



BV-BRC

Bacterial and Viral (BV) -
Bioinformatics Resource Center (BRC)

Test Plan

BV-BRC Beta Integrated Data and Tools Testing

Issued to:

National Institute of Allergy and Infectious Diseases
National Institute of Health

Contract No.: 75N93019C00076

Contract Title: Bioinformatics Resource Centers for Infectious Diseases

Revision Date:

July 6, 2022

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Purpose

This Test Plan is intended to **assess correct integration and operation of PATRIC and IRD/ViPR data, tools, and visualizations into the BV-BRC system (Beta version)**, which is based on the PATRIC system framework.

Description

Beta Testing is performed by representative and real users of the system to assess its functionality, usability, reliability and compatibility. It also ensures that there are no major failures in the system, and that it satisfies requirements and representative use cases from an end-user perspective. Beta Testing is performed by BV-BRC team members, external collaborators, and users who are early adopters of the new BV-BRC. To fully assess the accuracy and performance of the various components, former PATRIC team members will perform the bacteria-oriented test cases, and JCVI team members will perform the virus-oriented test cases since each group is most familiar with the legacy systems, data, and tools.

Since the BV-BRC is implemented using components of the constituent PATRIC and IRD/ViPR systems, both of which have long performance histories and validated performance, **the critical aspect of this testing is to ensure that data and tools are correctly integrated and producing results comparable with the validated legacy systems**. To do so, the test cases are composed of representative use cases, with results to be compared to validated results, typically from the legacy systems, modified if needed to account for updated backend data or presentation style. Where appropriate, links to test data sets, system documentation, and reference information is provided.

Test results will be recorded in a corresponding **Test Report** (as a separate deliverable), and triaged as follows:

- “Passed” - Test results meet all expected criteria.
- “Partial Success” - Test results meet key expected criteria, but have minor issues or suggestions for future enhancements. Issues and suggestions are recorded and tracked in the BV-BRC GitHub repository. These are used for implementation in the next version of the component.
- “Failed” - Test results do not meet key expected criteria due to a critical issue or software bug. These issues are recorded and tracked in the BV-BRC GitHub repository. These are queued for prompt remediation, then fixed, tested, and re-deployed in the system.

Organization

The Test Plan is organized into logical sections based on the types of components: Organism Data (menu), Searches, Data Pages, Tools & Services, and Workspace. Each Component has a Test Description with a basic description, test parameters and/or data, and criteria for success. Where appropriate, links to reference material are provided. Finally, each Item to Test is assigned a person(s) with Responsibility for performing the test and evaluating the results.

Searches, Global and Advanced

ID	Component	Test Description	Responsibility
S1	Global Search	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/, top right on any page <p>Procedure:</p> <ul style="list-style-type: none"> Test the Global Search using representative search criteria for bacterial and viral data. Test using example Keywords. Test using combinations Data Type and Keywords. Inspect search results to verify that they match search criteria. For viral data, test using representative criteria and verify that returned genome counts are comparable with IRD/ViPR. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/global_search.html 	ARW, CZ, AN, RK
S2	Taxa Search	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/searches/TaxaSearch <p>Procedure:</p> <ul style="list-style-type: none"> Test the Taxa Search using representative search criteria for bacterial and viral data. Test using example Keywords. Test using Taxon ID, Genetic Code, and Taxon Name. Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/searches_menu.html https://www.bv-brc.org/docs/quick_references/organisms_taxon/taxonomy.html 	ARW, EL, DD, RK
S3	Genomes Search	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/searches/GenomeSearch <p>Procedure:</p> <ul style="list-style-type: none"> Test the Genome Search using representative search criteria for bacterial and viral data. Test using example Keywords Test using combinations of Pathogen Group, Host Name, Host Group, Taxon Name, and Geographic Group. Inspect search results to verify that they match search criteria. For viral data, test using representative criteria and verify that returned genome counts are comparable. <p>References:</p>	ARW, CZ, AN, RK

		<ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/genome_table.html 	
S4	Strains Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/StrainSearch <p>Procedure</p> <ul style="list-style-type: none"> • Test Strains search and results page for segmented viruses: (Influenza, Arenaviridae, Lassa). • Inspect search results to verify that they match search criteria. • Test using representative criteria and verify that returned genome counts are comparable with IRD/ViPR. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html 	CZ, AN, RK
S5	Proteins Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/GenomicFeatureSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Proteins Search using representative search criteria for bacterial and viral data. • Test using example Keywords. • Test using BRC ID, Product Name, and Taxon Name. • Inspect search results to verify that they match search criteria. • For viral data, test using representative criteria and verify that returned protein counts are comparable with IRD/ViPR. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/proteins.html 	ARW, CZ, AN, RK
S6	Specialty Genes Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/SpecialtyGeneSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Specialty Genes Search using representative search criteria. • Test using example Keyword, Pathogen Group, and Taxon Name. • Test using Keyword and Pathogen Group only. • Inspect search results to verify that they match search criteria. <p>References:</p>	ARW, RK

		<ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/specialty_genes.html 	
S7	Domains and Motifs Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/DomainAndMotifSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Domains and Motifs Search using representative search criteria. • Test using example Keyword. • Test using example Taxon Name. • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html 	ARW, RK
S8	Epitopes Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/EpitopeSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Domains and Motifs Search using representative search criteria. • Test using example Pathogen Group and Protein Name. • Test using example Keyword • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html 	RK
S9	Protein Structures Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/ProteinStructureSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Protein Structures Search using representative search criteria. • Test using example Taxon Name and PDB ID. • Test using example Keyword • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/protein_structures.html 	ARW, RK
S10	Pathways Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/PathwaySearch 	ARW, RK

		<p>Procedure:</p> <ul style="list-style-type: none"> • Test the Pathways Search using representative search criteria. • Test using example Pathway Name and Genome ID. • Test using example Keyword • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/pathways.html 	
S11	Subsystems Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/SubsystemSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Subsystems Search using representative search criteria. • Test using example Keyword and Taxonomy Name. • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/subsystems_tab.html 	ARW, RK
S12	Surveillance Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/SurveillanceSearch <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Surveillance Search using representative search criteria. • Test using example Pathogen Test Type and Collection Year range. • Test using example Host Common Name and Collection Country. • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/surveillance_data.html 	YZ, GT, RK
S13	Serology Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/searches/SerologySearch 	ZW, RK

		<p>Procedure:</p> <ul style="list-style-type: none"> • Test the Serology Search using representative search criteria. • Test using example Test Type and Host Type. • Test using example Host Species and Geographic Group. • Inspect search results to verify that they match search criteria. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/searches_menu.html • https://www.bv-brc.org/docs/quick_references/organisms_taxon/serology_data.html 	
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Data Pages

ID	Component	Test Description	Responsibility
D1	Overview Tab (Bacteria Landing Pages)	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Bacteria/2 <p>Procedure:</p> <ul style="list-style-type: none"> • Test the bacterial data Overview Tabs. • Verify that links from each group go to the correct location. • Check other links to verify correct operation. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/overview.html 	ARW, RK
D2	Overview Tab (Viruses Landing Page)	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Virus/10239 <p>Procedure:</p> <ul style="list-style-type: none"> • Test the viral data Overview Tabs. • Verify that links from each group go to the correct location. • Check other links to verify correct operation. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/overview.html 	AN, RK
D3	Phylogeny Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/662#view_tab=phylogeny <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Phylogeny Tab. 	ARW, RK

		<ul style="list-style-type: none"> • Test the data tab with example bacterial genus. • Test phylogram and cladogram view. • Test node selection. • Test Genome Group and Genome View action buttons. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/overview.html 	
D4	Taxonomy Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/629#view_tab=taxontree <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Taxonomy Tab with example bacterial and viral data. • Verify the correct Taxonomy. • Test branch expansion and collapse • Test Taxon Overview action button. • Test Genomes and Features action buttons. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/taxonomy.html 	ARW, RS, RK
D5	Strains Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/11320#view_tab=strains_orthomyxoviridae <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Strains Tab with example viral data. • Verify the correct strains. • Test no strains reported for non-segmented viruses <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_menu.html#browsing-bv-brc-by-taxon 	CZ, AN, RK
D6	Genomes Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/1301#view_tab=genomes <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Genomes Tab with example bacterial and viral data. • Verify the correct genomes. • Test Keyword Filter • Test Advanced Search • Test Filters • Test Genome, Genomes, and Genome Group action buttons 	ARW, CZ, AN, RK

		<p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/genomes.html 	
D7	AMR Phenotypes Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/1301#view_tab=amr <p>Procedure:</p> <ul style="list-style-type: none"> • Test the AMR Phenotypes Tab with example bacterial data. • Verify display of Antibiotic, Resistant Phenotype, Evidence, Laboratory Typing Method • Test Antibiotic action button including Overview, AMR Phenotypes, AMR Genes, and AMR Regions displays. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/amr_phenotypes.html 	ARW, RK
D8	Sequences Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/194#view_tab=sequences <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Sequences Tab with example bacterial data. • Verify appropriate genomes • Test FASTA and Genome Browser action buttons. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/sequences.html 	ARW, RK
D9	Proteins (Features) Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/194#view_tab=features <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Proteins Tab with example bacterial and viral data. • Verify appropriate genomes. • Test Genome action button. • Test FASTA action buttons. • Test ID Map. • Test MSA. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/proteins.html 	ARW, CZ, AN, RK
D10	Protein Structures Tab	<p>URL:</p>	ARW, RS, RK

		<ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/194#view_tab=features <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Protein Structures Tab with example bacterial and viral data. • Verify structures exist. • Test Structure action button. • Test example structure operations. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/protein_structures.html 	
D11	Specialty Genes Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/32008#view_tab=specialtyGenes&filter=false <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Specialty Genes Tab with example bacterial data. • Verify display of Specialty Genes category • Test Pathway action button. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/specialty_genes.html 	ARW, RK
D12	Domains and Motifs Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/234#view_tab=proteinFeatures&filter=false <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Specialty Genes Tab with example bacterial data. • Verify display of Domain Source categories. • Test filtering by a Source. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_menu.html#browsing-bv-brc-by-taxon 	ARW, RK
D13	Epitopes Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/11320#view_tab=epitope&filter=false <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Epitopes Tab with example bacterial and viral data. • Verify appropriate genomes. 	ARW, RS

		<ul style="list-style-type: none"> • Test Filter. • Test Epitope action button. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_menu.html#browsing-bv-brc-by-taxon 	
D14	Surveillance Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/11320#view_tab=surveillance <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Epitopes Tab with example bacterial and viral data. • Verify appropriate genomes. • Test Filter. • Test Epitope action button. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/surveillance_data.html 	YZ, GT, RK
D15	Serology Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/11320#view_tab=serology <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Serology Tab with example viral data. • Test Filters. • Test Serology Record. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/serology_data.html 	ZW, RK
D16	Experiments Tab	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Taxonomy/1763#view_tab=experiments • https://www.bv-brc.org/view/Genome/10090.24#view_tab=experiments <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Experiment Data Tab with example bacterial and mouse host response datasets. • Test Filters / facets on the experiment page. • Select single experiment and review experiment details. • Test interactive gene list, heatmap viewer, and clustering tool. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_taxon/experiments.html 	ARW, ZW, RK

D17	Interactions Tab	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/view/Taxonomy/209#view_tab=interactions <p>Procedure:</p> <ul style="list-style-type: none"> Test the Interactions Tab with example bacterial data. Test Filters. Test Graph View. Test Graph functions. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/organisms_taxon/interactions.html 	ARW, RK
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Analysis Tools and Services

Preparation for testing of the BV-BRC Tools and Services consists of identifying exemplar use cases, including input data and parameters, and creating validated test result(s) using established instances of the tool(s) either in legacy systems or standalone versions. The test itself is then performed in the BV-BRC system using the specified input data and parameters, and the results are compared using key output content and metrics to verify that the tool or service is operating properly.

ID	Component	Test Description	Responsibility
A1	Genome Assembly Service - Bacteria	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/app/Assembly <p>Procedure:</p> <ul style="list-style-type: none"> Test the Genome Assembly Service using exemplar reads sets for bacterial genomes. Test input options, i.e., single-end and paired-end read sets using files uploaded to the workspace and using an SRA run accession as input. Test the assembly strategies, i.e., Auto, Unicycler, SPAdes, Canu, MetaSPAdes, PlasmidSPAdes, and MDA. For each job submitted, verify successful completion of the job, presence of output files, and quality of the assembled contigs by comparing them with the same or closely related public genome. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/services/genome_assembly_service.html https://www.bv-brc.org/docs/tutorial/genome_assembly/assembly.html 	ARW, MS

A2	Genome Annotation Service - Bacteria	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Annotation <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Genome Annotation Service using exemplar genome sequences for bacterial genomes. • Test input options, i.e., contig file from the user's machine or from the user's workspace. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the annotations by comparing them with the same, or closely related, public genome. • Verify successful integration of the genome in BV-BRC by reviewing the genome overview pages and other genome-level tabs. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/genome_annotation_service.html • https://www.bv-brc.org/docs/tutorial/genome_annotation/genome_annotation.html 	ARW, MS
A3	Genome Annotation Service - Phages	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Annotation <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Genome Annotation Service using exemplar genome sequences for phage genomes. • Test input options, i.e., contig file from the user's machine or from the user's workspace. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the annotations by comparing them with the same, or closely related, public genome. • Verify successful integration of the genome in BV-BRC by reviewing the genome overview pages and other genome-level tabs. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/genome_annotation_service.html • https://www.bv-brc.org/docs/tutorial/genome_annotation/genome_annotation.html 	ARW, MS
A4	Genome Annotation Service - Viruses	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Annotation <p>Procedure:</p>	MS

		<ul style="list-style-type: none"> • Test the Genome Annotation Service using exemplar genome sequences for viral genomes. • Test input options, i.e., contig file from the user's machine or from the user's workspace. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the annotations by comparing them with the same, or closely related, public genome. • Verify successful integration of the genome in BV-BRC by reviewing the genome overview pages and other genome-level tabs. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/genome_annotation_service.html • https://www.bv-brc.org/docs/tutorial/genome_annotation/genome_annotation.html 	
A5	Comprehensive Genome Analysis Service (Bacteria)	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/ComprehensiveGenomeAnalysis <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Comprehensive Genome Analysis service using exemplar bacterial datasets. • Test input options, i.e., single end or paired end read files from workspace, sear sets using SRA accessions, or assembled contigs from workspace. • Test assembly strategies, i.e., Auto, Unicycler, SPAdes, Canu, MetaSPAdes, and PlasmidSPAdes. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the assembly and annotations by comparing them to the same or similar public genome. • Verify successful integration of the genome in BV-BRC by reviewing genome overview pages and other genome level tabs. • Review the quality and accuracy of the comprehensive genome report by comparing the summary stats with those available on the genome overview page. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/comprehensive_genome_analysis_service.html • https://www.bv-brc.org/docs/tutorial/comprehensive_genome_analysis/comprehensive_genome_analysis.html 	ARW, MS

A6	BLAST (Homology) Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Homology <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Homology / BLAST service using exemplar queries for both bacteria and viruses. • Test input options, i.e. various BLAST programs, nucleotide and protein query sequences. • Test various BLAST databases, i.e. reference genome databases, taxon level databases, and genome specific databases for both bacterial and viruses. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the results. • Test result selection and action from the result table. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/blast.html • https://www.bv-brc.org/docs/tutorial/blast/blast.html 	ARW
A7	BLAST (Homology) Service - Short Peptide Search	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Homology <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Homology / BLAST service using exemplar short peptide queries. • Test various BLAST databases, i.e. reference genome databases, taxon level databases, and genome group databases. • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the results. • Test result selection and action from the result table, including MSA to visualize variations. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/blast.html • https://www.bv-brc.org/docs/tutorial/blast/blast.html 	ZW, MS
A8	Similar Genome Finder Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/GenomeDistance <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Similar Genome Finder Service using exemplar bacterial genome. • Test input options, i.e., genome id / genome name, fasta contig file, and fastq file as input. • Test different databases, i.e., reference / representative genomes and all public genomes. • For each job submitted, verify successful completion of the job and the quality of the search results. • Test the selection and actions from the search result 	ARW

		<p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/similar_genome_finder_service.html • https://www.bv-brc.org/docs/tutorial/similar_genome_finder/similar_genome_finder.html 	
A9	Meta-CATS Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/MetaCATS <p>Procedure:</p> <ul style="list-style-type: none"> • Test the service using exemplar datasets • Test input options, i.e. feature groups and alignment files. • Test auto grouping using various metadata attributes • For each job submitted, verify successful completion of the job, availability of the output files in the workspace, and quality of the results. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/metacats.html • https://www.bv-brc.org/docs/tutorial/metacats/metacats.html 	YZ, MS
A10	Phylogenetic Tree Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/PhylogeneticTree <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Phylogenetic Tree Service using exemplar bacterial genome groups. • Test input options, i.e. selecting genomes one by one or using genome groups. • Test the tree parameters, i.e. building trees with 10, 50, or 100 conserved genes. • For each job submitted, verify successful completion of the job, presence of output files, and quality of the phylogenetic tree. • Review the interactive tree viewer and verify all functions are working as expected. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/phylogenetic_tree_building_service.html • https://www.bv-brc.org/docs/tutorial/phylogenetic_tree/phylogenetic_tree.html 	ARW, MS
A11	Genome Alignment Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/GenomeAlignment <p>Procedure:</p>	ARW, MS

		<ul style="list-style-type: none"> • Test the Genome Alignment Service using exemplar bacterial genomes. • Test input options, i.e., adding one genome at a time and using a genome group. • For each job submitted, verify successful completion of the job and presence of output files. • Review the interactive genome alignment viewer and ensure the quality of the results. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/genome_alignment_service.html 	
A12	Primer Design Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/PrimerDesign <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Primer Design Service using exemplar bacterial and viral genes. • Test input options, i.e., using FASTA sequence as query or a FASTA sequence file from the workspace. • For each job submitted, verify successful completion of the job, presence of output files, and quality of the results. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/primer_design_service.html • https://www.bv-brc.org/docs/tutorial/primer_design/primer_design.html 	ARW, YZ, MS
A13	Variation Analysis Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Variation <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Variation Analysis Service using exemplar bacterial datasets. • Test input options, i.e., read files in FASTQ format and SRA accessions. • Test different alignment strategies, i.e. BWA-mem, BWA-mem-strict, Bowtie2 and LAST. Test different SNP callers, such as FreeBayes and SAMtools. • For each job submitted, verify successful completion of the job, presence of output files, and the quality of results. • Review SNPs and alignment files in interactive genome browser. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/variation_analysis_service.html 	ARW, MS

		<ul style="list-style-type: none"> • https://www.bv-brc.org/docs/tutorial/variation_analysis/variation_analysis.html 	
A14	Tn-Seq Analysis Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Tnseq <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Tn-seq Analysis Service using exemplar reads sets for bacterial genomes. • Test different strategies, i.e., essential genes and conditionally essential genes. • For each job submitted, verify successful completion of the job, presence of output files, including bam, wig, counts, transit stats files. Verify the quality of the results. • Review the results in the genome browser. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/tn_seq_analysis_service.html • https://www.bv-brc.org/docs/tutorial/tn-seq/tn-seq.html 	ARW, MS
A15	MSA and SNP Analysis Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/MSA <p>Procedure:</p> <ul style="list-style-type: none"> • Test the MSA and SNP Service using exemplar bacterial and viral gene and protein sequences. • Test input options, i.e., feature groups, fasta sequence files, and sequence input box. • Test different MSA algorithms, MAFFT and Muscle. • For each job submitted, verify successful completion of the job, presence of output files in various formats, review resulting MSA and SNPs. • View MSA using interactive MSA viewer and verify all functionality. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/msa_snp_variation_service.html • https://www.bv-brc.org/docs/tutorial/msa_snp_variation/msa_snp_variation.html 	CZ, YZ, MS
A16	Phylogenetic Tree (Gene Tree) Service and Viewer	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/GeneTree <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Gene Tree Service using exemplar bacterial and viral gene and protein sequences. • Test input options, i.e., genome group, feature group, and fasta sequence files. 	CZ, MS

		<ul style="list-style-type: none"> • Test different tree algorithms, i.e. RAXML, PHYML, and FASTTREE. • For each job submitted, verify successful completion of the job, presence of output files in various formats, review resulting gene trees. • View tree using interactive phylogenetic tree viewer and verify all functionality. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/genetree.html • https://www.bv-brc.org/docs/tutorial/genetree/genetree.html 	
A17	Proteome Comparison Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/SeqComparison <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Proteome Comparison Service using exemplar bacterial genomes. • Test input options, i.e., selecting genomes one by one, using genome group, and protein fasta file. • For each job submitted, verify successful completion of the job, presence of output files, and quality of results. • Review interactive proteome comparison viewer and its functions as expected. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/proteome_comparison_service.html • https://www.bv-brc.org/docs/tutorial/proteome_comparison/proteome_comparison.html 	ARW, MS
A18	Metagenomic Read Mapping Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/MetagenomicReadMapping <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Metagenomic Read Mapping Service using exemplar bacterial and metagenomic reads sets. • Test input options, i.e., single-end / paired-end read files and SRA accessions. • Search against antimicrobial resistance gene database (CARD) and virulence factor database (VFDB). • For each job submitted, verify successful completion of the job and presence of output files. • Review the list of AMR and virulence genes detected. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/metagenomic_read_mapping_service.html 	ARW, MS

		<ul style="list-style-type: none"> • https://www.bv-brc.org/docs/tutorial/metagenomic_read_mapping/metagenomic_read_mapping.html 	
A19	Taxonomic Classification Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/TaxonomicClassification <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Taxonomic Classification Service using exemplar metagenomic reads sets. • Test input options, i.e., single-end / paired-end read files and SRA accessions. • For each job submitted, verify successful completion of the job, presence of output files and their format. • Review and verify the taxonomic classification results using tabular reports. • Review and verify the results using interactive taxonomic classification viewer, Krona. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/taxonomic_classification_service.html • https://www.bv-brc.org/docs/tutorial/taxonomic_classification/taxonomic_classification.html 	ARW, MS
A20	Metagenomic Binning Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/MetagenomicBinning <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Metagenomic Binning Service using exemplar metagenomic reads sets. • Test input options, i.e., single-end / paired-end read files and SRA accessions. • For each job submitted, verify successful completion of the job and presence of output files. • Review the resulting metagenomic bins and their quality, including completeness and contamination. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/metagenomic_binning_service.html • https://www.bv-brc.org/docs/tutorial/metagenomic_binning/metagenomic_binning.html 	ARW, MS
A21	Expression Import Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/Expression <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Expression Import Service using exemplar bacterial gene expression datasets. 	ARW, MS

		<ul style="list-style-type: none"> For each job submitted, verify successful completion of the job and presence of output files. Review the dataset using interactive gene list. Test filters to identify differentially expressed genes. Test interactive heatmap viewer and clustering tool to identify genes with similar expression patterns across one or more samples. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/services/expression_data_import_service.html https://www.bv-brc.org/docs/tutorial/expression_import/expression_import.html 	
A22	RNA-Seq Analysis Service	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/app/Rnaseq <p>Procedure:</p> <ul style="list-style-type: none"> Test the RNA-seq Analysis using exemplar transcriptomic reads sets. Test input options, i.e., single-end / paired-end read files. Test different strategies, i.e. Tuxedo and HTSeq. Test differential and non-differential analysis options. For each job submitted, verify successful completion of the job, presence of output files, their content and format. Review differential expression results using interactive gene list and heatmap viewer. <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/services/rna_seq_analyses_service.html https://www.bv-brc.org/docs/tutorial/rna_seq/rna_seq.html 	ARW, MS
A23	ID Mapper Service	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/app/IDMapper <p>Procedure:</p> <ul style="list-style-type: none"> Test the ID Mapper Service using exemplar BVBR and RefSeq gene identifiers. Test input options, i.e mapping BVBR identifiers to external identifiers and reverse. For each job submitted, review and verify mapped identifiers. Select mapped records and test various actions available in the action bar, i.e. download results as table 	ARW, MS

		<p>or fasta, view corresponding features or genomes, and create genome or feature groups.</p> <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/id_mapper.html • https://www.bv-brc.org/docs/tutorial/id_mapper/id_mapper.html 	
A24	Fastq Utilities Service	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/app/FastqUtil <p>Procedure:</p> <ul style="list-style-type: none"> • Test FASTQ Utilities service using exemplar reads sets. • Test input options, i.e. read files and SRA accession as input. • Test different processing options, i.e. trim, fastqc, and real alignment to a reference genome. • For each job submitted, verify successful completion of the job, presence of output files, and quality of the results from various processing steps. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/fastq_utilities_service.html • https://www.bv-brc.org/docs/tutorial/fastq_utilities/fastq_utilities.html 	ARW, MS
V1	Genome Browser (JBrowse) Tool	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Genome/224914.11#view_tab=browser&loc=NC_003317%3A1..228800&tracks=refseqs%2CPATRICGenes%2CRefSeqGenes&highlight= <p>Procedure:</p> <ul style="list-style-type: none"> • Test the linear Genome Browser using an example bacterial and viral genome. • Test genome display. • Test zoom and pan. • Test feature flyovers/links. • Show track show/hide <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_genome/genome_browser.html 	ARW, ZW
V2	Circular Genome Browser	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/Genome/224914.11#view_tab=circular <p>Procedure:</p>	ARW

		<ul style="list-style-type: none"> • Test the circular Genome Browser using an example bacterial genome. • Test genome display. • Turn off tracks. • Change track colors. • Add custom track. • Upload your own data. • Test feature flyovers/links. • Test track show/hide. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/organisms_genome/genome_browser.html 	
V3	Proteome Comparison Viewer	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/workspace/public/BVBRC@patricbrc.org/BVBRC%20Tests/Proteome%20Comparison/Brucella%20Representative%20Genomes%20Comparison <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Proteome Comparison Viewer using an example set of bacterial genomes. • Test proteome viewer. • Test sequence identity colors. • Test links. • Test download. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/proteome_comparison_service.html 	ARW
V4	Genome Alignment Viewer	<p>URL:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/view/GenomeAlignment/ARWattam@patricbrc.org/BV-BRC%20Workshop/Genome%20Alignment/Brucella/.Brucella%20genome%20group%20alignment%20alpha%20test/alignment.json <p>Procedure:</p> <ul style="list-style-type: none"> • Test the Genome Alignment Viewer using an example set of bacterial genomes. • Test zoom. • Test ordering. • Test flyovers and links. <p>References:</p> <ul style="list-style-type: none"> • https://www.bv-brc.org/docs/quick_references/services/proteome_comparison_service.html 	ARW

V5	Heatmap	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/view/TranscriptomicsExperiment/?&wsExpId=/BV/BRC@patricbrc.org/BVBRC%20Tests/RNA-seq%20Analysis/.Abaumannii%20-%20Tuxedo%20-%20DE2/Abaumannii%20-%20Tuxedo%20-%20DE2_diffexp <p>Procedure:</p> <ul style="list-style-type: none"> Test the Heatmap using an example expression data set. Test zoom. Test filter (genome, keyword, log ratio, z-score). Test row/column rearrangement. Test clustering. Test show significant/all genes. Test flyovers and links. Test area selection and corresponding data access <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/services/rna_seq_analysis_service.html 	ARW
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Workspace

	Component	Test Description	Responsibility
W1	Workspace	<p>URL:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/, then login using the “Sign In” button on the top right of the page. Choose the Workspaces/Home menu option (URL to personal workspace is dependent on username) <p>Procedure:</p> <ul style="list-style-type: none"> Test storing, retrieval, and access of datasets Test creation and usage of folders Test upload of sequence data and associated meta-data using test files Test creation, deletion, copy, move and set operations (union, intersect, subtract) Test sharing with other users and making public Test if public workspaces are available to public <p>References:</p> <ul style="list-style-type: none"> https://www.bv-brc.org/docs/quick_references/workspace_groups_upload.html 	IS, ARW

