SUPPORTING INFECTIOUS DISEASE RESEARCH

### Toxoplasma gondii, Clone G2 AF

#### Catalog No. NR-10239

**Product Description:** *Toxoplasma gondii* (*T. gondii*), G2 AF is a non-virulent recombinant F1 clone selected from progeny of two parallel genetic crosses between a highly virulent Type I parental strain, GT1-FUDR3.3 and a non-virulent Type III parental strain, CTG.11 ARA-SNF.

#### Lot<sup>1</sup>: 58499194

#### Manufacturing Date: 07APR2009

TEST	SPECIFICATIONS	RESULTS
Genotyping <sup>2,3</sup>		
AK16 locus ( <i>Hinf</i> l digestion)	Consistent with parental Type III strain	Consistent with parental Type III strain
L358 locus (HaelII digestion)	Consistent with parental Type III strain	Consistent with parental Type III strain
Drug susceptibility <sup>4</sup>		
Sinefungin (SNF)	Sensitive	Sensitive
Adenine arabinoside (Ara-A)	Resistant	Resistant
Viable Cell Count by Hemacytometry (pre-freeze)	> 10 <sup>6</sup> cells/mL	3.6 x 10 <sup>7</sup> cells/mL
Viability (post-freeze) <sup>5</sup>	Growth	Growth
Sterility (21-day incubation)		
Harpo's HTYE broth <sup>6</sup> , 37°C and 26°C, aerobic	No growth	No growth
Trypticase soy broth, 37°C and 26°C, aerobic	No growth	No growth
Sabouraud broth, 37°C and 26°C, aerobic	No growth	No growth
Brain Heart Infusion, 37°C and 26°C, aerobic	No growth	No growth
Sheep blood agar, 37°C, aerobic	No growth	No growth
Sheep blood agar, 37°C, anaerobic	No growth	No growth
Thioglycollate broth, 37°C, anaerobic	No growth	No growth
Mycoplasma Contamination		
DNA Detection by PCR	None detected	None detected

<sup>1</sup>NR-10239 was produced by cultivation of the deposited material in human foreskin fibroblast cells (ATCC<sup>®</sup> CRL-1634<sup>™</sup>) with cell cultivation medium for parasites (<u>ATCC medium 2222</u>: adjusted to contain 10% heat-inactivated fetal bovine serum). The culture was propagated in 95% air, 5% CO<sub>2</sub> for 8 days at 37°C, until lysis of the host cell monolayer was reached.

<sup>2</sup>PCR amplification was performed separately for the two loci AK16 and L358. Where appropriate, samples were subjected to restriction enzyme digestion typing by agarose gel electrophoresis.

<sup>3</sup>Primer sequences, annealing temperatures, and conditions for restriction enzyme digestion may be obtained at the *Toxoplasma* Genome Map website (<u>Toxoplasma Genome Map</u>).

<sup>4</sup>Sinefungin was used at a concentration of 2.7 x 10<sup>-7</sup> M and ara-A was used at a concentration of 1.3 x 10<sup>-4</sup> M, as described (Sibley, L. D., et al. "Generation of a Restriction Fragment Length Polymorphism Linkage Map for *Toxoplasma gondii.*" <u>Genetics</u> 132 (1992): 1003-1015. PubMed: 1360931.)

<sup>5</sup>Viable cells and signs of infection were seen after 24 days under cultivation conditions at 37°C.

<sup>6</sup>Atlas, Ronald M. Handbook of Microbiological Media. 3rd ed. Ed. Lawrence C. Parks. Boca Raton: CRC Press, 2004, p. 798.

Date: 31 AUG 2009

## Signature: Signature on File

# Title: Technical Manager, BEI Authentication or designee

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