Genome Annotation Submission Cover Sheet

Preliminary Annotation Review Checklist 5-15-2018

Phage Name:

Odesza

Y	Your Name:		Warren Sconiers
Your Institution:		stitution:	University of the Ozarks
Υ	our em	nail:	wsconiers@ozarks.edu
		litional emails: r correspondence)	
	1. [1. [2. / 3. 4. 5. / 6. 7.	tee the Online Does the gence ame as the performance all the general Have the general Have all old Blace the locus the the the locus the	ox indicating completion of each task. If you are not sure how to do something, Bioinformatics manual page "How to Pass Preliminary Review". In the sequence in your final contain the same number of bases and is it the osted sequence on phagesdb.org? In the ses "valid" when you click the "validate" button? In the ses been renumbered such that they go sequentially from 1 to the highest that they go sequentially from 1 to the highest that they seem cleared, and all gene features reBLASTed? In the season of the sequence
	9. F	or the items I	below, generate a genome profile, and review the following. For the
		a. Have and b. Does even c. Do the find the formula of the find th	me_CompleteNotes.dnam5 file: ny duplicate genes (or any with the same stop coordinate?) been removed? very gene have one and only one complete set of Notes functions in the Notes match the official function list? hree lines of functional evidence described for EVERY gene? notes contain the initial Glimmer/GeneMark data from the autoannotation? PhageName .dnam5 file: ny duplicate genes (or any with the same stop coordinate?) been removed? otes field empty (including hidden marks?) function names in the Product field either match the official function list or rpothetical Protein"? unction field empty (including hidden marks?)
	 Did you use PECAAN to annotate your phage? If, so please describe how in the text field after question 11. Describe any issues or specific genes that you were unable to satisfactorily resolve, and warrant further inspection in the Quality Control review. We did not find any tRNAs using Aragorn and tRNAscan SE. We checked Kerry, Gravy, Duffy, and Rickmore genomes from our gene cluster (DJ) and we		
did not find any tail assembly chaperones, so therefore did not check for frameshifts.			

We added gene 8 so it does not have the initial Glimmer/GeneMark data.

We did not use PECAAN.