
C0NSG9	Tyrosine-tRNA ligase (EC 6.1.1.1) (Tyrosyl-tRNA synthetase)	HCBG_06099
C0NP46	Uncharacterized protein	HCBG_04926
C0NZF6	Uncharacterized protein	HCBG_08536
CONIA9	Uncharacterized protein	HCBG_03081
C0NMF3	Uncharacterized protein	HCBG_04683
CONPI9	Uncharacterized protein	HCBG_05069
CONKI6	Uncharacterized protein	HCBG_03666
CONF97	Uncharacterized protein	HCBG_01563
CONEJI	Uncharacterized protein	HCBG_01307
CONEC3	Uncharacterized protein	HCBG_01239
CONJN9	Uncharacterized protein	HCBG_03369
	Uncharacterized protein	
	Uncharacterized protein	
CONKEA	Uncharacterized protein	HCBC 03624
CONGR7	Uncharacterized protein	HCBG 07380
CONM01	Uncharacterized protein	HCBG_04531
	protein	



TABLE 3 (Continued)

Majority protein ID	Protein name	Gene name
C0NG47	Uncharacterized protein	HCBG_01863
C0NEU7	Uncharacterized protein	HCBG_01413
C0NG27	ValyI-tRNA synthetase	HCBG_01843
C0P019	Vip1 protein	HCBG_08749
C0NG23	Ribosome biogenesis protein RPF2	HCBG_01839
C0NGE8	Ribosome biogenesis protein TSR3	TSR3 HCBG_02420
CONAE4	Ribosome biogenesis protein YTM1	YTM1 HCBG_00090

regulate stability, localization, and translation of mRNAs in target cells (30). In *Mucor circinelloides* cells, the presence of the RNA silencing pathway (sRNA) resulted in the production of both sense and antisense sRNAs (31–33). Sequencing analysis of the sRNA content of this fungus showed the existence of exonic small interfering RNAs (exo-siRNAs) as a new type of sRNA. They were produced from exons of the same genes that are later regulated through the repression of the corresponding mRNA (34). This result agrees with our observation of short reads in the exonic regions of the transcripts. We therefore hypothesize that, similarly to what was described for *M. circinelloides* cells, *H. capsulatum* EV fragments can regulate expression of their own mRNAs. Of note, we also found a highly represented population of putative exonic RNA in *Paracoccidioides* strains (R. Peres da Silva, L. V. G. Longo, J. P. C. da Cunha, T. J. P. Sobreira, H. Faoro, M. L. Rodrigues, S. Goldenberg, L. R. Alves, and R. Puccia, unpublished data).

As H. capsulatum EVs contain different RNA molecules, it is reasonable to hypothesize that proteins that regulate RNA metabolism are also present in the EVs, probably associated with RNA. If validated, this hypothesis could indicate how the RNAs in a specific subset are directed to the vesicles and exported. RNA-binding proteins (RBPs) participate in several biological processes, from RNA transcription to decay (24). We detected a number of RNA-binding proteins in H. capsulatum EVs (25). These proteins were also identified in association with EVs in other systems. For example, in the EVs produced by human epithelial cells, 30 RBPs were identified (35), including heterogeneous nuclear ribonucleoproteins (hnRNPs). These proteins are responsible for directing pre-mRNAs in the maturation processes that culminate in transcriptional regulation, alternative splicing, transport, and localization (35). In addition, RBPs in EVs were identified in distinct models as hepatocytes, human embryonic kidney (HEK) cells, and mouse myoblast cells (35–37). Interestingly, one of the RBPs identified in EVs was SND1 (staphylococcal nuclease domain-containing protein 1), which is a main component of the RNA-induced silencing complex (RISC) that plays an important role in miRNA function (37).

Another example of a protein identified in the EVs of *H. capsulatum* and distinct organisms is an endonuclease of the Ago2 family. An infection model with *Plasmodium falciparum* demonstrated that infected red blood cells released EVs containing functional miRNA-Argonaute 2 complexes (38). Moreover, endothelial cells internalized the

TABLE 4 Proteins associated with the RNAi machinery in H. capsulatum G186AR EVs compared to S. pombe and N. crassa

H. capsulatum	G186AR		%	%
product	ID	E value	identity	positives
QDE2 protein	HCBG_03944	1.00E-85	28	45
QDE2 protein	HCBG_03944	1.00E-178	37	53
Dicer-like protein	HCBG_01751	1.00E-113	28	44
Dicer-like protein 2	HCBG_01136	3.00E-97	31	49
RNA-dependent RNA	HCBG_06604	3.00E-92	31	46
polimerase				
Dicer-like protein	HCBG_01751	0.00E + 00	45	60
QDE-2-interacting	HCBG_07373	2.00E-50	27	43
	H. capsulatum product QDE2 protein QDE2 protein Dicer-like protein 2 RNA-dependent RNA polimerase Dicer-like protein QDE-2-interacting protein (OIP)	H. capsulatum productG186AR IDQDE2 protein QDE2 proteinHCBG_03944QDE2 proteinHCBG_03944Dicer-like protein 2 Dicer-like protein 2 RNA-dependent RNA polimeraseHCBG_01751 HCBG_0604Dicer-like protein polimeraseHCBG_01751 HCBG_0604Dicer-like protein polimeraseHCBG_01751 HCBG_01751 HCBG_07373	H. capsulatum productG186AR IDE valueQDE2 protein QDE2 proteinHCBG_039441.00E-85 1.00E-178Dicer-like proteinHCBG_017511.00E-113 3.00E-97Dicer-like protein 2 polimeraseHCBG_011363.00E-97 3.00E-92Dicer-like proteinHCBG_066043.00E-92 2.00E+00Dicer-like proteinHCBG_017510.00E + 00 2.00E - 50	H. capsulatum product G186AR ID % QDE2 protein HCBG_03944 1.00E-85 28 QDE2 protein HCBG_03944 1.00E-178 37 Dicer-like protein HCBG_01751 1.00E-113 28 Dicer-like protein 2 HCBG_01136 3.00E-97 31 RNA-dependent RNA polimerase HCBG_01751 0.00E + 92 31 Dicer-like protein HCBG_01751 0.00E + 00 45 QDE-2-interacting HCBG_07373 2.00E-50 27





FIG 3 Gene ontology analysis. The pie charts present the gene ontology of mRNA sequences enriched in *H. capsulatum* cells (A) and in EVs isolated from *H. capsulatum* (B).

P. falciparum EVs, and the miRNA-Argonaute 2 complexes were transferred to the cells and acted in regulation of gene expression and in the barrier properties of the recipient cells (38). The Argonaute protein named QDE2 in *H. capsulatum* was identified as enriched in the EVs of the G217B strain.

The small silencing RNAs include a variety of molecules, such as microRNAs (miRNAs) and various small interfering RNAs (siRNAs), including exo-siRNAs, endogenous siRNAs (endo-siRNAs), and Piwi-interacting RNAs (piRNAs) (39). Previous studies of small RNAs in fungi identified the RNAi machinery in the fission yeast species Schizosaccharomyces pombe, in the budding yeast species Saccharomyces castellii and C. albicans, and in filamentous fungi (26, 27, 40). One of the best-characterized models is represented by the filamentous fungus N. crassa (27, 41-45). The RNAi machinery in that organism functions in defense against transposons (46). A similar process has been described in C. neoformans, where RNAi is involved in the regulation of transposon activity and genome integrity during vegetative growth (47). In N. crassa, the QDE2 gene encodes an Argonaute protein that is homologous to the rde-1 gene in C. elegans, encoding a protein required for double-stranded RNA (dsRNA)-induced silencing (27). The characterization of RNAs associated with QDE2 in N. crassa led to the identification of miRNA-like RNAs (milRNAs) in this organism (48). The identification of QDE2 in H. capsulatum EVs in association with the small RNAs indicated that the QDE2-milRNA complex might be directed to the EVs and possibly delivered to recipient cells, with the potential to interfere with gene expression regulation and/or cell-cell communication.

Fungal EVs have been implicated in a number of communication processes, including transfer of virulence (49) and antifungal resistance (50). In *Cryptococcus gattii*, pathogen-to-pathogen communication via EVs resulted in reversion of an avirulent phenotype through mechanisms that required vesicular RNA (49). The sequences required for this process, however, remained unknown. This is an efficient illustration of the potential derived from the characterization of EV-associated RNA in fungi. In this context, our study results provide information from the *H. capsulatum* model that will allow the design of pathogenic experimental models aiming at characterizing the role of extracellular RNAs in fungal pathogenesis.

MATERIALS AND METHODS

Fungal strains and growth conditions. The *H. capsulatum* strains were subjected to long-term storage at -80° C. Aliquots were inoculated into Ham's F-12 media (Gibco; catalog no. 21700-075) supplemented with glucose (18.2 g/liter), L-cysteine (8.4 mg/liter), HEPES (6 g/liter), and glutamic acid (1 g/liter) and cultivated at 37°C with constant shaking at 150 rpm. Viability assessments were performed using Janus green 0.02%, and all aliquots used had >99% live yeast cells. EVs were then isolated from fungal culture supernatants as previously described (12).

sRNA isolation. Small RNA-enriched fractions were isolated using a miRNeasy minikit (Qiagen) and were then treated with an RNeasy MinElute cleanup kit (Qiagen), according to the manufacturer's protocol, to obtain small RNA-enriched fractions. The sRNA profile was assessed in an Agilent 2100 Bioanalyzer (Agilent Technologies).

RNA sequencing. Purified sRNA (100 ng) was used for RNA-seq analysis with two independent biological replicates. The RNA-seq analysis was performed using a SOLiD 3 Plus platform and an RNA-Seq kit (Life Sciences) according to the manufacturer's recommendations.

In silico data analysis. The sequencing data were analyzed using version 10.1 of CLC Genomics Workbench. The reads were trimmed on the basis of quality, with a threshold Phred score of 25. The reference genomes used for mapping were obtained from the NCBI database (H. capsulatum G186AR strain ABBS02 and G217B strain ABBT01). The alignment was performed using the following parameters: additional number of bases of upstream and downstream sequences, 100; minimum number of reads, 10; maximum number of mismatches, 2; nonspecific match limit, -2, minimum fraction length, 0.7 for the genome mapping or 0.8 for the RNA mapping. The minimum proportion of read similarity mapped on the reference genome was 80%. Only uniquely mapped reads were considered in the analysis. The libraries were normalized per million, and the expression values for the transcripts were recorded in RPKM (reads per kilobase per million). We also analyzed other expression values, including TPM (transcripts per million) and CPM (counts per million). The statistical test applied was the DGE (differential gene expression) test. For the ncRNA analysis, the database used was the ncRNA database from Histoplasma capsulatum (EnsemblFungi G186AR GCA_000150115 assembly ASM15011v1). The secondary structure analysis was performed using the PPFold plugin in CLC Genomics Workbench v. 10.1 and the default parameters. The entire RNA-seq database was subjected to PPFold analysis, and the putative structures were determined. Analysis of the relationship between the profile of RNA sequences detected in this study and the protein composition of H. capsulatum EVs was based on results recently obtained with strain G217B using a proteomic approach (25). The cellular RNA used in this analysis was assessed using the Sequence Read Archive (SRA) database (accession numbers SRR2015219 and SRR2015223) (28).

Data availability. The data were deposited into the SRA database under study accession number PRJNA514312.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/ mSphere.00176-19.

 TABLE S1, XLSX file, 1.4 MB.

 TABLE S2, XLSX file, 0.01 MB.

 TABLE S3, XLSX file, 0.1 MB.

 TABLE S4, XLSX file, 2.1 MB.

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We declare that we have no conflicts of interest.



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