

HCB

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Title Human Cultural Boundaries

Version 0.0.0

Description Creates seed populations with phoneme inventories that can grow, migrate, and create off-shoot populations. Phoneme inventories mutate when populations establish a new territory.

License What license it uses

Encoding UTF-8

LazyData true

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R topics documented:

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| | |
|-----------------|--------------------------|
| AddBeringStrait | <i>Add Bering Strait</i> |
|-----------------|--------------------------|

Description

Removes connections at the FirstStep stage of the Local structure to create "barriers" between cells. Bering Strait Barriers are designed to create structures similar to the Bering Strait entering North America, going through Central America, then opening up into South America.

Usage

AddBeringStrait(P, firstStep)

Arguments

| | |
|-----------|--|
| P | A list of parameters. |
| firstStep | The local directions created by OneStepDirections(). |

| | |
|------------|--------------------|
| AddSegment | <i>Add Segment</i> |
|------------|--------------------|

Description

Creates lines to show the Bering Straight borders.

Usage

```
AddSegment(P, a, b, top = FALSE)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|----------|-------------------------------|
| AddShift | <i>Add or Shift a Phoneme</i> |
|----------|-------------------------------|

Description

Allows a language to either gain a new phoneme or shift an existing phoneme to better match another population's phoneme inventory.

Usage

```
AddShift(P, targetLanguage, languages, local, phonemeRelatedness, index)
```

Arguments

| | |
|----------------|--|
| P | A list of parameters. |
| targetLanguage | The target language to be modified if possible. |
| languages | All languages |
| local | The local territories data structure. |
| index | The target territory whose language may change. |
| phonemeProbab | The probability of gaining each phoneme in the population. |

| | |
|------------------|---------------------------|
| AddSnakeBarriers | <i>Add Snake Barriers</i> |
|------------------|---------------------------|

Description

Removes connections at the FirstStep stage of the Local structure to create "barriers" between cells. Snake Barriers are lines with length and spacing defined by the parameters. The barriers jut out from the east and west walls, alternating east, west, east, west. This creates a snaking, zig-zag pattern, hence the name.

Usage

```
AddSnakeBarriers(P, firstStep)
```

Arguments

| | |
|-----------|--|
| P | A list of parameters. |
| firstStep | The local directions created by OneStepDirections(). |

| | |
|------------------|---------------------------|
| BeringStraitPlot | <i>Bering Strait Plot</i> |
|------------------|---------------------------|

Description

Creates a plot that shows the Bring strait boundaries.

Usage

```
BeringStraitPlot(P, Data)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|--------------------|----------------------------|
| CardinalDirections | <i>Cardinal Directions</i> |
|--------------------|----------------------------|

Description

Calculates the territory indicies of locations around a target territory (also used for phoneme relat-
edness in the same way).

Usage

```
CardinalDirections(target, R, start, round, South, North, East, West, SE,  
NE, SW, NW)
```

Arguments

| | |
|--------|--|
| target | The territory around which to get local territoies. |
| R | The number of rows. |
| start | How much to offset numbers (for phoneme structures). |
| round | Whether to get a "round" set of territories (N, S, E, W only) for phonemes or a square set of territories (includes diagonals) for distance. |
| SE | Whether to get the southeasrern territory. |
| NE | Whether to get the northeastern territory. |
| SW | Whether to get the southwestern territory. |
| NW | Whether to get the northwestern territory. |
| south | Whether to get the southern territory. |
| north | Whether to get the northern territory. |
| east | Whether to get the eastern territory. |
| west | Whether to get the western territory. |

| | |
|------------------|--------------------------|
| DefineParameters | <i>Define Parameters</i> |
|------------------|--------------------------|

Description

Creates a parameter data structure for running simulations.

Usage

```
DefineParameters(Rows = 40, Cols = 50, ChanceExpand = 0.8,  
PopulationStartIndex = c(1, 2), NumPopulationPhonemes = rep(NA,  
length(PopulationStartIndex)), UsePopSize = TRUE,  
IndividualsStEmSuEM = c(1000, 10, 20, NA), MutationRate = 15,  
PhonemeDitribution = c(12, 24, 133), Consonants = 750,  
Vowels = 100, MinConsonant = 6, MinVowel = 6,  
PhonemeProbabilityType = "RealMimic", GrowthRate = 5,  
Barriers = FALSE, BarrierLength = 30, BarrierBreaks = 4,
```

```

MutationTypeChance = rep(1/5, 5), HorizontalRate = 0.1,
Bias = TRUE, Steps = 1, HorizontalLocal = TRUE,
NumberRandomHorizontal = 8, UpRoot = TRUE, Death = TRUE,
Bering = FALSE, MigrationSimSteps = 300, HorizontalSimSteps = 400,
Waves = FALSE, Seed = NA)

```

Arguments

| | |
|------------------------|---|
| Rows | The number of rows in the world matrix. |
| Cols | The number of columns in the world matrix. |
| ChanceExpand | The chance that a population will either move or send off a group of individuals to found a new population. |
| PopulationStartIndex | The position in the matrix where each seed population starts. The number of seed populations is defined by the number of starting indices. |
| NumPopulationPhonemes | The number of phonemes in each starting population. If set to NA, this is decided by sampling from a distribution with min, mode, and mean on the values from the PhonemeDistribution argument. |
| UsePopSize | Whether to take into account the the population size (number of people) when making decisions about moving, immigrating, and phoneme loss/addition biases. |
| IndividualsStEmSuEM | Four related parameters: 1) The number of individuals a seed population starts with, 2) the minimum number of individuals required to make a founder party to settle a new territory, 3) the minimum number of individuals that must stay behind when a founder party is sent off, and 4) the maximum number of individuals allowed to be in one founder party. |
| MutationRate | The rate at which phonemes mutate. E.g., if MutationRate==0.1, each phoneme in a population's phoneme inventory has a 10% chance to mutate. |
| Consonants | The number of possible consonants in existence. Default based on real phoneme data. |
| Vowels | The number of possible vowels in existence. Default based on real phoneme data. |
| MinConsonant | The minimum number of consonants that can be in a population's phoneme inventory. Default based on real phoneme data. |
| MinVowel | The minimum number of vowels that can be in a population's phoneme inventory. Default based on real phoneme data. |
| PhonemeProbabilityType | The method by which phoneme probabilities are established. Can be Real (uses the real data verbatim, and requires the correct number of consonants and phonemes), RealMimic (uses the real data to generate a new distribution of probability similar to the real data, can be used with any number of phonemes), Equal (all phonemes are equally likely to be known), Frequency (based on how common phonemes are across populations in the simulation), or Random (randomly generated). |
| GrowthRate | When an integer, the number of individuals added to each population every time step. When a fraction, the percent that a population increases each timestep. |
| Barriers | Whether to create "snake barriers" that limit the direction of migration in the matrix. |

| | |
|------------------------|---|
| BarrierLength | The width of snake barriers. |
| BarrierBreaks | The height of the space between snake barriers. |
| MutationTypeChance | The chance that each mutation type occurs. 1) Add, 2) Lose, 3) Split, 4) Join, and 5) Shift. |
| HorizontalRate | The fraction of the population that attempts to modify its phoneme inventory every horizontal timestep. |
| Bias | Whether to randomly bias mutations towards either gains or losses when populations are small. Set to true based on previously published data. |
| Steps | The number of distance steps away from a target location that are considered "local." Includes all 8 cardinal and ordinal directions around a target, so the local area is always a rectangle around the target location. |
| HorizontalLocal | Whether horizontal transfer occurs between local populations or globally. Set to FALSE as a control, as global horizontal transfer should abolish local patterns. |
| NumberRandomHorizontal | The number of locations to compare when HorizontalLocal==FALSE. Should be 8 when Steps==1, 24 when steps==2, 48 when Steps=3, ect. |
| UpRoot | Whether established populations can move (TRUE) or they remain in place for the entire simulation (FALSE). |
| Death | Whether populations can die out. |
| Bering | Whether to employ barriers that mimick the Bering Strait and Americas. |
| MigrationSimSteps | The number of time steps to run each wave of migration. |
| HorizontalSimSteps | The number of time steps to spend on horizontal transfer. |
| Waves | Whether migration occurs in waves or all seed populations are added at the same time. If TRUE, there is one wave for each seed population. |
| Seed | Sets a seed for reproducibility if an integer instead of NA. |
| PhonemeDistribution | The 1) min, 2) mode, and 3) max number of phonemes a population can have when sampling for seed population sizes and when preventing languages from gaining or losing too many phonemes. Defaults based on real phoneme data. |

Emigrate

Emigrate

Description

Picks which populations migrate, whether the entire population migrates or a founder party is sent off, and where the population migrates to. Allows only one population to enter a territory. When multiple populations attempt to enter the same territory, one is randomly chosen to do so while the rest stay put.

Usage

Emigrate(P, occupied, local, populations)

Arguments

| | |
|-------------|--|
| P | A list of parameters. |
| occupied | The territories with a population on them. |
| local | The local territories data structure. |
| populations | The data for all existing populations. |

Extinction

Extinction

Description

Tests which populations will die based on population size and random chance.

Usage

Extinction(populations, occupied)

Arguments

| | |
|-------------|--|
| populations | The data for all existing populations. |
| occupied | The territories with a population on them. |

GeneratePhonemeProbabilities

Generate Phoneme Probabilities

Description

Generates a vector of the probability to know each phoneme.

Usage

GeneratePhonemeProbabilities(P)

Arguments

| | |
|---|-----------------------|
| P | A list of parameters. |
|---|-----------------------|

| | |
|----------------------|-------------------------------|
| GenerateSeedLanguage | <i>Generate Seed Language</i> |
|----------------------|-------------------------------|

Description

Creates a language for a seed population.

Usage

GenerateSeedLanguage(P, phonemeProbab, seedNum)

Arguments

| | |
|---------------|--|
| P | A list of parameters. |
| phonemeProbab | The probability of gaining each phoneme in the population. |
| seedNum | Which population seed is having it's language generated. |

| | |
|--------------|-----------------------|
| GetAMutation | <i>Get A Mutation</i> |
|--------------|-----------------------|

Description

Returns a new Add, Loss, Split, Join, or Shift mutation. Biases the mutations towards gaining or losing syllables when the parameter Bias==TRUE.

Usage

GetAMutation(P, phonemes, phonemeProbab, phonemeRelatedness, gain)

Arguments

| | |
|--------------------|--|
| P | A list of parameters. |
| phonemes | The phonemes currently in the language. |
| phonemeProbab | The probability of gaining each phoneme in the population. |
| phonemeRelatedness | The phoneme relatedness list. |
| gain | Whether to bias the mutations towards gaining syllables (TRUE) or losing them (FALSE). |

GetASplitShiftJoinMut *Get A Split Shift Join Mutation*

Description

Recursively calls itself until a phoneme is found that can be used to generate the mutation type of interest. Returns Null if no phoneme can mutate appropriately.

Usage

```
GetASplitShiftJoinMut(phonemes, phonemeRelatedness, unusable = NULL,
                      type)
```

Arguments

| | |
|--------------------|--|
| phonemes | Phonemes that can be mutated. |
| phonemeRelatedness | The phoneme relatedness list. |
| unusable | Phonemes that cannot be used to obtain the correct type of mutation. |
| type | Which kind of mutation to create: Split, Join, or Shift. |

GetBeringCoords *Get Bering Strait Coordinates*

Description

Returns the hardcoded locations of the Bering Strait boundaries.

Usage

```
GetBeringCoords()
```

Getcolors *Get Colors*

Description

Creates a color gradient.

Usage

```
Getcolors(P, Data, i, colors = c("coral1", "coral4"))
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

GetDist*Get Distance*

Description

Get the euclidian distance between two territories.

Usage

```
GetDist(P, point1, point2)
```

Arguments

| | |
|--------|-----------------------------|
| P | A list of parameters. |
| point1 | One territory location. |
| point2 | Another territory location. |

GetFactorDim*Get Factor Dimentions*

Description

Given a number of consonants or vowels, creates a data structure that is as square as possible.

Usage

```
GetFactorDim(nPhonemes)
```

Arguments

| | |
|-----------|--|
| nPhonemes | The number of Phonemes (vowels or consonants). |
|-----------|--|

GetGroups*Get Groups*

Description

Returns the territories descended from each seed. Includes detailed ancestry data. Only works when Uproot and Death are FALSE.

Usage

```
GetGroups(P, Data)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|---------------|-----------------------|
| GetImmigrants | <i>Get Immigrants</i> |
|---------------|-----------------------|

Description

Tests which populations immigrate, removes those that did not migrate from the migraton data. Splits the data into populations that sent off founder parties and those that moved as a single population.

Usage

```
GetImmigrants(P, occupied, local, populations)
```

Arguments

| | |
|-------------|--|
| occupied | The territories with a population on them. |
| local | The local territories data structure. |
| populations | The data for all exising populations. |

| | |
|--------------------|------------------------------|
| GetRealPhonemeData | <i>Get Real Phoneme Data</i> |
|--------------------|------------------------------|

Description

Uses the real Phoneme data from Creanza..... UPDATE THIS!!!! to determine the phoneme probabilities.

Usage

```
GetRealPhonemeData(nPhoneme, actual, vowel = FALSE)
```

Arguments

| | |
|----------|--|
| nPhoneme | The number of phonemes (vowels or consonants). |
| actual | Whether the data is Real (TRUE) or RealMimic (FALSE). |
| vowel | If true load the vowel data, otherwise load the cosonant data. |

| | |
|--------------|----------------------|
| GetTerritory | <i>Get Territory</i> |
|--------------|----------------------|

Description

Returns a territory a population can migrate to or NA if none are available.

Usage

```
GetTerritory(local, open)
```

Arguments

| | |
|-------|---|
| local | Territories that are within reach of the target territory. |
| open | Which territoies can be migrated to (i.e. no other population currently resides there). |

| | |
|-------------|---------------------------|
| GetXYCoords | <i>Get XY Coordinates</i> |
|-------------|---------------------------|

Description

CONverts territory numbers into X,Y corrdinates.

Usage

```
GetXYCoords(P, territories)
```

Arguments

| | |
|-------------|---|
| P | A list of parameters. |
| territories | A vector of territory indicies to convert into X,Y coordinates. |

| | |
|-------------|----------------------|
| GroupBySeed | <i>Group By Seed</i> |
|-------------|----------------------|

Description

Returns the territories descended from each seed. Works for all sims, but lacks ancestry data.

Usage

```
GroupBySeed(P, Data)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|---------------|---|
| HCBSimulation | <i>Human Cultural Boundaries Simulation</i> |
|---------------|---|

Description

Runs a simulation.

Usage

HCBSimulation(P)

Arguments

P A list of parameters.

| | |
|----------------------------|-------------------------------------|
| HoritontalTransferRepeater | <i>Horitontal Transfer Repeater</i> |
|----------------------------|-------------------------------------|

Description

A wrapper for the horizontal transfer process. After migration, allow populations to exchnage phoneme information, losing or gaining syllables based on other populations in the simulation. Occurs HSims number of time steps.

Usage

HoritontalTransferRepeater(P, S)

Arguments

P A list of parameters.
S A list of the data structures.

| | |
|--------------------|----------------------------|
| HorizontalTransfer | <i>Horizontal Transfer</i> |
|--------------------|----------------------------|

Description

A function wrapper that get the language to modify and allows the phoneme change to either add/shift or remove a phoneme if this can be done.

Usage

HorizontalTransfer(P, languages, local, phonemeRelatedness, phonemeProbab,
index)

Arguments

| | |
|--------------------|--|
| P | A list of parameters. |
| languages | All languages. |
| local | The local territories data structure. |
| phonemeRelatedness | The phoneme relatedness list. |
| phonemeProbab | The probability of gaining each phoneme in the population. |
| index | The target territory whose language may change. |

Initialize

Initialize

Description

The function wrapper that makes calls to create the population and phoneme data structures and then populates them with initial data.

Usage

```
Initialize(P)
```

Arguments

| | |
|---|-----------------------|
| P | A list of parameters. |
|---|-----------------------|

Lose

Lose a Phoneme

Description

Allows a language to either lose a phoneme to better match other populations.

Usage

```
Lose(P, targetLanguage, phonemeProbab)
```

Arguments

| | |
|----------------|--|
| P | A list of parameters. |
| targetLanguage | The target language to be modified if possible. |
| phonemeProbab | The probability of gaining each phoneme in the population. |

| | |
|-----------------|--------------------------|
| MakeDistanceMap | <i>Make Distance Map</i> |
|-----------------|--------------------------|

Description

Creates a distance map based on the euclidian distances between territories.

Usage

MakeDistanceMap(P)

Arguments

| | |
|---|-----------------------|
| P | A list of parameters. |
|---|-----------------------|

| | |
|--------------|----------------------|
| MakeLanguage | <i>Make Language</i> |
|--------------|----------------------|

Description

Copies and mutates the parent's language to create a new language for a founder party.

Usage

MakeLanguage(P, phonemeProbab, phonemeRelatedness, language, popSize)

Arguments

| | |
|--------------------|--|
| P | A list of parameters. |
| phonemeProbab | The probability of gaining each phoneme in the population. |
| phonemeRelatedness | The phoneme relatedness list. |
| language | The parent language to mutate into a new language. |
| popSize | The number of individuals in the parent population. |

| | |
|----------------|------------------------|
| MakePopulation | <i>Make Population</i> |
|----------------|------------------------|

Description

Generates new population based on the parent population.

Usage

MakePopulation(P, population)

Arguments

| | |
|------------|--|
| P | A list of parameters. |
| population | The population data used to make a new population. |

| | |
|-----------|------------------|
| Migration | <i>Migration</i> |
|-----------|------------------|

Description

Main simulation function. Allows populations to migrate, split, and die.

Usage

```
Migration(P, S)
```

Arguments

| | |
|---|----------------------------|
| P | A list of parameters. |
| S | A list of data structures. |

| | |
|---------------|-----------------------|
| MigrationPlot | <i>Migration Plot</i> |
|---------------|-----------------------|

Description

Shows the expansion of populations from the seed population. Only works when Uproot and Death are FALSE.

Usage

```
MigrationPlot(P, Data, groups = NA, colorSet = NA)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|--------------------|-----------------------------|
| NextStepDirections | <i>Next Step Directions</i> |
|--------------------|-----------------------------|

Description

Expands the Steps list one more step out.

Usage

```
NextStepDirections(firstStep, currentStep, start = 0)
```

Arguments

| | |
|-------------|--|
| firstStep | The original StepOne. |
| currentStep | StepOne in its current state. |
| start | How much to offset numbers (for phoneme structures). |

NextWave

Next Wave

Description

Adds the seed data for the next wave to the population and language dataframes.

Usage

```
NextWave(P, S, i)
```

Arguments

| | |
|---|------------------------------|
| P | A list of parameters. |
| S | A list of data structures. |
| i | The number of the next wave. |

OneStepDirections

One Step Directions

Description

Creates the FirstStep data structure.

Usage

```
OneStepDirections(R, C, start = 0, round = FALSE)
```

Arguments

| | |
|-------|---|
| R | The number of rows. |
| C | The number of columns. |
| start | How much to offset numbers (for phoneme structures). |
| round | whether to make the spacing Round (Phonemes) or Square (Territories). |

PhonemeFrequencyPlots

Phoneme Frequency Plots

Description

Shows how common each phonemes is in the simulation color coded by population.

Usage

```
PhonemeFrequencyPlots(P, Data, groups = NA, colorSet = NA)
```

Arguments

| | |
|------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |

| | |
|---------------|-----------------------|
| PhonemeMantel | <i>Phoneme Mantel</i> |
|---------------|-----------------------|

Description

Performs both a Hamming and Jaccard Mantel test.

Usage

```
PhonemeMantel(P, Data, repeats = 100)
```

Arguments

| | |
|---------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |
| repeats | How many times to repeat the analysis. |

| | |
|---------------------------------|---|
| PhonemePopulationFrequencyPlots | <i>Phoneme Population Frequency Plots</i> |
|---------------------------------|---|

Description

Shows how common each phonemes is in the simulation color coded by population.

Usage

```
PhonemePopulationFrequencyPlots(P, Data, groups = NA, colorSet = NA,  
sort = TRUE)
```

Arguments

| | |
|----------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |
| groups | Group structure of which territories were descended from what population seed. |
| colorSet | The colors to use. |
| sort | Whether to sort the data from most to least frequent phoneme. |

| | |
|------------------|--------------------------|
| PopulationGrowth | <i>Population Growth</i> |
|------------------|--------------------------|

Description

Adds new individuals to existing populations when population size is used in the simulation.

Usage

```
PopulationGrowth(growthRate, populationSizes, occupied)
```

Arguments

| | |
|-----------------|--|
| growthRate | The population growth rate parameter. |
| populationSizes | The number of people live on each territory. |
| occupied | The indices of territories with people living on them. |

| | |
|----------------|------------------------|
| PopulationPlot | <i>Population Plot</i> |
|----------------|------------------------|

Description

Population Plot

Usage

```
PopulationPlot(P, seedGroups, colors)
```

Arguments

| | |
|------------|--|
| P | A list of parameters. |
| seedGroups | Group structure of which territories were descended from what population seed. |
| colors | A vector of colors, one for each seed. |

`RemoveHorizontalConnections`*Remove Horizontal Connections*

Description

Affects local territories below/South (and perhaps to the Southeast and Southwest) the target territory (index) and above/North (perhaps Northwest and Northeast) of index +1.

Usage

```
RemoveHorizontalConnections(R, index, firstStep, right = TRUE,  
                           left = TRUE)
```

Arguments

| | |
|------------------------|--|
| <code>R</code> | The number of rows in the population matrix. |
| <code>index</code> | The target territory. |
| <code>firstStep</code> | The local directions created by <code>OneStepDirections()</code> . |
| <code>right</code> | Whether to remove the right diagonal. |
| <code>left</code> | Whether to remove the left diagonal. |

`RemoveVerticalConnections`*Remove Vertical Connections*

Description

Affects local territories right/East (and perhaps to the Northeast and Southeast) the target territory (index) and left/West (perhaps Northwest and Southwest) of index + R.

Usage

```
RemoveVerticalConnections(R, index, firstStep, above = TRUE,  
                          below = TRUE)
```

Arguments

| | |
|------------------------|--|
| <code>R</code> | The number of rows in the population matrix. |
| <code>index</code> | The target territory. |
| <code>firstStep</code> | The local directions created by <code>OneStepDirections()</code> . |
| <code>above</code> | Whether to remove the upper diagonal. |
| <code>below</code> | Whether to remove the lower diagonal. |

| | |
|-----------------|-------------------------|
| ResetPopulation | <i>Reset Population</i> |
|-----------------|-------------------------|

Description

Resets phoeneme and population data in the original territory when an entire population moes to a new territory.

Usage

```
ResetPopulation()
```

| | |
|-----------------|-------------------------|
| ShiftDirections | <i>Shift Directions</i> |
|-----------------|-------------------------|

Description

Returns the relationships between phonemes with an offset of start.

Usage

```
ShiftDirections(nPhonemes, start = 0)
```

Arguments

| | |
|-----------|--|
| nPhonemes | The number of phonemes. |
| start | Where to start number the phonemes (0 for consonants, number of consonants +1 for vowels). |

| | |
|--------------|----------------------|
| SnapshotPlot | <i>Snapshot Plot</i> |
|--------------|----------------------|

Description

Shows which territories are populated and from which seed they decended.

Usage

```
SnapshotPlot(P, Data, colors)
```

Arguments

| | |
|--------|--|
| P | A list of parameters. |
| Data | The Pre or Post output from an HBC simulation. |
| colors | A vector of colors, one for each seed. |

UpdateStructuresMove *Update Structures Move*

Description

Copies population data from one territory to another when the entire population migrates and then erases the original data.

Usage

UpdateStructuresMove(S, move, former)

Arguments

| | |
|--------|----------------------------|
| S | A list of data structures. |
| move | The indices of territories |
| former | The indices of territories |

UpdateStructuresRemove
Update Structures Remove

Description

Deletes population and language information for specified territories.

Usage

UpdateStructuresRemove(S, remove)

Arguments

| | |
|--------|---|
| S | A list of data structures. |
| Remove | The indices of territories whose data should be erased. |

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