

## Sets

- $I$  : set of grid cells, indexed by  $i$ .
- $S = \{A, M, E, O\}$  : set of species.
- $\text{Adj} \subseteq I \times I$  : adjacency relation;  $(i, j) \in \text{Adj}$  if and only if cells  $i$  and  $j$  share a side (4-neighborhood).
- $K_s$  : set of original colonies (territories) of species  $s$ . Each  $k \in K_s$  represents one connected component of originally occupied cells.

## Parameters

- $\text{area}_i \in \mathbb{R}_+$  : area of cell  $i$ .  $\text{area}_i \geq 0.56$  is maximum.
- $\text{orig\_has}_{i,s} \in \{0, 1\}$  : 1 if species  $s$  originally occurs in cell  $i$ .
- $\text{cost}_{i,s}^T \in \mathbb{R}_+$  : adaptation (territory expansion) cost for species  $s$  to occupy cell  $i$ .
- $\text{cost}_i^C \in \mathbb{R}_+$  : corridor creation cost in cell  $i$ .
- $\text{minArea}_s \in \mathbb{R}_+$  : minimum viable total territory area for species  $s$ .
- $\text{meanArea}_s \in \mathbb{R}_+$  : “satisfactory” target area for species  $s$ .
- $\text{maxArea}_s \in \mathbb{R}_+$  : maximum allowed territory area for species  $s$ .
- $\text{minArea}_{k,s} \in \mathbb{R}_+$  : minimum required area for colony  $k \in K_s$ .
- $\text{meanArea}_{k,s} \in \mathbb{R}_+$  : satisfactory target area for colony  $k \in K_s$ .
- $\text{maxArea}_{k,s} \in \mathbb{R}_+$  : maximum allowed area for colony  $k \in K_s$ .
- $\text{conflict}(s, s') \in \{0, 1\}$  : equals 1 if species  $s$  and  $s'$  cannot overlap (e.g.  $M$  with  $E$  or  $O$ ), otherwise 0.
- $\text{suit}_{i,s} \in [0, 1]$  : suitability score of cell  $i$  for species  $s$  (1 = ideal habitat, 0 = unsuitable).
- $\text{priority}_s \in \mathbb{R}_+$  : priority coefficient of species  $s$  (higher means greater conservation priority).
- $L_{\max} \in \mathbb{Z}_+$  : maximum allowed corridor length (in cells).

## Decision Variables

- $T_{i,s} \in \{0,1\}$  : 1 if cell  $i$  belongs to the *final* territory of species  $s$  (original or expanded), 0 otherwise.
- $E_{i,s} \in \{0,1\}$  : 1 if species  $s$  newly expands into cell  $i$  (i.e.  $i$  was not originally occupied), 0 otherwise.
- $C_i \in \{0,1\}$  : 1 if a corridor is built in cell  $i$ , 0 otherwise.
- $\lambda_{(i,j),s} \in \{0,1\}$  : 1 if species  $s$  uses adjacency  $(i,j) \in \text{Adj}$  as a corridor connection between two territories.
- $\text{Reach}_{i,s,k} \in \{0,1\}$  for  $i \in I$ ,  $s \in S$ ,  $k \in K_s$ : 1 if colony  $k$  of species  $s$  can reach cell  $i$  via corridors, 0 otherwise.
- $g_{i,j,s,k} \geq 0$ : corridor flow on arc  $i \rightarrow j$  for colony  $k$  of species  $s$ .
- $\text{Inject}_{s,i,k} \geq 0$ : flow injected at corridor cell  $i$  for colony  $k$  of species  $s$ .
- $use_{i,s} \in \{0,1\}$ : species-level corridor-use; optional.
- $dist_{i,s,k} \in \{0, 1, \dots, L_{\max}\}$  for  $i \in I$ ,  $s \in S$ ,  $k \in K_s$ : distance (in cells) from colony  $k$  of species  $s$  to cell  $i$  via corridors.

## Auxiliary Variables

- $\text{Area}_s = \sum_{i \in I} area_i T_{i,s}$  : total final area of species  $s$ .
- $\text{ExpArea}_s = \sum_{i \in I} area_i E_{i,s}$  : total expanded area for species  $s$ .
- $\text{Area}_{k,s} = \sum_{i \in I} area_i T_{i,s,k}$  : area of colony  $k$  of species  $s$ , where  $T_{i,s,k}$  indicates membership of  $i$  in colony  $k$  (colony-assignment variables introduced in the model section).
- $y_s = \sum_{(i,j) \in \text{Adj}} \lambda_{(i,j),s}$  : number of corridor links used for species  $s$ .
- $n_s = |K_s|$  : number of original colonies of species  $s$ .

## Stage-Related Quantities

- $\text{MinFeasible}_s = minArea_s$  : minimum feasible area requirement for species  $s$  (Stage 1).
- $\text{MeanTarget}_s = meanArea_s - minArea_s$  : area above minimum toward satisfactory target (Stage 2).
- $\text{MaxPotential}_s = maxArea_s - minArea_s$  : maximum possible expansion above minimum (Stage 3).
- $\text{ConnTarget}_s = n_s - 1$  : maximum number of meaningful inter-colony connections for species  $s$  (tree structure).

# 1 Objective function

## 1.1 Stage 1: Minimum viable territory

### Precomputed parameter

- $\rho_{i,s} := \frac{area_i}{cost_{i,s}^T}$  (precompute for each 0 and species; assume  $cost_{i,s}^T > 0$ ).  
*This is a constant parameter used to linearize area-per-cost preference.*

### Stage 1 objective (minimal territory expansion, no priority)

Stage 1 requires that each species attains its minimal territory. To achieve the required minima with best cost-effectiveness and habitat quality, we maximize the sum of per-cell benefit coefficients for newly expanded cells:

$$(Stage\ 1)\quad \max Z^{(1)} = \sum_{s \in S} \sum_{i \in I} \underbrace{suit_{i,s}}_{\text{suitability}} \underbrace{\frac{\rho_{i,s}}{area_i}}_{\text{cost}_{i,s}^T} E_{i,s}$$

subject to Stage 1 feasibility constraints, e.g.:

(a) Minimal area per species:

$$\sum_{i \in I} area_i E_{i,s} \leq R_{\min,s} + A_{\max}^{\text{cell}} \quad \forall s \in S,$$

where  $A_{\max}^{\text{cell}} := \max_{i \in I} area_i$ .

(b) Expansion fe

(constraints ensuring each expanded cell is adjacent / reachable to its original patch)

(c) No-merge and conflict rules:

(constraints preventing merges between original patches of same species and forbidding expansion that violates conflicts)

(d) Binary domains:

$$E_{i,s} \in \{0, 1\}, T_{i,s} \in \{0, 1\}, C_i \in \{0, 1\}, \dots$$

(Note)  $\rho_{i,s}$  and  $suit_{i,s}$  are model parameters; the objective is linear in  $E_{i,s}$ .

## 1.2 Stage 2 objective (satisfactory territory expansion with proportional corridor creation)

**Stage 2 objective (balanced territory & corridors)**

**Auxiliary definitions (per species  $s$ ).**

$$A_s = \sum_{i \in I} area_i T_{i,s} \quad (\text{total final area of species } s),$$

$$A_{1,s} + A_{2,s} = A_s, \quad 0 \leq A_{1,s} \leq n_s \cdot meanArea_s, \quad A_{2,s} \geq 0,$$

$$TB_s \leq A_{1,s} - m_s, \quad TB_s \geq 0, \quad TB_s \leq Rmean_s \quad (\text{territory between-min-and-mean}),$$

choose  $Y_s$  as either: 
$$\begin{cases} Y_{cells,s} = \sum_{i \in I} use_{i,s} & (\text{corridor cells assigned to species } s - \text{easier}), \\ Y_{edges,s} = \sum_{(i,j) \in \text{Adj}} \lambda_{(i,j),s} & (\text{corridor edges used by species } s - \text{preferred}). \end{cases}$$

**Soft proportional constraint (per species  $s$ ).** (If  $n_s \leq 1$  or  $Rmean_s \leq 0$ , omit this constraint for that  $s$  and set deviations to zero.)

$$Y_s \cdot Rmean_s - TB_s \cdot (n_s - 1) = dev_s^+ - dev_s^-, \quad dev_s^+, dev_s^- \geq 0.$$

**Connectivity certificate variables (recall / definitions).**

$$ConnectedCount_s = \sum_{k \in K_s} conn_{s,k},$$

where  $conn_{s,k} \in \{0, 1\}$  indicates colony  $k$  of species  $s$  is certified connected (via corridor flow constraints introduced in the model).

**Connectivity reward scaling parameter. (An example choice.)**

$$\gamma_{conn} = \frac{n_s \cdot meanArea_s - minArea_s}{n_s - 1}$$

**Stage 2 objective (maximize).**

$$\begin{aligned}
 (\text{Stage 2}) \quad \max Z^{(2)} = & \underbrace{\frac{\sum_{s \in S} (A_{1,s} + 0.7 A_{2,s})}{0.56}}_{\text{territory range-reward (full cell units)}} + \gamma_{\text{conn}} \underbrace{\sum_{s \in S} \text{ConnectedCount}_s}_{\text{connectivity reward}} \\
 & - \underbrace{\gamma_{\text{cost}} \left( \sum_{s \in S} \sum_{i \in I} \text{cost}_{i,s}^T E_{i,s} + \sum_{i \in I} \text{cost}_i^C C_i \right)}_{\text{adaptation \& corridor costs}} \\
 & - \underbrace{\gamma_{\text{edge}} \sum_{(i,j) \in \text{Adj}} \sum_{s \in S} \lambda_{(i,j),s}}_{\text{edge penalty (favor tree-like)}} \\
 & - \underbrace{\gamma_{\text{prop}} \sum_{s \in S} (dev_s^+ + dev_s^-)}_{\text{penalize proportion deviation}}
 \end{aligned}$$

### Implementation notes and recommended parameter choices

- **Choice of  $Y_s$ :** for true tree semantics select  $Y_s := Y_{edges,s}$ . If corridor costs and practical construction are cell-based, the simpler option is  $Y_s := Y_{cells,s}$ . Use edges for proportionality to  $n_s - 1$ .
- $\gamma_{\text{conn}}$  should reflect how valuable connecting one colony is relative to adding area. As a baseline, it can be set to  $\gamma_{\text{conn}} \approx$  average per-patch area-equivalent (e.g. mean cell-area).
- $\gamma_{\text{cost}}$  scales monetary costs into the same units as territory reward. If costs are already monetary and territory reward is in area units, normalize by converting one to the other or choose  $\gamma_{\text{cost}}$  to produce desired trade-offs. Suggested equation:  $\gamma_{\text{costs}} = \frac{1}{|I|} \sum_{i \in I} \rho_{i,s} = \frac{1}{|I|} \sum_{i \in I} \frac{\text{cost}_{i,s}}{\text{cost}_{i,s}^T}$ .
- $\gamma_{\text{edge}}$  should be small (e.g. 0.01–0.1 in normalized units) to prefer trees but not overrule connectivity gains.
- $\gamma_{\text{prop}}$  controls how tightly the proportion is enforced: large values  $\Rightarrow$  near-equality, small values  $\Rightarrow$  loose. Start with a moderate value and tune.
- For species with  $n_s \leq 1$  or  $Rmean_s \leq 0$ , set  $dev_s^\pm = 0$  (no proportional constraint) and either fix  $Y_s = 0$  (no corridors) or treat corridor reward as zero for that species.
- Maintain Stage 1 hard constraints (minimal territory) when solving Stage 2 (do not relax them).

### Stage 3 objective (top-up expansion to maxima)

Auxiliary parameters (for Stage 3).

- $Budget_{\text{rem}} \in \mathbb{R}_+$  : remaining monetary budget available for Stage 3 expansions.
- $Rmax_s := n_s k_s - m_s$  : maximum remaining area (in cell-units or area-units) for species  $s$  above the minimum.
- (Optional)  $\delta \in (0, 1]$  : diminishing-return coefficient for expansion beyond the "mean" level (use  $\delta < 1$  to downweight overly large expansions). If not used, set  $\delta = 1$ .

Stage 3 objective (maximize additional expansion benefit).

$$(\text{Stage 3}) \quad \max Z^{(3)} = \sum_{s \in S} \sum_{i \in I} \underbrace{\text{suit}_{i,s}}_{\text{suitability}} \frac{\rho_{i,s}}{\underbrace{\text{area}_i}_{\text{cost}_{i,s}^T}} E_{i,s}^{(3)}$$

where  $E_{i,s}^{(3)}$  are the Stage 3 decision binaries that equal 1 if cell  $i$  is chosen for additional expansion for species  $s$  during Stage 3 (cells already committed in Stages 1–2 should be fixed and excluded from  $E_{i,s}^{(3)}$ ).

Subject to (core) Stage 3 constraints.

(a) Per-species maximum cap:

$$\sum_{i \in I} \text{area}_i E_{i,s}^{(3)} \leq Rmax_s \quad \forall s \in S,$$

(b) Budget constraint:

$$\sum_{s \in S} \sum_{i \in I} \text{cost}_{i,s}^T E_{i,s}^{(3)} + \sum_{i \in I} \text{cost}_i^C C_i^{(3)} \leq Budget_{\text{rem}},$$

(c) Feasibility (adjacency / no-merge / conflict / corridor rules):

(Stage-3 expansions must obey the same adjacency, connectivity and conflict constraints)

(d) Binary domain:

$$E_{i,s}^{(3)} \in \{0, 1\}, \quad C_i^{(3)} \in \{0, 1\}, \dots$$

**Optional: diminish returns beyond mean.** If you wish to reduce marginal benefit for expansions beyond the satisfactory (mean) level, split each species' additional expansion into two parts:

$$A_{1,s}^{(\text{tot})} = \sum_{i \in I} \text{area}_i T_{i,s} \quad (\text{current total after Stages 1–2})$$

$$\text{Let } Exceed_s = \max\{0, A_{1,s}^{(\text{tot})} - n_s \cdot meanArea_s\}$$

and then weight expansions beyond the mean by  $\delta$  in the objective:

$$Z^{(3)} = \sum_s \sum_i suit_{i,s} \rho_{i,s} E_{i,s}^{(3, \leq mean)} + \delta \sum_s \sum_i suit_{i,s} \rho_{i,s} E_{i,s}^{(3, > mean)}$$

(Implementation note: implement the split  $E^{(3, \leq mean)}$  and  $E^{(3, > mean)}$  by comparing cumulative area to  $n_s \cdot meanArea_s$  using standard linearization tricks.)

### Notes.

- In a sequential workflow, variables/decisions already fixed by Stage 1 and Stage 2 should be fixed (constants) during Stage 3; the Stage 3 binary  $E_{i,s}^{(3)}$  should only be available for cells not already selected.
- If corridor additions are permitted in Stage 3, include  $C_i^{(3)}$  and associated corridor feasibility constraints. Often corridors are set in Stage 2 and not expanded further — adapt per your workflow.
- The objective uses  $\rho_{i,s} = area_i / cost_{i,s}^T$  to prefer cells giving most area per monetary unit, adjusted by suitability.
- If you prefer to prioritize particular species in Stage 3, include a multiplicative weight  $w_s$  (priority) inside the summand.

## 2 Constraints

### Constraints — General feasibility & geometry

**Notation reminder (used below).** We keep the naming conventions introduced earlier:

$$\begin{aligned} I, S, \text{Adj}, K_s, \\ T_{i,s} \in \{0, 1\}, E_{i,s} \in \{0, 1\}, C_i \in \{0, 1\}, use_{i,s} \in \{0, 1\}, \\ area_i, cost_{i,s}^T, cost_i^C, suit_{i,s} \in [0, 1], \\ n_s, m_s, \ell_s, k_s, Rmean_s := n_s \ell_s - m_s, \\ \rho_{i,s} := \frac{area_i}{cost_{i,s}^T} \text{ (precomputed),} \\ conflict(s, s') \in \{0, 1\} \text{ indicates pairwise conflict.} \end{aligned}$$

#### 1.1 Variable domains and basic identities

$$T_{i,s} \in \{0, 1\}, \quad E_{i,s} \in \{0, 1\}, \quad C_i \in \{0, 1\}, \quad use_{i,s} \in \{0, 1\}, \quad (1)$$

$$\lambda_{(i,j),s} \in \{0, 1\} \quad \forall (i, j) \in \text{Adj}, s \in S \quad (\text{corridor-edge usage binaries, optional}). \quad (2)$$

Basic preservation of original presence:

$$T_{i,s} \geq orig\_has_{i,s} \quad \forall i \in I, s \in S. \quad (3)$$

Link expanded-only to final membership:

$$E_{i,s} \geq T_{i,s} - orig\_has_{i,s} \quad \forall i, s, \quad (4)$$

$$E_{i,s} \leq T_{i,s}, \quad E_{i,s} \leq 1 - orig\_has_{i,s} \quad \forall i, s. \quad (5)$$

These ensure  $E_{i,s} = 1$  exactly when cell  $i$  is newly added for species  $s$ .

## 1.2 Area aggregates

Total (final) area by species:

$$A_s = \sum_{i \in I} area_i T_{i,s} \quad \forall s \in S. \quad (6)$$

Total expanded area (for cost accounting) by species:

$$ExpArea_s = \sum_{i \in I} area_i E_{i,s} \quad \forall s \in S. \quad (7)$$

## 1.3 Budget / cost accounting (global / stage-level)

Total adaptation (expansion) cost and corridor cost:

$$Cost^{\text{adapt}} = \sum_{s \in S} \sum_{i \in I} cost_{i,s}^T E_{i,s}, \quad (8)$$

$$Cost^{\text{corr}} = \sum_{i \in I} cost_i^C C_i. \quad (9)$$

Example global budget constraint (use stage-specific budget where appropriate):

$$Cost^{\text{adapt}} + Cost^{\text{corr}} \leq Budget_{\text{total}}. \quad (10)$$

## 1.4 Adjacency / geometry precomputation and usage note

We assume the adjacency relation  $\text{Adj}$  is precomputed externally (parameter):  $(i, j) \in \text{Adj}$  iff cells  $i$  and  $j$  share a side (4-neighborhood). Use the set  $\text{Adj}$  in later constraints that require adjacency (expansion adjacency, corridor continuity, flow conservation, etc.).

## 1.5 Corridor logical linking

A species may use a corridor cell only if the cell is built as corridor:

$$use_{i,s} \leq C_i \quad \forall i, s.$$

**Note:** Corridor usage does *not* require territory overlap. Connectivity of a species to a corridor is handled solely through the corridor-patch gateway and flow constraints (Section 3).

## 1.6 Corridor-edge linking (optional)

If using edge binaries  $\lambda_{(i,j),s}$  to represent corridor edges used by species  $s$ , link them to corridor cells:

$$\lambda_{(i,j),s} \leq C_i, \quad \lambda_{(i,j),s} \leq C_j, \quad \lambda_{(i,j),s} \geq C_i + C_j - 1, \quad (11)$$

and if per-species edge usage must be tied to species accessibility:

$$\lambda_{(i,j),s} \leq use_{i,s}, \quad \lambda_{(i,j),s} \leq use_{j,s}. \quad (12)$$

## 1.7 Suitability-based hard exclusions (optional)

If you want to **forbid** selection of cells that are unsuitable for a species (hard exclusion), define a binary parameter  $allowed_{i,s} \in \{0, 1\}$  by preprocessing:

$$allowed_{i,s} = \begin{cases} 1 & \text{if } suit_{i,s} \geq \tau_s, \\ 0 & \text{otherwise,} \end{cases}$$

for some threshold  $\tau_s$ . Then impose:

$$T_{i,s} \leq allowed_{i,s}, \quad E_{i,s} \leq allowed_{i,s} \quad \forall i, s. \quad (13)$$

If you prefer soft exclusion, instead multiply suitability into the objective and do not enforce (13).

## 1.8 Conflict / predator-prey overlap protection (do not worsen initial conflicts)

Let  $P$  be the set of unordered species-pairs with conflicts:

$$P = \{(s, s') \in S \times S : conflict(s, s') = 1, s < s'\}.$$

Precompute the original overlap indicator for each cell and pair:

$$orig\_overlap_{i,s,s'} := orig\_has_{i,s} \cdot orig\_has_{i,s'}$$

(this is a parameter equal to 1 if both species originally co-occur in cell  $i$ , 0 otherwise).

Introduce linearized overlap binaries for final map:

$$O_{i,s,s'} \in \{0, 1\}, \quad O_{i,s,s'} \leq T_{i,s}, \quad O_{i,s,s'} \leq T_{i,s'}, \quad O_{i,s,s'} \geq T_{i,s} + T_{i,s'} - 1,$$

for all  $(s, s') \in P$ ,  $i \in I$ . These enforce  $O_{i,s,s'} = 1$  iff both species occupy cell  $i$  in the final plan.

Constraint: do not increase the number of overlapping cells relative to the original map:

$$\sum_{i \in I} O_{i,s,s'} \leq \sum_{i \in I} orig\_overlap_{i,s,s'} \quad \forall (s, s') \in P. \quad (14)$$

This enforces your rule: "If there is any overlap between initial conflicting colonies at the beginning, we ignore it and expand in a way that does not worsen the situation."

## 1.9 Corridor exclusivity for conflicting species

Prevent conflicting species from sharing the *same* corridor cell (if required):

$$use_{i,s} + use_{i,s'} \leq 1 \quad \forall (s, s') \in P, i \in I.$$

## 1.10 Bookkeeping / useful derived quantities

Number of corridor cells usable by species  $s$ :

$$Y_{cells,s} = \sum_{i \in I} use_{i,s}.$$

Number of corridor edges used by species  $s$  (if edges used):

$$Y_{edges,s} = \sum_{(i,j) \in \text{Adj}} \lambda_{(i,j),s}.$$

## 1.11 Soft enforcement note for tree-like corridors

The strict constraint “no cycles; corridor count  $\leq n_s - 1$ ” can be computationally heavy. We therefore recommend implementing this concept as a *soft* preference:

$$\text{penalize } \sum_{(i,j) \in \text{Adj}} \sum_{s \in S} \lambda_{(i,j),s} \text{ in the objective (small weight)}$$

rather than enforcing a combinatorial cycle-elimination constraint. If you prefer a hard tree constraint later, add connectivity-flow formulations with cut/MTZ-style subtour elimination (but be aware of model size).

# Constraints — Territory formation & expansion

**Notation reminder (new items used here).**

$T_{i,s,k} \in \{0, 1\}$  for  $k \in K_s$  : cell  $i$  is assigned to final territory of original patch  $k$ ,  
 $patchOf_{i,s}$  : parameter mapping original cell  $i$  to its original patch  $k \in K_s$   
(defined only when  $orig\_has_{i,s} = 1$ ),  
 $f_{i,j,s,k} \geq 0$  : continuous flow on directed arc  $(i \rightarrow j)$  for patch  $k$  of species  $s$ ,  
 $U^{\text{flow}}$  : a large upper bound for flow (suggest  $U^{\text{flow}} = |I|$ ).

## 2.1 Patch assignment and no-merge

**(a) Decompose species membership by original patch.**

$$T_{i,s} = \sum_{k \in K_s} T_{i,s,k} \quad \forall i \in I, s \in S, \tag{15}$$

with the interpretation that every final membership  $T_{i,s}$  is assigned to exactly one original patch of the same species.

**(b) Prevent a cell being assigned to more than one original patch of the same species (no-merge).**

$$\sum_{k \in K_s} T_{i,s,k} \leq 1 \quad \forall i \in I, s \in S. \quad (16)$$

**(c) Fix original patch assignments for original cells.** For each original cell  $i$  and species  $s$  let  $k = \text{patchOf}_{i,s}$  be its original patch id (parameter). Then:

$$T_{i,s,k} = \text{orig\_has}_{i,s} \quad \forall i \in I, s \in S \text{ with } \text{orig\_has}_{i,s} = 1. \quad (17)$$

(If a species does not originally occupy cell  $i$ , there is no enforced equality in (17).)

## 2.2 Per-patch area accounting and A1/A2 split (for Stage 2 reward)

**(a) Patch area variables.**

$$A_{k,s} = \sum_{i \in I} \text{area}_i T_{i,s,k} \quad \forall s \in S, k \in K_s.$$

**(b) Per-patch A1/A2 split that implements "reward = 1 up to mean, 0.7 beyond".** For every original patch  $k \in K_s$  we define:

$$A1_{k,s} + A2_{k,s} = A_{k,s}, \quad (18)$$

$$0 \leq A1_{k,s} \leq \text{meanArea}_s, \quad A2_{k,s} \geq 0. \quad (19)$$

(At optimum,  $A1_{k,s} = \min\{A_{k,s}, \text{meanArea}_s\}$  and  $A2_{k,s} = \max\{0, A_{k,s} - \text{meanArea}_s\}$ .)

**(c) Species-level aggregates (useful in objectives).**

$$A_s = \sum_{k \in K_s} A_{k,s}, \quad A1_s = \sum_{k \in K_s} A1_{k,s}, \quad A2_s = \sum_{k \in K_s} A2_{k,s}. \quad (20)$$

## 2.3 Minimal / maximum area per patch / species (Stage 1 & 3)

**(a) Per-species Stage 1 minimal requirement (hard):**

$$A_s \geq n_s \cdot \text{meanArea}_s - m_s = R\text{mean}_s \quad \forall s \in S, \quad (21)$$

or, if you want to enforce per-patch minima instead, require:

$$A_{k,s} \geq \text{minArea}_{k,s} \quad \forall s \in S, k \in K_s. \quad (22)$$

**(b) Per-species maximum cap (Stage 3 constraint):**

$$A_s \leq n_s \cdot k_s - m_s \quad \forall s \in S. \quad (23)$$

## 2.4 Adjacency-based expansion feasibility (local rule)

New expansions must be adjacent to existing territory (original or newly expanded). For each cell  $i$  and species  $s$ :

$$E_{i,s} \leq \sum_{j: (i,j) \in \text{Adj}} T_{j,s}. \quad (24)$$

This enforces: if  $E_{i,s} = 1$  then at least one neighbor  $j$  has  $T_{j,s} = 1$  (original or expanded) ensuring local growth only.

## 2.5 Flow-based connectivity certification for each original patch (guarantees contiguity)

Rationale: to ensure that all cells assigned to a particular original patch  $k \in K_s$  form a connected region reachable from the chosen patch root  $r = \text{patchroot}_{s,k}$ , we use a single-commodity flow construction.

### Definitions / bounds.

$f_{i,j,s,k} \geq 0 \quad \forall (i,j) \in \text{Adj}, s \in S, k \in K_s$  (directed flow variable on edge  $i \rightarrow j$ ), choose  $U^{\text{flow}} = |I|$  (or  $\sum_i 1$ ) as an upper bound.

**Edge capacity / activation linking (optional)** Optionally, introduce binary  $Z_{(i,j),s,k} \in \{0, 1\}$  to indicate edge activation for patch  $k$ . Link flows to activations if desired:

$$f_{i,j,s,k} \leq U^{\text{flow}} Z_{(i,j),s,k}, \quad Z_{(i,j),s,k} \leq T_{i,s,k}, \quad Z_{(i,j),s,k} \leq T_{j,s,k}. \quad (25)$$

(Using  $Z$ -binaries strengthens the formulation but increases binary count; the formulation remains valid without  $Z$  by using bounds on  $f$  tied to  $T$ .)

**Flow conservation / demand definition.** Let  $r = \text{patchroot}_{s,k}$ . For each node  $i \in I$  and each  $s, k$ :

$$\sum_{j: (j,i) \in \text{Adj}} f_{j,i,s,k} - \sum_{j: (i,j) \in \text{Adj}} f_{i,j,s,k} = d_{i,s,k}, \quad (26)$$

where the right-hand side  $d_{i,s,k}$  is defined as:

$$d_{i,s,k} = \begin{cases} - \sum_{i' \in I \setminus \{r\}} T_{i',s,k} & \text{if } i = r \quad (\text{root supplies total demand}), \\ T_{i,s,k} & \text{if } i \neq r \quad (\text{each selected non-root node demands 1 unit}). \end{cases}$$

This enforces that for each selected node  $i \neq r$  exactly one unit of flow must reach it from the root  $r$ .

**Flow-variable bounding by selection (to prevent flow through unselected nodes).** For every arc  $(i, j)$  and patch  $k$ :

$$f_{i,j,s,k} \leq U^{\text{flow}} T_{i,s,k}, \quad (27)$$

$$f_{i,j,s,k} \leq U^{\text{flow}} T_{j,s,k}. \quad (28)$$

Thus flows can be nonzero only on arcs whose both endpoints are selected for the same patch (either enforced via these bounds or via the optional  $Z$  variables in (25)).

**Interpretation.** If a node  $i$  is selected for patch  $k$  then the flow constraints force a path of positive flow from the root  $r$  to  $i$  using only nodes/edges where  $T_{*,s,k} = 1$ . This certifies contiguity of the final patch  $k$ .

## 2.6 Species absence and preventing expansions for absent species

If a species  $s$  has no original patches ( $n_s = 0$ ), disallow expansion and territory:

$$T_{i,s} = 0, \quad E_{i,s} = 0 \quad \forall i \in I, \text{ if } n_s = 0. \quad (29)$$

## 2.7 Symmetry-breaking (optional, helps solver)

To reduce symmetric solutions among interchangeable patches, you may impose ordering on patch indices using e.g. root-cell ids or patch areas:

$$A_{k,s} \geq A_{k+1,s} \quad \forall s \in S, k \text{ ordered by patch id}$$

(Use only if necessary — adds constraints but can speed up branch-and-bound.)

# 3 Corridor Design and Connectivity Constraints

This section introduces binary corridor-cell variables, species-specific usage variables, patch-corridor overlap indicators, and the patch-level connectivity system based on the reachability variables  $\text{Reach}_{i,s,k}$ . Connectivity is certified deterministically by propagating reachability through activated corridor edges.

## 3.1 Notation (new variables)

$C_i \in \{0, 1\}$	corridor built in cell $i$ ,
$use_{i,s} \in \{0, 1\}$	species $s$ may use corridor cell $i$ ,
$\lambda_{(i,j),s} \in \{0, 1\}$	corridor edge $(i, j) \in Adj$ usable by species $s$ ,
$Overlap_{i,s,k} \in \{0, 1\}$	corridor cell $i$ overlaps patch $k$ of species $s$ ,
$Gateway_{s,k} \in \{0, 1\}$	patch $k$ of species $s$ has at least one corridor gateway,
$Reach_{i,s,k} \in \{0, 1\}$	cell $i$ is corridor-reachable from patch $k$ of species $s$ .

### 3.2 Corridor Cell Usage

A species may use a corridor cell only if the corridor cell exists:

$$use_{i,s} \leq C_i \quad \forall i \in I, s \in S.$$

Optionally, require that species may use a corridor cell only if it overlaps their territory (unless explicitly allowed):

$$use_{i,s} \leq T_{i,s} + Allow_{i,s}, \quad Allow_{i,s} \in \{0, 1\}.$$

### 3.3 Corridor Edges

Corridor edges may be used only if both endpoint corridor cells are built:

$$\lambda_{(i,j),s} \leq C_i, \quad \lambda_{(i,j),s} \leq C_j, \quad (30)$$

$$\lambda_{(i,j),s} \geq C_i + C_j - 1, \quad \forall (i,j) \in Adj, s \in S. \quad (31)$$

Species may use an edge only if they may use both endpoint cells:

$$\lambda_{(i,j),s} \leq use_{i,s}, \quad \lambda_{(i,j),s} \leq use_{j,s} \quad \forall (i,j) \in Adj, s \in S. \quad (32)$$

### 3.4 Corridor–Patch Overlap and Gateways

Corridor cell  $i$  overlaps patch  $k$  of species  $s$  if and only if both conditions hold:

$$Overlap_{i,s,k} \leq C_i, \quad (33)$$

$$Overlap_{i,s,k} \leq T_{i,s,k}, \quad (34)$$

$$Overlap_{i,s,k} \geq C_i + T_{i,s,k} - 1, \quad \forall i \in I, s \in S, k \in K_s. \quad (35)$$

A patch has a corridor gateway if at least one corridor cell overlaps it:

$$\sum_{i \in I} Overlap_{i,s,k} \geq Gateway_{s,k}, \quad (36)$$

$$\sum_{i \in I} Overlap_{i,s,k} \leq U^{gw} Gateway_{s,k}, \quad (37)$$

where  $U^{gw}$  is an upper bound on possible gateway cells (e.g. patch area).

### 3.5 Corridor Reachability and Patch Connectivity

**Initial reachability from a patch.** Any gateway cell of patch  $k$  is initially reachable from patch  $k$ :

$$Reach_{i,s,k} \geq Overlap_{i,s,k} \quad \forall i, s, k.$$

**Propagation of reachability through corridor edges.** Reachability spreads from  $i$  to  $j$  only if the corridor edge  $(i, j)$  is active and usable by species  $s$ :

$$Reach_{j,s,k} \geq Reach_{i,s,k} + \lambda_{(i,j),s} - 1, \quad (38)$$

$$Reach_{i,s,k} \geq Reach_{j,s,k} + \lambda_{(i,j),s} - 1, \quad \forall (i, j) \in Adj, s \in S, k \in K_s. \quad (39)$$

**Patch-to-root connectivity certification.** Let  $k_r$  denote the chosen root patch of species  $s$ . Patch  $k$  is considered corridor-connected to the root if a corridor cell that overlaps patch  $k$  is reachable from any corridor cell overlapping the root:

$$conn_{s,k} \leq \sum_{i \in I} Reach_{i,s,k_r} \cdot Overlap_{i,s,k}, \quad \forall s \in S, k \in K_s. \quad (40)$$

Additionally, a patch cannot be connected unless it has a gateway:

$$conn_{s,k} \leq Gateway_{s,k} \quad \forall s, k. \quad (41)$$

### 3.6 Corridor Exclusivity for Predator–Prey Species

For each predator–prey pair  $(s, s') \in P$ , disallow simultaneous use of a corridor cell:

$$use_{i,s} + use_{i,s'} \leq 1 \quad \forall i \in I, (s, s') \in P. \quad (42)$$

### 3.7 Corridor Construction Cost

Corridor construction cost is accumulated as:

$$Cost^{corr} = \sum_{i \in I} cost_i^C C_i. \quad (43)$$

### 3.8 Soft Penalty for Excessive Corridor Edges

To encourage sparse, tree-like corridor networks while still allowing cycles:

$$Penalty_{edges} = \gamma_{edge} \sum_{s \in S} \sum_{(i,j) \in Adj} \lambda_{(i,j),s}. \quad (44)$$

### 3.9 Gateways have distance 0 (if Overlap exists).

This forces any corridor cell that overlaps a patch  $k$  to be a root (distance 0) for propagation.

$$dist_{i,s,k} \leq L_{\max}(1 - Overlap_{i,s,k}) + 0 \quad \forall i, s, k.$$

Explanation: when  $Overlap_{i,s,k} = 1$ ,  $dist_{i,s,k} \leq 0 \Rightarrow dist = 0$ .

### 3.10 Upper bound for all dist variables (redundant but explicit)

$$dist_{i,s,k} \leq L_{\max} \quad \forall i, s, k.$$

### 3.11 Propagation constraint across an active corridor edge:

if edge  $(i, j)$  is active for species  $s$ , then  $dist_j \leq dist_i + 1$ . Use  $Big-M = L_{\max}$  to deactivate when edge not active.

$$dist_{j,s,k} \leq dist_{i,s,k} + 1 + L_{\max} (1 - \lambda_{(i,j),s}) \quad \forall (i, j) \in Adj, s \in S, k \in K_s.$$

### 3.12 Symmetric direction (ensure propagation both ways if needed):

$$dist_{i,s,k} \leq dist_{j,s,k} + 1 + L_{\max} \cdot (1 - \lambda_{(i,j),s}) \quad \forall (i, j) \in Adj, s \in S, k \in K_s.$$

### 3.13 Link reachability to distance:

if cell  $i$  is reachable ( $Reach_{i,s,k} = 1$ ) from patch  $k$ , its distance must be  $\leq L_{\max}$ . (The `dist` variable is already bounded by  $L_{\max}$ , but this constraint ties Reach to dist.)

$$dist_{i,s,k} \leq L_{\max} - (1 - Reach_{i,s,k}) \cdot L_{\max} \quad \forall i, s, k.$$

This constraint is equivalent to:  $Reach_{i,s,k} = 1 \Rightarrow dist_{i,s,k} \leq L_{\max}$ . (When  $Reach = 0$  the  $RHS = 0$ , so it does not enforce anything active beyond nonnegativity; the dist variable's domain still applies.)

### 3.14 (Optional) force Reach to be consistent with finite distances:

if  $dist \leq L_{\max}$  then Reach may be 1; otherwise Reach must be 0. We enforce the implication:  $dist_{i,s,k} \leq L_{\max} * Reach_{i,s,k}$ .

$$dist_{i,s,k} \leq L_{\max} Reach_{i,s,k} \quad \forall i, s, k.$$

## Constraints — Predator-prey and species interaction

**Interpretation and scope.** We assume a set of predator-prey pairs  $P_{\text{pred}} \subseteq S \times S$  where the first element is the predator (e.g.  $M$ ) and the second is the prey (e.g.  $E$  or  $O$ ). For all  $(p, r) \in P_{\text{pred}}$  the model must ensure that final planning does not worsen predator-prey co-occurrence at the cell level: no new cell-level overlaps between predator  $p$  and prey  $r$  may be created (though existing overlaps that are present in the original map are tolerated).

## 4.1 Cell-level no-new-overlap for predator–prey pairs (strict)

Precompute original co-occurrence parameter:

$$\text{orig\_overlap}_{i,p,r} = \text{orig\_has}_{i,p} \cdot \text{orig\_has}_{i,r}, \quad \forall i \in I, (p, r) \in P_{\text{pred}}.$$

Linearized final overlap indicator (repeated for clarity):

$$\begin{aligned} O_{i,p,r} &\in \{0, 1\}, \\ O_{i,p,r} &\leq T_{i,p}, \quad O_{i,p,r} \leq T_{i,r}, \\ O_{i,p,r} &\geq T_{i,p} + T_{i,r} - 1. \end{aligned}$$

**Strict no-new-overlap constraint (per cell):**

$$O_{i,p,r} \leq \text{orig\_overlap}_{i,p,r} \quad \forall i \in I, (p, r) \in P_{\text{pred}}. \quad (45)$$

This enforces that a predator  $p$  and prey  $r$  can co-occur in cell  $i$  in the final plan only if they already co-occurred there originally.

## 4.2 No expansion of predator into prey-occupied cells (alternative / redundant)

An equivalent linear form (redundant if (45) is used) that explicitly prevents the predator from occupying any cell that the prey occupies unless originally co-occupied:

$$T_{i,p} \leq 1 - (T_{i,r} - \text{orig\_has}_{i,p} \cdot \text{orig\_has}_{i,r}) \quad \forall i, (p, r) \in P_{\text{pred}}.$$

This is stronger-looking in algebraic form but logically equivalent to the cell-level overlap prohibition above.

## 4.3 Corridor exclusivity reiterated

Predator and prey should not be permitted to use the same corridor cells:

$$\text{use}_{i,p} + \text{use}_{i,r} \leq 1 \quad \forall i \in I, (p, r) \in P_{\text{pred}}.$$

This was already included in general constraints but is repeated here for emphasis.

## 4.4 Prevent predator expansions that simultaneously overlap original and newly-expanded prey territory (optional special rule)

You requested a special rule that prevents predator  $p$  from expanding into cells that would cause it to intersect both original and newly-expanded territories of prey  $r$ . One linear implementation is:

For each cell  $i$ :  $E_{i,p} + orig\_has_{i,r} + E_{i,r} \leq 2$ .

Explanation: If both  $orig\_has_{i,r} = 1$  (prey originally present) and  $E_{i,r} = 1$  (prey newly expanded into the cell), then  $orig\_has_{i,r} + E_{i,r} \geq 1$ ; the inequality above prevents  $E_{i,p} = 1$  when both are present in a way that would make predator expansion coincide with prey original+expanded presence. This rule is optional and should be used only if the biological justification is necessary — otherwise the strict per-cell no-new-overlap (Section 4.1) is sufficient.

#### 4.5 Optional buffer-zone (neighborhood) constraints for prey protection

If you wish to enforce a one-cell buffer (no predator expansion into any cell adjacent to a prey's final territory), include this neighbor-based linear constraint for all  $(p, r) \in P_{\text{pred}}$ :

$$E_{i,p} \leq 1 - \max_{j: (i,j) \in \text{Adj}} T_{j,r}, \quad \forall i \in I.$$

Linearized as:

$$E_{i,p} \leq 1 - B_{i,r}, \quad B_{i,r} \geq T_{j,r} \quad \forall j : (i,j) \in \text{Adj}, \quad B_{i,r} \in \{0, 1\}.$$

This forces predator  $p$  to avoid any cell adjacent to the prey's final territory. Use only if buffer protection is required.

#### 4.6 Remarks on alternatives and model behavior

- Using per-cell no-new-overlap (45) is conservative and safe: predators never newly co-occur with prey in any cell. If you prefer a more flexible policy (e.g. allow some controlled new overlaps as long as total overlap does not increase), replace the per-cell constraint by the earlier sum-level constraint:

$$\sum_{i \in I} O_{i,p,r} \leq \sum_{i \in I} orig\_overlap_{i,p,r}.$$

- Corridor exclusivity is required to prevent predator and prey from using the same corridor cells and potentially transmitting predation through the same route. This is enforced by the 'use' variables constraint above.
- If some initial overlaps exist ( $orig\_overlap = 1$ ), the model will not remove them (we preserve originals) but will not create any new overlapping cells involving predator-prey pairs.