LANDIS-II Output Biomass Reclassification v3.0

Extension User Guide

Robert M. Scheller, North Carolina State University

Last Revised: July 31, 2018

# Table of Contents

[1 Introduction 2](#_Toc520790561)

[1.1 Major Versions 2](#_Toc520790562)

[1.1.1 Version 3.0 (August 2018) 2](#_Toc520790563)

[1.1.2 Version 2.1 (June 2017) 2](#_Toc520790564)

[1.1.3 Version 2.0 (June 2012) 2](#_Toc520790565)

[1.2 Minor Versions 3](#_Toc520790566)

[1.2.1 Version 2.1.1 (May 2018) 3](#_Toc520790567)

[1.3 Acknowledgements 3](#_Toc520790568)

[2 Input File 4](#_Toc520790569)

[2.1 LandisData 4](#_Toc520790570)

[2.2 Timestep 4](#_Toc520790571)

[2.3 Reclass Map Descriptions 4](#_Toc520790572)

[2.4 Map Names 5](#_Toc520790573)

# Introduction

This document describes the **Output Biomass Reclassification**  extension for the LANDIS-II model. Readers should read the *LANDIS-II Model User Guide* prior to reading this document. Other output extensions produce biomass output data (see *LANDIS-II Biomass Output User Guide*).

The output extension described herein uses cohort information at each site to classify every active site into a forest type defined by the User (see below). A dominance value is calculated for each species. The dominance value will be equal to the total biomass (B) (Mg/ha) for the species or [B \* -1], as determined by the User.

Each forest type has an associated group of species. The species dominance values are summed for each forest type to which they are assigned. Some species may have a negative dominance for a given forest type, as determined by the User. A site is assigned to (given the value of) the forest type with the highest total dominance value.

## Major Versions

### Version 3.0 (August 2018)

Output Biomass Reclassification is now compatible with Core v7.

### Version 2.1 (June 2017)

Added compatibility with the Metadata library. The Metadata Library outputs metadata for all model outputs, allowing compatibility with visualization tools.

Added compatibility with succession extensions that support the cohort interfaces from Biomass Cohorts through the new dependency on the Biomass Library. Any succession extension that uses a cohort structure that supports the interfaces from Biomass Cohorts should be able to be compatible with this extension.

### Version 2.0 (June 2012)

Output Biomass Reclassification is now compatible with Core v6.0.

## Minor Versions

### Version 2.1.1 (May 2018)

Updated cohort biomass library.

## Acknowledgements

Funding for the development of LANDIS-II has been provided by the North Central Research Station (Rhinelander, Wisconsin) of the U.S. Forest Service. Valuable contributions to the development of the model and extensions were made by Brian R. Sturtevant, Eric J. Gustafson, and David J. Mladenoff.

# Input File

The input parameters for this extension are specified in one input file. This text file must comply with the general format requirements described in section 3.1 Text Input Files in the *LANDIS-II Model User Guide*.

## LandisData

The first parameter is the title of the input file:

LandisData “Reclass Biomass Output”

## Timestep

The second parameter is the time step in years. For example:

Timestep 15

## Reclass Map Descriptions

The next suite of parameters defines the desired reclassification(s) and must be preceded by the keyword ReclassMaps. The input is a table or tables describing the name of the reclassification, the forest types for the reclassification, and the species within (or excluded from) each forest type (Table 1).

|  |  |  |
| --- | --- | --- |
| Table 1. |  |  |
| Parameter | Data type | Example |
| Reclassification Name | string | Reclass1 |
| Forest Type | string | MapleHardwood |
| Species | string | acersacc -pinubank |

First, the Reclassification name must be given, followed by the symbol ‘->’. Do not use spaces in the reclassification name. Immediately following is the first forest type, such as NorthernPine or “Southern Oak” (quotes required if spaces used). Each subsequent forest type is listed on a separate line. Following the name of each forest type is a list of species to be included or excluded. If a species should contribute to the dominance value of a forest type, list the species name. If a species should be subtracted from a forest type, list the species name preceded by a ‘-‘ (negative) sign. **Not all species need be included and a species may be listed in more than one forest type.** An example reclassification:

ReclassMaps

>> Map Name Forest Type Species

>> -------- ------------- -------

reclass1 -> MapleHardwood acersacc betualle -pinubank

NorthernPines pinubank pinuresi pinustro

Oaks querelli querrubr

If a site can not be classified, or is unforested, the default value is zero (0). Otherwise, the map numbering will follow the order of the forest types (e.g., MapleHardwood = 1, NorthernPines = 2, Other = 3).

**Note: If the reclassification calculates equal values for more than one forest type (a tie), then the forest type listed first among equals will be assigned to that cell.**

Multiple reclassifications can be defined. Simply create a new reclassification name with forest types and species below the first reclassification defined.

## Map Names

The next parameter, MapFileNames, describes where output maps are placed and their format. The first portion lists the directory where the maps should be places, relative the location of the scenario text file (e.g., reclass/). The second portion includes two variables for creating file names. {reclass-map-name} will be replaced with the reclass map name. {timestep} will be replaced with the output time step. Other characters can be inserted as desired. A meaningful file extension (e.g., .gis) should also be included. For example:

MapFileNames output/bio-{reclass-map-name}-{timestep}.img