# The grainscape package for R: Tutorials using simulated landscapes

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# Introduction

Are animals free to move across the landscape? What, if anything, may limit or encourage their movement and dispersal. These are the questions of landscape connectivity research, and natural resource managers and practitioners of conservation are increasingly asking them in the course of their work. What are the effects of habitat fragmentation, and can organisms find new habitat when they need to? Can corridors help? Is it possible to stop the spread of pests and pathogens by managing the connectivity of their hosts?

The grainscape package was developed for landscape connectivity analyses. The approach comes from the patch-based landscape graphs tradition (Urban and Keitt 2001; A. Fall et al. 2007; Galpern, Manseau, and Fall 2011) where a mathematical graph or network is used to represent the relationships among habitat patches. grainscape provides two types of landscape connectivity models. The first, a minimum planar graph (A. Fall et al. 2007) is an efficient approximation of the potential for connectivity among a set of habitat patches.

The second model type is the grain of connectivity, which is based on the minimum planar graph, but extends in a way that may be useful for highly mobile terrestrial animals, such as ungulates and carnivores. For these organisms the patch may be not be a discrete and definable feature or landcover type, but is rather defined probabilistically (e.g., as the probability of resource selection). By modelling the relationships among a polygons in a Voronoi tessellation of the landscape rather than discrete patches, grains of connectivity offers improvements in the ability to model highly-mobile organisms (Galpern, Manseau, and Wilson 2012). Importantly it provides continuous coverage of the entire landscape surface, in a way that a typical patch-based graph does not, and the ability to measure landscape connectivity at multiple scales. The scalability, in particular, offers flexibility to accommodate uncertainty in how species may perceive landscape features.

grainscape provides functions to extract the minimum planar graph and create two types of grains of connectivity: patch-based and lattic forms. The minimum planar graph, an essential step in all cases, is calculated based on an implementation used in SELES (A. Fall and Fall 2001).

#### **Tutorials**

There are four tutorials. The first demonstrates how to extract a minimum planar graph and illustrates how it may be used to ask questions about landscape connectivity. The second demonstrates patch-based grains of connectivity modelling where animals are known to have some affinity to a resource patch, and the third uses lattice grains of connectivity modelling, an approach that is more appropriate where a patch concept does not clearly apply. A fourth tutorial demonstrates how to produce more advanced visualizations of grains of connectivity and the corridors connecting locations.

In each case we draw on artificial landscape data included with the package. All analyses presented here can be run with the basic grainscape installation, but require additional packages be loaded:

```
library(igraph)
library(sp)
library(raster)
library(grainscape)
```

# Tutorial 1: The minimin planar graph

The minimum planar graph (hereafter MPG) is a spatial representation of a graph or a network that provides an efficient approximation of all possible pairwise connections between graph nodes (A. Fall et al. 2007). In graph-based landscape connectivity analyses, graph nodes have typically been patches of habitat that are demonstrably important for the species in question (A. Fall et al. 2007).

An MPG has links (*i.e.*, models the possibility for organism movement and dispersal) between spatially adjacent habitat patches. In some cases spatially adjacent patches may not be linked, if the shortest connection between them can be made through a third patch. In practice, this property means that the MPG can be used to make a simple and easily visualized picture of how a set of habitat patches is connected. The alternative, the complete graph, can quickly become challenging to interpret because these may contain a dense set of graph links making the pattern difficult to discern. A second advantage of the MPG is the much reduced set of graph links; this can be valuable where computational efficiency is important, and essential where the number of habitat patches being modelled numbers in the thousands.

However, there are some types of connectivity analyses where the MPG approximation of the complete graph is not appropriate. For example, assessing community structure within a landscape patch network (*i.e.*, finding sets of patches that are densely connected) is not possible as redundant connections have been removed intentionally. Equally, the MPG is a poor choice for prioritizing the influence of a patch for connectivity, the objective in a number of landscape graph studies (*e.g.* Pascual-Hortal and Saura 2006). Please see Galpern, Manseau, and Fall (2011) for further discussion of these limitations and of the MPG.

# Step 1: Preparing the resistance surface

The MPG has typically been constructed using least-cost path links between the perimeters of habitat patches. This implies that landscape structure in the "matrix" between patches is influencing movement, and that the organism in question is on average minimizing its costs when moving through this matrix (an assumption possibly appropriate for terrestrial animals, and terrestrial animal-dispersed plants). Equally, MPGs can be constructed using Euclidean links, where the only influence of the matrix on movement is the effect of spatial separation (*i.e.*, distance it presents between neighbouring habitat). In the following example we illustrate just the case where links are least-cost paths on a resistance surface. Euclidean links can be produced by passing a uniform cost surface (a constant raster), and a raster describing the patches. The only difference in analysis is at the first step below.

We begin by loading a landscape raster distributed with the package. Note that any raster format readable using the raster package can be used here. The .asc format rasters distributed with the package are ESRI ArcASCII format.

```
patchy <- raster(system.file("extdata/patchy.asc", package = "grainscape"))</pre>
```

Then, for convenience, we use R to turn this raster into a resistance surface. In this example we will assume the feature class 1 are the patches, so we will set them to resistance value also equal to 1 (*i.e.*, no additional resistance to movement than distance alone). The river, feature class 2, is assigned the highest resistance of 10. Other features are assigned values in between. The parameterization of resistance surfaces is itself a big topic (Zeller, McGarigal, and Whiteley 2012).

patchyCost <- reclassify(patchy, rcl = cbind(c(1,2,3,4,5), c(1, 10, 8, 3, 6)))
plot(patchyCost)</pre>

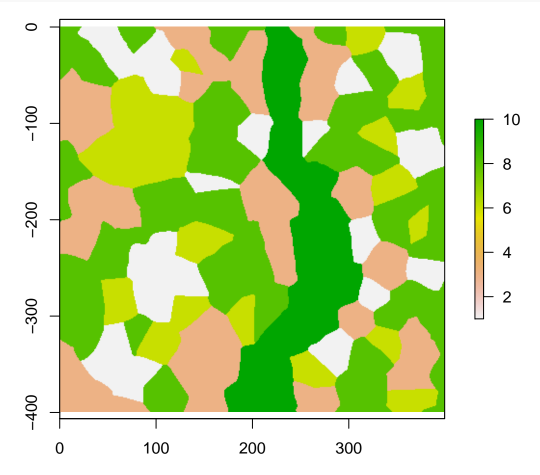


Figure 1: Input raster resistance surface to create MPG. Features with value of 1 will be the patches in the graph. A river (dark green) has the highest resistance in this example.

The result is shown in Figure 1.

## Step 2: Extracting the MPG

With a resistance surface in hand the next step is to create the MPG. Basic use of the function, MPG() to do this is shown here. Here for simplicity we assume that all areas with the resistance value equal to 1 on the raster are patches.

```
patchyMPG <- MPG(patchyCost, patch = (patchyCost == 1))</pre>
```

Note that we defined the patches by passing a binary raster patchyCost==1 as input. If patches are to be defined in a different way (e.g., using resistance classes 1 and 2) then a binary patch raster can be produced as patch=patchyCost %in% c(1,2). Equally, we could use the raster package to load a binary raster produced manually in GIS software and specify the resulting raster object as the patch parameter.

#### Step 3: Quick visualization of the MPG

A quick way to visualize the MPG is provided in the MPG object produced by MPG().

```
plot(patchyMPG)
```

However, I've done something a little more advanced so that it shows up cleanly here. The result is shown in Figure 2.

```
## Plot the patches / nodes
plot(patchyMPG@patchId, col = "black", main = "", legend = FALSE)

## Before plotting the links we'll make them wider so they print nicely
links <- patchyMPG@lcpPerimWeight > 0
links[boundaries(links, type = "outer")] <- 1
plot(links, col = "darkgrey", add = TRUE, legend = FALSE)</pre>
```

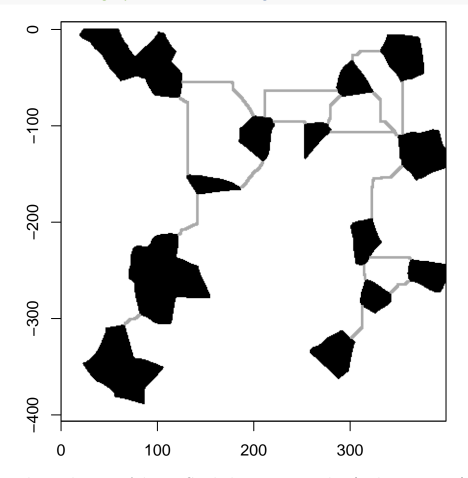


Figure 2: A quick visualization of the MPG. Black areas are patches (nodes or vertices) in the graph, and grey lines are links (edges) showing the least-cost path between the perimeters of the patches. In depth discussion of how the MPG is generated can be found elsewhere (A. Fall et al. 2007).

## Step 4: Reporting on the MPG

Following extraction, the MPG is available as an igraph object (see patchyMPG@mpg) and can be analyzed using any of the functions in this package. A quick way to report on the structure of the graph in tabular format is provided by the function graphdf():

```
graphdf(patchyMPG)
```

The output shows the structure of the vertices (nodes) and their attributes under the list element \$v\$ as well as the structure of the graph in the form of an edge list (i.e., pairs of edges e1 and e2 that are connected) and associated edge (link) attributes under the list element \$e\$. Note that only the first three lines of each has been reproduced here. Please see the manual for the interpretation of the attributes.

Note, however, that the attribute eucPerimWeight is not the same as extracting the MPG on a uniform (Euclidean) cost surface. This attribute describes the Euclidean distance between patch perimeters given the Voronoi tessellation of the cost surface (the dual of the MPG). Please see the next tutorial for information about the Voronoi tessellation.

## Step 5: Thresholding the MPG

A frequent step in the analysis of a patch-based landscape graph is to threshold the graph into a series of subgraphs or components representing connected areas for an organism (Urban and Keitt 2001; Galpern, Manseau, and Fall 2011; Galpern, Manseau, and Wilson 2012). This has sometimes been called a scalar analysis (Brooks 2003).

The function threshold() provides a way to conduct a scalar analysis at multiple scales. Here we ask for 5 thresholds, and the function finds five approximately evenly-spaced threshold values in link length:

```
scalarAnalysis <- threshold(patchyMPG, nThresh = 5)
scalarAnalysis$summary</pre>
```

```
##
     maxLink nComponents
## 1
            0
                         13
## 2
          204
                          4
                          2
## 3
          408
                          1
## 4
          612
## 5
          816
                          1
```

The \$summary of this analysis can be plotted to explore scales of aggregation in the landscape. Figure 3 shows a scalar analysis of this landscape with 100 thresholds, where the response variables is the number of components or subgraphs created by the thresholding.

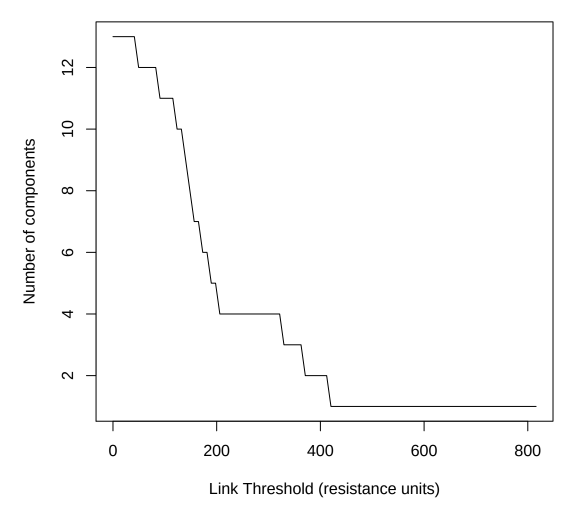


Figure 3: A scalar analysis at 100 thresholds of the MPG in Figure 2. When the landscape is a single component at higher link thresholds all patches are completely connected. As an example, an organism able to disperse 200 resistance units would experience this landscape as 6 connected regions.

Other independent variables about the components (e.g., area of patches) could, of course, be calculated by processing the thresholded graphs scalarAnalysis\$th and their attributes using the igraph function clusters().

# Step 6: Visualizing a thresholded graph

Consider an organism able to disperse a maximum of 200 resistance units. According to Figure 3 this organism would experience this landscape as 6 connected regions. This is visualized in Figure 4.

```
## Plot the patches
plot(patchyMPG@patchId, main = "", col = "black", legend = FALSE)
```

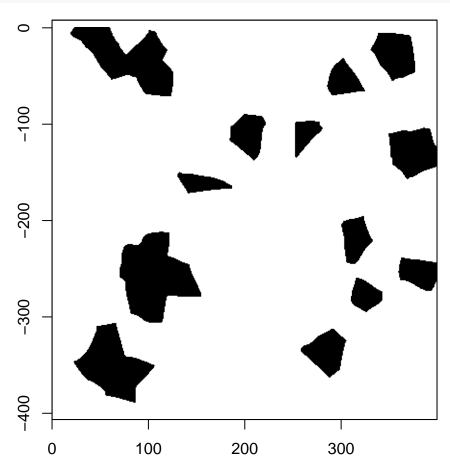


Figure 4: The thresholded MPG depicted with a link length of 200 resistance units. An organism that can disperse a maximum of 200 resistance units would experience this landscape as 6 connected regions in the depicted spatial configuration. Note that the plotting has been made a little more complicated than it needs to be in order to emphasize the links for printing.

```
## Plot the links, but emphasize them first for printing
#thresholdLink <- (patchyMPG@lcpPerimWeight <= 200)
#thresholdLink[thresholdLink == 0] <- NA
#thresholdLink[boundaries(thresholdLink, type = "outer")] <- 1
#plot(thresholdLink, col = "darkgrey", add = TRUE, legend = FALSE)</pre>
```

#### Step 7: Next steps

With the MPG in hand, all sorts of subsequent analyses about landscape connectivity are possible. Grains of connectivity (GOC), the subject of the next three tutorials, is an example of using the MPG and its complement the Voronoi tessellation.

When programming your own analyses in R based on the MPG it is helpful to observe that the patchyMPG@patchId and patchyMPG@lcpLinkId rasters contain the numerical IDs of the patches (nodes) and links respectively. These are are also contained as attributes in the igraph object patchyMPG@mpg. Using these three data objects together gives flexibility to visualize any graph analysis. To explore the attributes further use graphdf().

## Tutorial 2: Patch-based grains of connectivity

Grains of connectivity (hereafter GOC) was initially developed in two papers (Galpern, Manseau, and Wilson 2012; Galpern and Manseau 2013). Please refer to these papers for much more detail on this method. In summary, grains of connectivity describes a tessellation of functionally-connected regions of the landscape. These may be especially useful for modelling landscape connectivity for highly-mobile terrestrial organisms that are not obligate patch occupants.

Patch-based GOC, shown in this tutorial, may be most relevant for oganisms that have patch affinities, but experience "habitat" as a probabilistic surface. Lattice GOC, shown in the next tutorial, is likely more appropriate when a patch affinity cannot be supported (*i.e.*, there is no resource for which proximity is demonstrably important). This may also be the case when modelling landscape connectivity for large terrestrial mammals.

#### Step 1: Begin with an MPG

Here, we repeat the steps in Tutorial 1 for the same resistance surface. Any of the variations imaginable for the MPG are equally valid here.

```
patchy <- raster(system.file("extdata/patchy.asc", package = "grainscape"))
patchyCost <- reclassify(patchy, rcl = cbind(c(1,2,3,4,5), c(1, 10, 8, 3, 6)))
patchyMPG <- MPG(patchyCost, patch = (patchyCost == 1))</pre>
```

#### Step 2: Exploring the Voronoi tessellation

Before we build the GOC graph, we should explore the essential building block of GOC, which is the Voronoi tessellation. This particular Voronoi tessellation was first described elsewhere (A. Fall et al. 2007) and is the complement of the MPG. It is found by finding the region of proximity in resistance units around a resource patch. In contrast to a typical Voronoi tessellation where the generators are points and distance is Euclidean, this tessellation uses two-dimensional patches as generators and distance is calculated in cost or resistance space. Ultimately, the tessellation is found in SELES using a marching algorithm.

The Voronoi tessellation was extracted by MPG() in order to build the MPG. It can be plotted simply as follows:

#### plot(patchyMPG@voronoi)

A plot with the Voronoi tessellation and the patches superimposed is shown in Figure 5.

```
plot(patchyMPG@voronoi, main = "")
plot(patchyMPG@patchId, add = TRUE, col = "black", legend = FALSE)
```

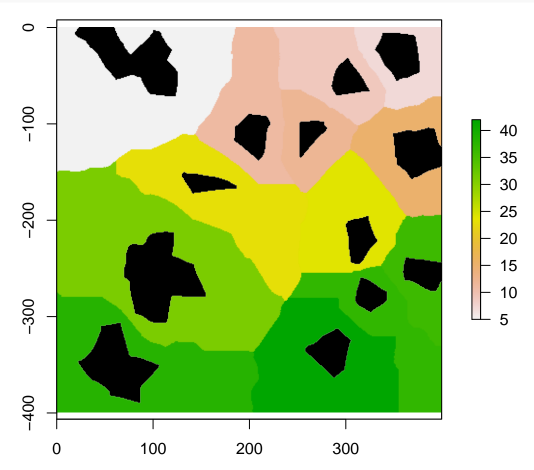


Figure 5: A Voronoi tessellation. This is the complement of the MPG. The patches are used as generators, and regions of proximity (polygons of different colours) are found in cost or resistance units. The method was first described by A. Fall et al. (2007).

# Step 3: Building GOC models

GOC analysis is essentially a scalar or thresholding analysis (see threshold() and 'Tutorial 1') except the graph being thresholded is one of Voronoi polygons rather than patches. In particular, it is the MPG of Voronoi polygons that is thresholded. As links are added, and Voronoi polygons linked, the relevant polygons are combined describing larger connected regions. An additional step is that the links connecting a pair of polygons are the mean value of all links connecting any patches in each of the two polygons from the MPG (Galpern, Manseau, and Wilson 2012).

The function GOC() builds GOC models at multiple thresholds. As with threshold() we can specify the number of thresholds, or grains of connectivity models, we want to create using the nThresh

parameter.

```
patchyGOC <- GOC(patchyMPG, nThresh = 100)</pre>
```

# Step 4: Visualizing a GOC model

To get a quick sense of the connected regions described by a GOC model at a given threshold (or scale of movement) we can use the function visualize(). This example uses the functions plotting mechanism to plot the 62nd threshold in the patchyGOC object.

Figure 6 shows the resulting plot.

```
visualize(patchyGOC, whichThresh = 62, doPlot = TRUE)
```

# whichThresh=62

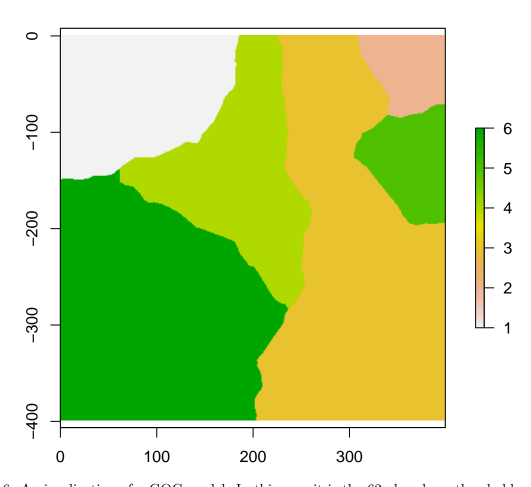


Figure 6: A visualization of a GOC model. In this case it is the 62nd scale or threshold extracted, which represents a link threshold of 168 resistance units, and a landscape of 6 components. Voronoi polygons imply regions that are functionally-connected at the given movement threshold.

```
## $summary
## maxLink nPolygon maxPolygonArea minPolygonArea meanPolygonArea
## 62 165 6 52381 5972 26666.67
```

```
##
      medianPolygonArea
                              ECS
                                   ECSCore
## 62
                23364.5 5507.376 4957.617
##
## $voronoi
## class
               : RasterLayer
## dimensions
               : 400, 400, 160000
                                    (nrow, ncol, ncell)
## resolution
               : 0.99875, 0.99875
                                    (x, y)
## extent
               : 0, 399.5, -399, 0.5 (xmin, xmax, ymin, ymax)
## coord. ref. : NA
## data source : in memory
## names
               : layer
## values
               : 1, 6 (min, max)
##
##
## $centroids
## SpatialPoints:
##
        coords.x1
                   coords.x2
## [1,] 360.40676
                   -35.83614
  [2,] 186.39126 -143.19656
         80.76057
                   -63.14828
## [3.]
## [4,] 362.72133 -131.81864
## [5,]
        98.54269 -286.92194
## [6,] 303.10301 -234.27488
## Coordinate Reference System (CRS) arguments: NA
```

#### Step 5: Identifying locations on a GOC model

Organisms have locations in space. It is often necessary to know which connected region contains a given location in a GOC model. This can be done using the function point(). First we create a thousand random locations that are on the landscape, and then run the function:

```
loc <- cbind(runif(1000)*399, runif(1000) * -399)
locGOC <- point(patchyGOC, loc)</pre>
```

The resulting object locGOC reports on the polygon which contains each location at each threshold (pointPolygon) as well as several other variables describing features of the polygons in which point locations are contained.

# Step 6: Finding the distance between points on a GOC model

As mentioned at an earlier step, at each threshold (or scale) there is also a GOC graph describing the relationships among the polygons. We can find the distance between the polygons containing pairs of points using the distance() function. This may be particularly useful in certain types of landscape genetic analyses where genetic distances are correlated with landscape distances (Galpern, Manseau, and Wilson 2012). A key advantage is that thresholding permits scaling of these distances for coarse representations of the landscape, removing variation from the calculation of distance that may be uncorrelated with dispersal.

Here we'll use four locations on the landscape.

```
loc <- rbind(c(100, -100), c(200, -200), c(300, -300), c(100, -300))
distGOC <- distance(patchyGOC, loc)
```

The resulting object locGOC reports on the pairwise distances in accumulated resistance units between each of those four points given the GOC model at each threshold. The results for the 62nd threshold are shown, where the row and column names give the polygonId containing each of the four points.

```
distGOC$th[[62]]
```

```
## $grainD

## 1 4 3 6

## 1 0.0000 402.0000 915.6667 729.0

## 4 402.0000 0.0000 513.6667 418.0

## 3 915.6667 513.6667 0.0000 676.5

## 6 729.0000 418.0000 676.5000 0.0
```

# Tutorial 3: Lattice grains of connectivity

A lattice grains of connectivity model uses a lattice of focal points superimposed over the landscape rather than patches on the landscape as nodes in the MPG. These points are also the generators used in the Voronoi tessellation. The result is a more generic way to scale up a resistance surface to coarser scales, without requiring assumptions about a particular habitat type that may be "focal" for connectivity. All that is required for analysis is an input resistance surface, reducing the ecological parameter set of the model.

#### Step 1: Building a lattice GOC model

For lattice GOC modelling a decision is required regarding the spacing of points in the lattice. This amounts to a decision about the finest scale, or grain, to be analyzed. Analysis time quickly increases as more nodes (lattice points) are included, so this decision should be made carefully.

The steps in extracting the MPG are the same as before, except instead of specifying a raster for the patch parameter, we specify an integer. This integer gives the spacing in cells between focal points in the square lattice. Thus a square raster of 400 cells in dimension, and patch=40 produces a lattice of 10 \* 10 = 100 focal points.

```
patchyMPGLattice <- MPG(patchyCost, patch = 40)</pre>
```

Let's check how many focal points were actually used in this case:

```
max(patchyMPGLattice@patchId[], na.rm = TRUE)
```

```
## [1] 104
```

Which is what we expected. A visualization of these focal points and the MPG is shown in Figure 7.

```
## Plot the links but make them wider so they print nicely
links <- (patchyMPGLattice@lcpPerimWeight > 0)
```

```
links[boundaries(links, type = "outer")] <- 1
plot(links, main = "", col = "darkgrey", legend = FALSE)

## Add the focal points. We'll get the locations from the MPG object
focalPts <- cbind(V(patchyMPGLattice@mpg)$centroidX, V(patchyMPGLattice@mpg)$centroidY)
points(focalPts, pch = "+", cex = 1.25)</pre>
```

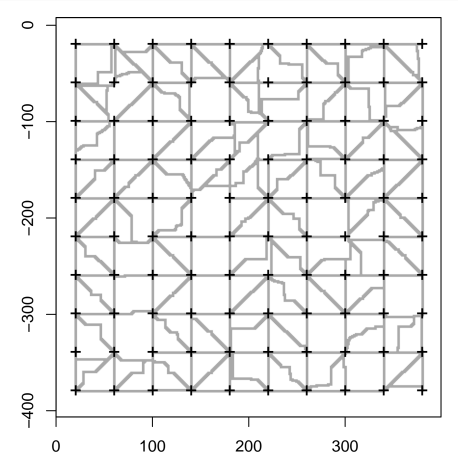


Figure 7: A visualization of an MPG of a lattice of focal points for the resistance surface in Figure 1. Again, we have complicated plotting a bit to emphasize the links better.

# Step 2: Producing and visualizing a lattice GOC model

These steps proceeds much as with the patch-based GOC model. The next step is to produce GOC model.

```
patchyGOCLattice <- GOC(patchyMPGLattice, nThresh = 100)</pre>
```

Visualization of the Voronoi tessellation for the 35th threshold is shown in Figure 8.

# whichThresh=35

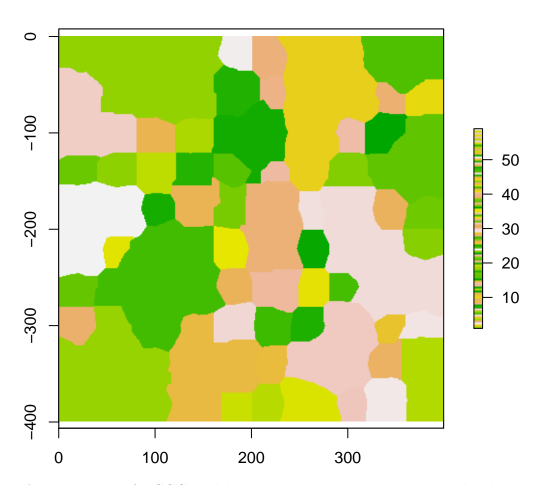


Figure 8: A visualization of a GOC model. In this case it is the 35th scale or threshold extracted, which represents a link threshold of 122 resistance units. Voronoi polygons imply regions that are functionally-connected at this movement threshold. It is easy to imagine, here, how using more lattice points makes it possible to define grains of connectivity with higher resolution.

#### Step 3: Next steps

Further analyses can proceed as with the patch-based GOC model. For example point and distance can be used, as can igraph analyses of the graphs contained in patchyGOCLattice@th and patchMPGLattice@mpg.

# Tutorial 4: Advanced visualization for grains of connectivity models

So far we have been visualizing the products of analysis using raster formats. Since all analyses take place in raster format this has the added advantage of speed. However, for publications, or on-screen visuals, it tends to be better to have high resolution vector representations; particularly of the Voronoi polygons, as this makes it easier to understand what they show without confusing blocks of colour.

In the two sections of this tutorial we demonstrate how to use the **rgeos** package in R to produce polygon representations of the Voronoi polygons in a GOC model, as well as using a **grainscape** function intended for advanced visualizations.

#### Section 1: Visualizing GOC models with vectorized Voronoi polygons

For final analyses where visualization is a goal, build the GOC model with the sp=TRUE parameter set. Also, the rgeos package must be installed. Here, we do this for both the patch-based and lattice GOC models of Tutorials 2 and 3, respectively. Note that this can be very slow, consuming orders of magnitude more computational time!

```
patchyGOC_SP <- GOC(patchyMPG, nThresh = 100, sp = TRUE)
patchyGOCLattice_SP <- GOC(patchyMPGLattice, nThresh = 100, sp = TRUE)</pre>
```

Note how we can now visualize the Voronoi tessellation also with the sp=TRUE parameter to make things sharper. Figure 9 shows this for the lattice GOC model at the finest threshold possible, superimposed over the input resistance surface used to generate the GOC model.

```
plot(patchyCost, legend = FALSE, main = "")
plot(visualize(patchyGOCLattice_SP, whichThresh = 1, sp = TRUE)$voronoiSP,
        add = TRUE)

focalPts <- cbind(V(patchyMPGLattice@mpg)$centroidX, V(patchyMPGLattice@mpg)$centroidY)
points(focalPts, pch = "+", cex = 1.25)</pre>
```

## Section 2: Visualizing the complete GOC model, corridors and shortest paths

grainscape provides a function called corridor() to simplify advanced visualizations when the goal is to show a GOC, the graph connecting Voronoi polygons and a shortest path or corridor between certain points on the landscape.

Figure 10 shows a visualization for the patch-based GOC model, and Figure 11 shows a visualization for the lattice GOC model. Finally Figure 12 demonstrates how to take manual control of the visualization to liven up the product!

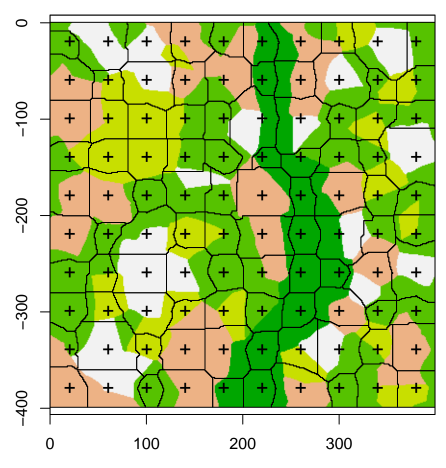
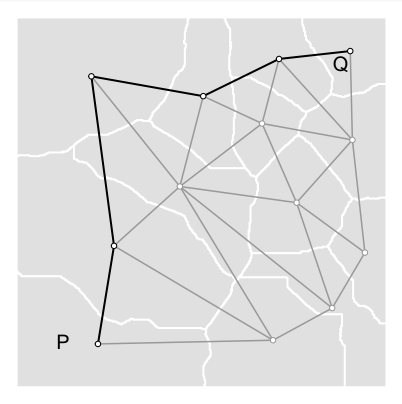
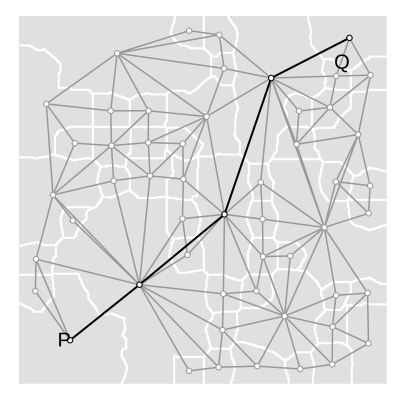


Figure 9: A visualization of a lattice GOC model using vectorized Voronoi polygons, superimposed over the input resistance surface. In this case the Voronoi polygons show the finest grain possible using the 100 point lattice (+) initially specified. Plotting only the outlines of the Voronoi polygons makes it easier to show other information on this map (e.g., resistance surface) and retain clarity. Recall, when interpreting this figure, that the polygons show regions of proximity from each focal lattice point given distance in resistance units.



Shortest path between P and Q: 1243 resistance units

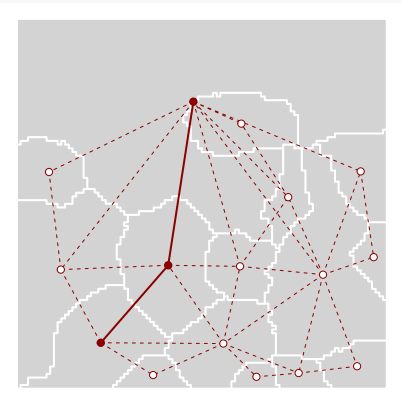
Figure 10: A visualization of the complete patch-based GOC model at its finest scale and the shortest path or least-cost corridor given the GOC graph between two locations on the map (P and Q) at this scale. Grey regions are Voronoi polygons in the patch-based GOC model.



Shortest path between P and Q: 1589.12 resistance units

Figure 11: A visualization of the complete lattice GOC model at a relatively coarse scale and the shortest path or least-cost corridor given the GOC graph between two locations on the map (P and Q) at this scale. Grey regions are Voronoi polygons in the lattice GOC model.

```
## Produce a patch-based MPG where patches are resistance features=1
tinyPatchMPG <- MPG(cost = tinyCost, patch = (tinyCost == 1))</pre>
## Extract a representative subset of 5 grains of connectivity using sp=TRUE
tinyPatchGOC <- GOC(tinyPatchMPG, nThresh = 5, sp = TRUE)</pre>
## Visualization of a corridor
corridorStartEnd <- rbind(c(10,10), c(90,90))
tinyPatchCorridor <- corridor(tinyPatchGOC, whichThresh = 3, coords = corridorStartEnd)</pre>
plot(tinyPatchCorridor$voronoiSP, col = "lightgrey", border = "white", lwd = 2)
plot(tinyPatchCorridor$linksSP, col = "darkred", lty = "dashed", add = TRUE)
plot(tinyPatchCorridor$nodesSP, col = "darkred", pch = 21, bg = "white", add = TRUE)
plot(tinyPatchCorridor$shortestLinksSP, col = "darkred", lty = "solid",
     lwd = 2, add = TRUE)
plot(tinyPatchCorridor$shortestNodesSP, col = "darkred", pch = 21,
     bg = "darkred", add = TRUE)
mtext(paste("Corridor shortest path length:",
            round(tinyPatchCorridor$corridorLength, 2),
            "resistance units"),
      side = 1)
```



Corridor shortest path length: 151.5 resistance units

Figure 12: More control over a corridor visualization using an example from the manual, and the tiny.asc resistance surface.

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