Mapping Load

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1 Purpose of Document

To describe the Mapping Load product.

2 Overview

The purpose of the Mapping Load is to:

- 1. To load new or update existing experiment records (one per chromosome) into MGI Mapping tables and set these attributes:
 - Mapping Reference (J:)
 - Experiment Type
 - Gene
 - Assay Type
- 2. To assign MGI Accession IDs to the new experiments.
- 3. To update a Gene's Chromosome and/or Band.

3 Inputs

- 1. Reference (J:)
- 2. Experiment Type
- 3. A tab-delimited file in the following format (no column headings):

Table 1: Input File

Field	Name	Example	Туре	Max Length
1*	MGI Accession ID	MGI:12345	string	30
2*	Chromosome	1	string	2
3*	Update Marker Chromosome?	yes no	string	3
4	Band	A1	string	20

Field	Name	Example	Туре	Max Length
5*	Mapping Assay	assembly	string	80
6	Description	A123456	string	255

Table 1: Input File

Fields with "*" require a non-null value. Other fields must be present but can be blank.

Field 2:

Any mouse chromosome

4 Outputs

- 1. 5 BCP files (inputfile name is prepended to each bcp file name):
 - MLD_Expts.bcp
 - MLD_Marker.bcp
 - MLD_Expt_Marker.bcp
 - MLD_Notes.bcp
 - ACC_Accession.bcp
- 2. Diagnostics file (inputfile.MMDDYYYY.diagnostics) that includes all SQL commands issued by the load.
- 3. Error file (inputfile.MMDDYYYY.error) that includes any data validation errors.

5 Processing

- 1. Verify Mode.
 - if mode = full: delete existing experiment records; create new experiments
 - if mode = incremental: append data to existing experiments
 - if mode = preview: set "DEBUG" to True
- 2. Verify the J:is valid.

If the verification fails, write the error to the error file and skip the record.

- 3. Create the master Experiment records and Mapping Accession IDs (MLD_Expts, ACC_Accession).
- 4. For each line in the input file:
 - Verify the MGI Accession ID is valid (not null).
 If the verification fails, write the error to the error file and skip the record.
 - 2. Verify the Mapping Assay is valid (not null).

If the verification fails, write the error to the error file and skip the record.

- 3. Verify the Chromosome is valid (not null).

 If the verification fails, write the error to the error file and skip the record.
- 4. Create the MLD bcp records (MLD_Marker, MLD_Expt_Marker).
- 5. Load the bcp files into the database.

6 Usage

To execute:

Python Program: mappingload.py

Parameters:

- -S server
- -D mgd database
- -U user
- -P password file
- -M mode (*full*, *incremental* or *preview*)
- -I input file
- -R reference (J: in format ####)
- -E experiment type ("TEXT-Physical Mapping")