Research Article

Efficient regionalization techniques for socio-economic geographical units using minimum spanning trees

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Regionalization is a classification procedure applied to spatial objects with an areal representation, which groups them into homogeneous contiguous regions. This paper presents an efficient method for regionalization. The first step creates a connectivity graph that captures the neighbourhood relationship between the spatial objects. The cost of each edge in the graph is inversely proportional to the similarity between the regions it joins. We summarize the neighbourhood structure by a minimum spanning tree (MST), which is a connected tree with no circuits. We partition the MST by successive removal of edges that link dissimilar regions. The result is the division of the spatial objects into connected regions that have maximum internal homogeneity. Since the MST partitioning problem is NP-hard, we propose a heuristic to speed up the tree partitioning significantly. Our results show that our proposed method combines performance and quality, and it is a good alternative to other regionalization methods found in the literature.

Keywords: Regionalization; Constrained clustering; Graph partitioning; Optimization; Zone design; Census data analysis

1. Introduction

In a significant number of geographical applications, such as socio-economic, health, and census data analysis, data are organized as a large set of spatial objects represented by areas. Examples of these spatial objects include census tracts, health districts, and municipalities. In many circumstances, it is desirable to group a large number of spatial objects into a smaller number of subsets of objects, which are internally homogeneous and occupy contiguous regions in space. This procedure is called regionalization, which results in new areas (called regions) with a more comprehensive geographic extent. The idea of regionalization applied to socio-economic units (also called the zone design) was pioneered by Openshaw (1977). Grouping a set of homogeneous areal units to compose a larger region can be useful for sampling procedures (Martin 1998). Regionalization is especially valuable for

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dealing with large data sets of areal units, removing fine-scale variations and preserving relevant patterns. If properly carried out, it can produce entities that are more useful for spatial pattern discovery than the original data set (Openshaw and Alvanides 1999).

There are three types of regionalization techniques proposed in the literature. The first is a two-step procedure that applies a non-spatial clustering algorithm, followed by a neighbourhood-preserving classification. The second uses the geographical coordinates of the location as extra attributes in the clustering procedure (Haining et al. 2000). In these two techniques, the clustering procedure is not constrained by the neighbourhood relationship between the areal units, being introduced either as a second step in the method or as an indirect constraint when using coordinates as attributes. Therefore, the resulting clusters may not properly depict the inherent spatial patterns of the data set (Openshaw and Wymer 1995). In the third approach, the neighbourhood relationship between the spatial objects is used explicitly in the optimization procedure. The best-known approach to regionalization of socio-economic units based on the latter approach is the AZP algorithm (Openshaw and Rao 1995). While the original AZP algorithm used ‘Monte Carlo’ optimization, Alvanides et al. (2002) developed an alternative implementation of AZP using simulated annealing.

In this paper, we follow the third approach, aiming for an efficient regionalization procedure. We represent the objects by a graph, and our proposed algorithm prunes the graph to get contiguous clusters. The advantage of graph-based techniques is to make spatial adjacency an essential part of the clustering procedure. The challenge for graph-based regionalization techniques is twofold: to bring down the computational cost of the optimization procedure and to reduce the sensitivity on the choice of the initial position for tree partitioning. To address these questions, we present a technique for graph-based regionalization in two steps. In the first step, we create a minimum spanning tree (MST) from the neighbourhood graph. This MST represents a statistical summary of the neighbourhood graph. Then, we employ a heuristic to prune the MST and obtain a set of spatial clusters. In the paper, section 2 presents a review of the literature. Section 3 describes the proposed method, showing how to build the MST and describing how to partition the tree efficiently. In section 4, we apply the method on a case study on the city of São Paulo. Section 5 presents a performance analysis in comparison with the AZP method.

2. Regionalization approaches: a review of the literature

There are three main approaches to regionalization. The first approach is a two-step algorithm. In the first step, a conventional clustering procedure is performed using non-spatial attributes. In the second step, objects in the same cluster with no spatial contact will be split, forming different regions. This solution enables a quick evaluation of spatial dependence among objects. However, this method does not capture the spatial adjacency condition directly, resulting in limited capacity to capture spatial patterns. Another inconvenience of this approach is the lack of control on the resulting regions. Data sets with a low spatial autocorrelation will result in more regions than is desirable (Haining et al. 2000).

The second approach used in regionalization considers both the geographical position and non-spatial features of those objects. The clustering algorithm uses the coordinates of the area’s centroid as extra attributes, and measures similarity
between objects as a weighted mean of nearness in the feature space and nearness in
the geographical space. To work well, the algorithm needs a choice of weights
that produces connected clusters. The analyst has to perform the procedure several
times, trying different weights until they obtain the connected regions. This
weighted means approach is used by the regionalization method of the SAGE
(Spatial Analysis in a GIS Environment) system (Haining et al. 2000). SAGE
employs an iterative k-means clustering procedure whose objective function has
three terms: (1) homogeneity: similarity in the feature space; (2) compactness:
nearness in the geographical space, based on centroid coordinates; (3) equality:
similarity of values of a selected attribute in the resulting clusters (population, for
example). SAGE employs the equality measure to produce balanced regions for a
given attribute, to allow proper comparisons across resulting regions. Typical
control attributes are total population or population at risk for a given disease.
Martin (1998) uses a similar approach to produce output areas for the analysis of
UK census data.

Openshaw et al. (1998) criticize the weighed means regionalization algorithm,
since the three conditions (homogeneity, compactness, and equality) are not strictly
comparable. According to them, a better and simpler strategy would be to select
homogeneity as the sole basis for the objective function. Compactness and equality
are constraints, and should not be combined with homogeneity measures. Instead,
compactness and equality should be handled by defining their expected values (e.g.
minimum population within a region).

The third approach includes algorithms that use adjacency relations as constraints
to the clustering procedure. The AZP (Automatic Zoning Procedure), proposed by
Openshaw (1997), is an example. This method starts with performing a random
partition of \( n \) objects in \( k \) regions. Then, through trial and error, it seeks to
reallocate objects over regions to minimize an objective function, subject to the
adjacency constraint. Improvements to AZP led to the ZDES automatic zoning
system (Alvanides et al. 2002). Since the AZP algorithm is computationally
expensive, it is useful to consider other techniques that use the adjacency relations of
the objects and are computationally efficient. This interest leads to the authors’
proposal, described in the next section.

3. Regionalization through graph partitioning

3.1 General strategy

In this section, we describe our strategy for transforming the regionalization
problem into a graph partitioning problem. We refer to the algorithm as SKATER
(Spatial ‘K’luster Analysis by Tree Edge Removal). We use a connectivity graph to
capture the adjacency relations between objects, as shown in figure 1(a). In the
graph, each object is associated with a vertex and linked by edges to its neighbours.
The cost of each edge is proportional to the dissimilarity between the objects it joins,
where we measure dissimilarity using the values of the attributes of the neighbouring
pair. By cutting the graph at suitable places, we get connected clusters. In this way,
we transform the regionalization problem in an optimal graph partitioning problem.
Since optimal graph partitioning is an NP-hard problem, we need a heuristic
applicable to large spatial data sets to achieve suboptimal solutions in acceptable
time. For a general discussion on heuristics for graph partitioning for other types of
application, see Even et al. (1997) and Karypis and Kumar (1998).
Our proposal is to limit the complexity of the graph by pruning edges with high dissimilarity. The pruning produces a reduced graph that defines a smaller class of possible partitions and whose edges join similar areas. In the reduced graph, further removal of any edge splits the graph into two unconnected subgraphs. To carry out this idea, we take our reduced graph to be a minimum spanning tree (MST), defined in the next section and shown in figure 1(b). After creating the MST, we obtain clusters by partitioning the tree. Since the optimal solution for partitioning an MST is also an NP-hard problem, we propose a heuristic procedure for tree partitioning that produces good-quality clusters at acceptable computational costs. We describe the method in detail below.

3.2 Definition of an MST

In this section, we define a minimum spanning tree and discuss the conditions for its uniqueness. Consider a set of areal spatial objects \( O \) with a set of attributes \( \{A_1, ..., A_n\} \). All objects have an attribute vector \( x = (a_1, ..., a_n) \) where \( a_1 \) is a possible value of the attribute \( A_1 \). The topology of the set determines a connectivity graph \( G=(V, L) \) with a set of vertices \( V \) and a set of edges \( L \). There is an edge connecting vertices \( v_i \) and \( v_j \) if areas \( i \) and \( j \) are adjacent (figure 1(a)). We associate a cost \( d(i, j) \) with the edge \( (v_i, v_j) \) by measuring the dissimilarity between objects \( i \) and \( j \) using their attribute vectors \( x_i \) and \( x_j \). The dissimilarity measure is context-dependent. When the attributes have comparable scales, such as when they are measured in standard deviation units, a usual choice is the square of the Euclidean distance between the attribute vectors \( x_i \) and \( x_j \):

\[
d(i, j) = d(x_i, x_j) = \sum_{l=1}^{n} (x_{il} - x_{jl})^2.
\]

A path from node \( v_1 \) to node \( v_k \) is a sequence of nodes \( (v_1, v_2, ..., v_k) \) connected by edges \( (v_1, v_2), ..., (v_{k-1}, v_k) \). A graph \( G \) is connected if, for any pair of nodes \( v_i \) and \( v_j \),
there is at least one path connecting them. A spatial cluster is a connected subset of nodes. Our aim is to partition the graph \( G \) into \( C \) disjoint spatial clusters \( G_1, \ldots, G_C \), where their union is \( G \), and each is a connected subgraph. This is an NP-hard optimization problem whose objective function is based on a measure of within-cluster homogeneity.

A circuit is a path where the first and the final nodes are the same, and a tree is a connected graph with no circuits. A spanning tree \( T \) of a graph \( G \) is a tree containing all \( n \) nodes of \( G \), where any two nodes of \( G \) are connected by a unique path, and the number of edges in \( T \) is \( n-1 \). The removal of any edge from \( T \) results in two disconnected subgraphs that are spatial clusters candidates. A minimum spanning tree is a spanning tree with minimum cost, where the cost is measured as the sum of the dissimilarities over all the edges of the tree. The minimum spanning tree is unique if the costs between any node and all its neighbours are distinct (Aho et al. 1983). Should the dissimilarity measure use categorical attributes, many edges would have the same cost, and this could lead to more than one minimum spanning tree. In spatial applications involving socio-economic units, we expect the attributes to be random variables with continuous variation, resulting in a unique minimum spanning tree for the usual choices of dissimilarity measures. The algorithm used to build the minimum spanning tree makes this point clear.

3.3 MST generation

In this section, we describe the algorithm for building the minimum spanning tree. We build the MST in a recursive way, based on Prim’s algorithm (Jungnickel 1999). Given a connectivity graph \( G=(V, L) \) with a set of vertices \( V \) and a set of edges \( L \), the algorithm starts with one \( T_1 \) tree, containing only one vertex. At each iteration, we add a new edge and a new vertex to the tree. In iteration \( n \), the \( T_n \) tree contains all \( n \) vertices of \( V \) and a subset \( L_m \) of \( L \) with \( n-1 \) edges. The sum of costs associated with edges in \( L_m \) is minimal. The steps for MST generation are:

- Step 1: Choose any vertex \( v_i \) in the complete set of vertices \( V \), setting \( T_k=T_1=(\{v_i\}, \phi) \).
- Step 2: Find the edge of lowest cost \( (l') \) in \( L \) that connects any vertex of \( T_k \) to another vertex, \( v_j \), belonging to \( V \) but not to \( T_k \).
- Step 3: Add \( v_j \) and \( l' \) to the tree \( T_k \), creating a new tree \( T_{k+1} \).
- Step 4: Repeat Step 2 until all vertices have been included in the tree \( (T_n) \).

Figure 2 shows the procedure for MST construction, showing the first three and the last iteration. If more than one edge of lowest cost could be inserted in step 2 of the algorithm, the MST would not be unique. In these cases, the MST would be dependent on the choice of the vertex \( v_i \) (Step 1) and of the order of evaluation of the links costs in the Step 2. However, as explained above, in the graphs associated to socio-economic data, this is unlikely to happen.

3.4 MST partitioning

This section describes the partitioning algorithm for breaking up the MST into a set of contiguous clusters. Using the MST, we transform the regionalization problem into a tree partitioning problem. To make a partition of \( n \) objects in \( k \) regions, it is necessary to remove \( k-1 \) edges from the MST. Each resulting cluster will be a tree, with all vertices connected and no circuits (cf. section 3.2). To make a partition of \( n \)
objects in $k$ trees, we use a hierarchical division strategy. Initially, all objects belong to a single tree. As we remove edges from the original MST, a set of disconnected trees appears, and each tree in this set has a univocal correspondence with a region. At each iteration, one of the trees is split into two by cutting out an edge, until we reach the number of clusters previously stipulated.

The partitioning algorithm produces a graph $G^*$ that contains a set of trees $T_1, \ldots, T_n$ where each tree is connected but has no common edges or vertices with the other trees. At the first iteration, $G^*$ has only one tree, which is the MST. At each iteration, we examine the $G^*$ graph, and take out one edge that will divide the tree $T_i$ into two trees $T_i^1$ and $T_i^2$. This is the same as splitting a connected region into two subregions. We select the edge that brings about the largest increase in the overall quality of the resulting clusters. The quality measure is the sum of the intracluster square deviations, which needs to be minimized:

$$Q(II) = \sum_{i=0}^{k} SSD_i,$$

(2)
where \( \Pi \) is a partition of objects into \( k \) trees; \( Q(\Pi) \) is a value associated with the quality of a \( \Pi \) partition; and \( \text{SSD}_i \) is the sum of square deviations in region \( i \).

The intracluster square deviation \( \text{SSD} \) is a measure of dispersion of attribute values for the objects in a region. Homogeneous regions have small \( \text{SSD} \) values. Thus, the smaller the \( Q(\Pi) \), the better the partition. The intracluster square deviation \( \text{SSD} \) is:

\[
\text{SSD}_k = \sum_{j=1}^{m} \sum_{i=1}^{n_k} (x_{ij} - \bar{x}_j)^2,
\]

where \( n_k \) is the number of spatial objects in tree \( k \); \( x_{ij} \) is the \( j \)th attribute of spatial object \( i \); \( m \) is the number of attributes considered in analysis; and \( \bar{x}_j \) is the average value of the \( j \)th attribute for all objects in tree \( k \).

At each iteration, we have to remove an edge from the graph \( G^* \) that contains a set of trees \( T_1, \ldots, T_n \). To do this, we compare the optimum solutions for each of the trees \( T_1, \ldots, T_n \). The solution that best subdivides a tree \( T \) is the optimum solution \( S^R \), according to an objective function:

\[
f_1(S^T) = \text{SSD}_T - (\text{SSD}_{Ta} + \text{SSD}_{Tb}),
\]

where: \( S^T \) is the arrangement produced by cutting out the edge \( l \) from the tree \( T \), and \( Ta \) and \( Tb \) are the two trees produced by diving \( T \) after cutting out the edge \( l \).

At each iteration, we divide the tree \( T_i \) that has the highest value of the objective function \( f_1(S^T_i) \). The idea is to get the greatest improvement of quality at each step. Starting from an MST, we produce the clusters as follows:

- **Step 1:** Start the graph \( G^* = (T_0) \) where \( T_0 = \text{MST} \).
- **Step 2:** Identify the edge that has the highest objective function \( S^T_{To} \).
- **Step 3:** While \( \#(G^*) < k \) (desired number of clusters), repeat steps 4 and 5.
- **Step 4:** For all trees in \( G^* \), select the tree \( T_i \) with the best objective function \( f_1(S^T_{Ti}) \).
- **Step 5:** Split \( T_i \) into two new subtrees and update \( G^* \).

Figure 3 shows the method, with the first three iterations. In the first iteration, there is only one tree in \( G^* \) (the MST). Then, we select the best objective function \( S^T_{To} \) and divide the tree by cutting out the edge matching \( S^T_{To} \). This originates two new trees, \( T_1 \) and \( T_2 \). By repeating the pruning, we will create an optimal partition of the MST. However, the exhaustive comparison of all possible values of the objective function is expensive computationally. To speed up the choice, we propose a heuristic described in detail in the next section.

### 3.5 A heuristic for fast tree partitioning

In this section, we describe a heuristic procedure that speeds up MST partitioning. In section 3.4 above, we described the MST partitioning algorithm. We pointed out that, for each subtree, we need to find the edge that best subdivides it. At each iteration, the algorithm chooses the edge that maximizes the objective function (equation (4)). However, we made no mention of the computational demands involved in choosing the best solution. In fact, choosing the best edge to partition each subtree is computationally demanding. In the exhaustive case, we would need to examine all the arrangements leading to \( k \) clusters, and select the edge that leads to the optimal solution. This is an NP-hard problem that leads to a combinational
explosion. Therefore, this section describes an efficient heuristic that approximates the optimal solution at acceptable speeds.

Taken as an optimization problem, the search for the edge that best subdivides a tree equals the search for the best candidate within a solution space $S$:

$$ S = \{ S_1, S_2, \ldots, S_{n-1} \}, $$

where the $S_i$ candidate is the removal of edge $l$ from the tree. The algorithm searches for an optimum solution by analysing the neighbours of candidates already visited. We start by evaluating the solution $S_i$ at the current vertex and the solutions in its neighbourhood. Figure 4 shows how we do the expansion by neighbourhood in the solution space. In the example, $S_i$ has four neighbours: $S_j$, $S_k$, $S_l$, and $S_m$. We
evaluate all five solutions, and keep track of the solution with the highest objective function \( f_1(S) \) (cf. equation 2).

After finding out the best solution in a given neighbourhood, the next step is to select a vertex for expanding our search for even better solutions. Contrary to common sense, the proper choice is not necessarily the vertex with the highest value of objective function. The main problem with all optimization techniques is to avoid choosing solutions that are local maxima, instead of the desired global maximum, by also examining candidates that do not bring an immediate improvement to the objective function. To do this, we use a second objective function, \( f_2 \), which selects solutions that will divide the tree into two groups that are more homogeneous and balanced. The \( f_2 \) function prevents the generation of subtrees that are very uneven in size. Its value for a vertex is the smaller of two differences between the SSD value of the current tree and SSD values of the two sub-trees resulting from cutting out this vertex:

\[
  f_2 = \min[(SSD_T - SSD_{Ta}), (SSD_T - SSD_{Tb})].
\]  

A good choice for the starting-point can reduce the iterations needed for the search strategy to find the optimum solution. Considering the character of hierarchical division methods, a satisfactory starting-point is a vertex located in the centre of the tree. To find out the central vertex, we go over all edges until we identify the edge that best splits the tree into two sub-trees of similar sizes. Then, we continue with the optimization using the two objective functions. The stopping condition (SC) of the exploration strategy is the maximum number of iterations without growth in the value of \( f_1 \). In summary, the optimization heuristic is:

- **Step 1:** Start from the central vertex, \( V_c \). Insert solutions associated to edges incident in \( V_c \) in the list of potential solutions, \( S_p \). Set \( n^* = n = 0 \) and \( f_1(S^*) = 0 \).
- **Step 2:** Evaluate solutions in \( S_p \) and store them in the list \( L \).
- **Step 3:** Update the information on best available solution. Select the solution \( S_j \) in \( S_p \) with the highest objective function \( f_1(S_j) \). If \( f_1(S_j) > f_1(S^*) \), then \( S^* = S_j \) and \( n^* = n \).
- **Step 4:** Set \( n = n + 1 \). Based on the balancing function \( f_2(S_j) \), select in the list \( L \) a solution that will have its neighbourhood expanded, originating a new list of potential solutions \( S_p \).
- **Step 5:** Check the stopping condition \( (n - n^* > SC) \). If it is false, go back to Step 2. Otherwise, choose \( S^* \) as the best available solution and finish.

In this procedure, we assume:

- \( S^* \) is the best interim solution, which on completion will represent the chosen solution;
- \( n \) is the number of iterations;
- \( n^* \) is the last iteration where an improvement of the objective function \( f_1 \) has occurred;
- \( S_p \) is the list of potential solutions in the current iteration;
- \( L \) is the list of candidates that have been evaluated but not yet expanded;
- \( S_j \) is the best solution in the current list of potential solutions \( S_p \). We identify this solution after considering all solutions in \( S_p \);
- SC is the stopping condition for the search, defined as the number of iterations without improvements in \( f_1 \).
To show the behaviour of the heuristic, we present one experiment where we chose a vertex far from the best solution as the starting-point. Figure 5 presents a bar chart with the values of the $f_1$ objective function. We note a local maximum, which was reached in the seventh iteration. Six additional iterations were necessary before the value of the objective function started to increase again. We found the optimum solution at the fourteenth iteration. In this experiment, the stopping condition was eight iterations without any improvement in the $f_1$ objective function.

In the experiment, an MST vertex far from the ideal solution was the seed of the exploration procedure. With this choice, we intended to show how the search strategy escapes from local maxima. In a real case, we should select a starting-point that reduces the iterations needed to find an optimum solution. Considering the character of hierarchal division methods, a good starting-point is a vertex located in the centre of the tree. Our tests show that adopting a central vertex as starting-point reduces the number of evaluations to find the best solution.

### 3.6 Performance impact of the heuristic

This section discusses the performance impact of the proposed heuristic, compared with the exhaustive search method. The data set has 415 municipalities in the Brazilian state of Bahia, which were grouped in 20 regions based on three attributes. These attributes measure the percentage of municipality area with three land cover types: crops, pasture, and woods. We used two different values for the stopping condition (SC=15 and SC=30). Higher values of the stopping condition SC result in better-quality partitions, at a higher computational cost. Table 1 presents results, comparing the exhaustive search (where no heuristic is applied) with the heuristic using two different stopping conditions.

![Figure 5. Values of the objective function during exploration of the solution space in the experiment.](image-url)

|                    | Exhaustive search | Heuristic with SC=30 | Heuristic with SC=15 |
|--------------------|-------------------|----------------------|----------------------|
| $Q(\Pi)$—partition quality | 380.22            | 380.22               | 390.88               |
| Number of evaluations | 8060              | 1818                 | 1121                 |
| Relative performance (no optimization=100) | 100               | 31                   | 17                   |

Table 1. Performance comparison of MST partitioning methods.
Table 1 compares the quality of the resulting partitions, the number of evaluations and the relative performance. The quality measure is the sum of the intracluster square deviations, which needs to be minimized (equation (2)). The exhaustive search produces the best quality, at a large performance cost. The heuristic procedure with SC=30 also achieves an optimal quality, at 30% of the running time of the exhaustive search. The heuristic with SC=15 approximates the optimal quality at 17% of the running time. The execution time in table 2 does not include time spent for MST generation, which is the same for all three cases. We also provide the number of evaluations performed by each method. An evaluation is the set of all operations in one loop of the MST partitioning algorithm described in section 3.4. Each evaluation selects one possible edge to be removed.

4. Applying SKATER: a case study in São Paulo

In this section, we show how to apply the SKATER algorithm for a case study in the city of São Paulo, Brazil. The study has two parts: the first study shows results without restrictions in the regionalization procedure. The second part shows how SKATER can incorporate restrictions. The data for the study consist of the ‘Social Exclusion/Inclusion Map of the City of São Paulo’ (Câmara et al. 2004). Based on data from the 1991 census and the legal division of São Paulo into 96 districts, the social exclusion/inclusion map has four socio-economic indexes: income distribution, quality of life, human development and gender equality. The indexes are normalized in a continuous scale of \([-1, 1]\).

The first study involves obtaining eight homogenous regions for São Paulo, starting from the 96 districts and using the four socio-economic indicators of the Social Exclusion/Inclusion Map. We used the SKATER algorithm as described in the previous section. Figure 1(a) shows the districts map and the connectivity graph, and figure 1(b) shows the MST. The tree partitioning requires cutting out seven edges (figure 6(a)). The result of the unrestricted regionalization (shown in figure 6(a)) has great variations in the number of districts of each region. The north-eastern region has one district only and has 12 408 inhabitants, which is 0.12% of total population in the municipality. The larger region next to it has 36 districts with a population of 3 533 082 inhabitants, which is 36.6% of the total population. For many applications, this unbalance is not convenient. Therefore, we need to add restrictions to the regionalization procedure. In studies involving rare events, for example, it may be necessary to establish minimum values for the population at risk in the resulting clusters, for resulting rates to be representative.

A simple way to add restrictions to SKATER is to set upper and lower limits for certain attributes of a region. If a solution is off limits, it will be considered invalid. Thus, the search strategy includes both a condition of homogeneity of regions and a condition for minimum or maximum value of attributes in a region. We performed a

| No. objects | Skater | AZP |
|-------------|--------|-----|
|             | Q(P)   | Time (s) | Q(P) | Time (s) | Q(P)% | Time% |
| 96          | 5.046  | 10   | 7.0584 | 92.2 | 0.715  | 0.108 |
| 415         | 403.32 | 216  | 408.072 | 2588.6 | 0.988  | 0.083 |
| 853         | 3.141  | 889  | 3.1874 | 9206  | 0.985  | 0.097 |
second regionalization procedure with a constraint of a minimum population of 500,000 inhabitants for a region. The result is shown in figure 6(b). Regions are more balanced in population, which now varies from 651,513 to 2,543,743 inhabitants.

5. Comparison between SKATER and AZP

In this section, we compare the SKATER and the AZP algorithms. We selected AZP, since it is a well-known algorithm and has been used for regionalization of the UK Census (Openshaw and Rao 1995, Openshaw and Alvanides 2001, Alvanides et al. 2002, Martin 2003). The AZP algorithm is also a constrained classification procedure of the same type as SKATER. We recognize that there is no decisive way to compare two classification methods, because performance is dependent on the characteristics of the data set used in the study. Nevertheless, we can detect some trends with well-selected experiments. We compared SKATER with AZP in a series of experiments with different arrangements of number of objects, number of attributes, and resulting regions. The grouping condition used by the two algorithms in the two experiments was the internal homogeneity of the regions. We used three data sets in the comparison: (1) 96 districts of the municipality of São Paulo; (2) 415 municipalities of the Brazilian state of Bahia state; and (3) 853 municipalities of the state of Minas Gerais. The attributes used in the experiments were socio-economic variables.

We implemented the AZP as described in Openshaw and Rao (1995). We used the same stopping condition for both methods in all experiments (SC=10). In our tests, the AZP method has been shown to be sensitive to the choice of the initial partition, changing the quality of resulting partitions in a significant way. This fact suggests that this method has a tendency to converge to local maxima. This problem has been pointed out in Openshaw and Rao (1995). Following their suggestion, we performed the AZP method five times for each experiment, using different initial partitions. We took the best value of the five runs as the partition quality for the AZP method.
We compared SKATER and AZP for processing speed and quality of the resulting partitions. We tested three cases: different number of objects, different number of attributes, and different number of resulting regions. The first experiment used the three data sets that have a different number of objects (96 for São Paulo, 415 for Bahia, and 845 for Minas Gerais). The number of resulting clusters and the number of attributes were the same (20 clusters and three attributes). The results are shown in table 2. The second experiment used the same data set (415 municipalities of state of Bahia) and same number of resulting regions (20), and varied the number of attributes (three, six, and nine attributes). The results are shown in table 3. The third experiment used the same data set (415 municipalities of Bahia state) and same number of number of attributes (three), and varied the number of resulting clusters (20, 40, 60). The results are shown in table 4.

The criterion for quality comparison was the homogeneity of the resulting regions, measured by equation (2), where a smaller value is better. SKATER was superior to AZP in all tests. The relation among the partition quality values \[ Q(P_{AZP})/Q(P_{SKATER}) \] varied between 0.678 and 0.988, and the average value was 0.856. The processing speed of SKATER was much superior to AZP, ranging from a gain of 10 to almost 30 times. The relative gain in processing speed of SKATER over AZP improved as the number of attributes increased (Table 3). The relative advantage in performance of SKATER over AZP also increased with the number of resulting clusters (table 4).

6. Conclusion

In this paper, we present SKATER (Spatial ‘K’luster Analysis by Tree Edge Removal), an efficient method for regionalization of socio-economic units represented as spatial objects, which combines the use of a minimum spanning tree with combinational optimization techniques. The main algorithm has a fast convergence towards a best-cost solution. The algorithms that use tree partitioning

| No. objects | Skater Q(P) | Time (s) | AZP Q(P) | Time (s) | Q(P)% | Time% |
|-------------|-------------|----------|----------|----------|-------|-------|
| 3           | 403.3       | 216      | 408.1    | 2588.6   | 0.988 | 0.083 |
| 6           | 806.6       | 225      | 869.8    | 3904.4   | 0.927 | 0.058 |
| 9           | 1210.0      | 234      | 1324.4   | 6011.0   | 0.914 | 0.039 |

aData set: 415 municipalities of the state of Bahia with 20 clusters.

Table 4. Comparison of AZP and SKATER for different numbers of clustersa.

| No. objects | Skater Q(P) | Time (s) | AZP Q(P) | Time (s) | Q(P)% | Time% |
|-------------|-------------|----------|----------|----------|-------|-------|
| 20          | 403.3       | 216      | 408.1    | 2588.6   | 0.988 | 0.083 |
| 40          | 245.7       | 256      | 314.0    | 4316     | 0.782 | 0.059 |
| 60          | 172.9       | 274      | 254.9    | 7988     | 0.678 | 0.034 |

aData set: 415 municipalities of the state of Bahia with three attributes.
heuristics are much faster without any quality degradation. In comparison with the well-known AZP regionalization method, SKATER produces good-quality results and is significantly faster. The algorithm also allows restrictions to be included in the regionalization procedure. In cases where homogeneity of regions is an important need, it produces a good-quality partition in a short time. We consider SKATER to be a good choice for regionalization, especially when applied to large data sets. The algorithm is available as open-source software, as part of the open-source GIS library TerraLib (http://www.terralib.org), and is included in the TerraView visualization and analysis GIS (http://www.terralib.org/terraview).

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