A local scale-sensitive indicator of spatial autocorrelation for assessing high- and low-value clusters in multi-scale datasets

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Georeferenced user-generated datasets like those extracted from Twitter are increasingly gaining the interest of spatial analysts. Such datasets oftentimes reflect a wide array of real-world phenomena. However, each of these phenomena takes place at a certain spatial scale. Therefore, user-generated datasets are of multi-scale nature. Such datasets cannot be properly dealt with using the most common analysis methods, because these are typically designed for single-scale datasets where all observations are expected to reflect one single phenomenon (e.g., crime incidents). In this paper, we focus on the popular local G statistic. We propose a modified scale-sensitive version of a local G statistic. Furthermore, our approach comprises an alternative neighborhood definition that enables to extract certain scales of interest. We compared our method with the original one on a real-world Twitter dataset. Our experiments show that our approach is able to better detect spatial autocorrelation at specific scales, as opposed to the original method. Based on the findings of our research, we identified a number of scale-related issues that our approach is able to overcome. Thus, we demonstrate the multi-scale suitability of the proposed solution.

\textbf{Keywords:} Scale, Spatial Autocorrelation, User-Generated Data, Social Media, Twitter

1. Introduction

Spatial patterns of geographic phenomena can be explored using indicators of spatial autocorrelation. Such indicators express the degree of dependence among different observations of some spatial variable (Getis 2010). In more general terms, spatial autocorrelation can be described as the correlation between a matrix of spatial relations (usually referred to as “spatial weights matrix”) and an attribute value matrix. Corresponding indices are often designed as test statistics. In such circumstances, their goal is to find unusually high degrees of spatial dependence by testing against the null hypothesis of spatial independence (Getis 2010). Typical fields where this kind of statistic is particularly helpful are human geography, epidemiology or criminology. In such fields, spatial autocorrelation statistics can, for instance, be used for finding areas of high economic prosperity, regions of elevated infectivity or crime hot spots.

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One recurring problem with spatial autocorrelation statistics is their sensitivity to spatial scale effects. Most geographic phenomena operate on a specific scale range. This typically includes both an upper and a lower distance bound. Some processes occur globally, while others are limited to small regions (Dungan et al. 2002). Therefore, geographic data acquisition requires adjusting the measuring scale to the phenomenon of interest. This is achievable with little effort in controlled experiments that rely on automated measuring devices. Appropriate geographic deployment of such devices leads to a correctly scaled dataset. However, adjusting the measurement scale becomes more difficult (or even impossible) when employing uncontrolled data acquisition methods, for instance when observing social activities through georeferenced human reports in social media feeds like Twitter. Such uncontrolled data acquisition does not allow for a priori scale adjusting and thus causes a potential misfit of the measuring scale. In addition, user-generated data often represents more than one phenomenon. Observations originating from such data sources reflect a wide array of underlying phenomena. Moreover, single contributors reporting about these phenomena typically do not interact directly. Thus, their contributions appear in a geometrically superimposed manner. A similar effect can be observed in census data, where processes operating at different scales are interacting crosswise and are aggregated to the respective datasets (Manley et al. 2006). The analysis of user-generated data in general is of ever increasing interest. Recently, social media in particular has been leveraged in diverse fields such as human mobility analysis (e.g., Hawelka et al. 2014), event detection (e.g., Crooks et al. 2013) or sentiment analysis (e.g., Mitchell et al. 2013).

However, most of the available spatial autocorrelation statistics have been developed in the context of controlled data acquisition processes. They assume some spatial variable to represent only one phenomenon, measured at a best fitting scale. In such case, it is possible to adopt a region-oriented point of view by asking the question “What region of a dataset is out of the ordinary?” Here, one just has to properly model the size and shape of the focal neighborhoods. However, multi-topic and thus multi-scale datasets like those extracted from social media are of heterogeneous nature. Every sub-region can contain observations at small scales being situated next to others at larger scales. These observations appear to be crosswise and overlapping. In fact, one region cannot be regarded as one coherent spatial unit in such cases. The question here changes to “Which observation at a certain scale in what region of a dataset is out of the ordinary?” Thus, the focus changes from being purely region-oriented towards a phenomenon-oriented viewpoint. The question is how to separate the extraordinary from the ordinary without drawing wrong conclusions from such heterogeneous mixed-scale regions.

Existing spatial autocorrelation approaches apply various strategies for coping with scale issues. One of these is to vary the spatial weights matrix in size, shape or topological configuration. A broad range of different approaches was developed over the last decades. Getis & Aldstadt (2004) figured out eleven different general schemes, without claiming completeness. A well-known scale-related issue that is related to neighborhood definition is that of topological invariance. Different topological configurations might comprise the same spatial weights matrix when being modelled by
simple binary contiguity. This effect even appears across different scales (Dacey 1965). One can avoid this kind of problem by recognizing topology in the neighborhood definition via applying an appropriate weighting scheme (Cliff & Ord 1969). Another way of dealing with scale is to use local statistics instead of global measures. These can account for non-stationary spatial processes and exogenous factors causing heterogeneity (such as topography). Thus, they can model local-scale characteristics more realistically (Fotheringham 2009).

In this paper, we propose a modified version of a local G statistic, which we call GS statistic. The “S” in the name reflects our emphasis on scale. Our version of the local G statistic is able to deal with multi-scale datasets. Spatial autocorrelation can be assessed by following a two-step approach: First, the scale range of interest is extracted by relying on a new neighborhood definition. Our neighborhood definition differs from common approaches in that all tuples of observations within the local focus are examined with respect to their scale. Furthermore, the principle of the statistic itself is modified towards operating at a certain scale, instead of mixing up different ones. This allows for unraveling the autocorrelation structure of all locally available scales separately. We further develop equations for assessing the variance and the expectation and we present a standardized version of our statistic. Finally, we test our approach by comparing it to the original method. We apply both the original and our method to a Twitter dataset consisting of a snapshot of an urban setting from the city of San Francisco and we discuss some scale-related issues.

We start the remainder of this article by giving background information on the ambiguous term of geographic scale in Section 2. Afterwards, in Section 3, we present a literature review on the field of spatial autocorrelation statistics, with special focus on scale. In Section 4, we define our modified statistic, which is being tested in Section 5. We end our paper with some concluding remarks in Section 6.

2. Background: some notes on geographic scale

The concept of geographic scale is central to this paper. Spatial phenomena are supposed to operate at a certain scale. Therefore, accounting for this property is crucial for obtaining realistic results from spatial autocorrelation analysis. However, scale is an ambiguous term. While the concept is of interest to several disciplines, each adopted a different meaning (see Gibson et al. 2000 for a multi-disciplinary overview). Ecologists use the term for describing levels in the hierarchical system of biological taxonomy or in the hierarchy of a food chain (Allen & Hoekstra 1992). Sociologists classify their research according to the scale of human relationships, i.e., into micro-, meso-, macro- and global-sociology (Smelser 1995). Scholars from political sciences or from urban planning use the term “scale” less from a quantitative than a conceptual point of view. In analogy to political jurisdictions, they classify their research into studies at the local, regional, national or international scale (Turner et al. 1989, Gibson et al. 2000).

Different notions of scale are also common even within the single discipline of Geography. Cartographic scale, for instance, refers to a ratio between model and reality. It is a proxy for the degree of spatial reduction during the process of reality abstraction
In contrast, phenomenon scale (or operational scale) describes the areal magnitude that a phenomenon covers in the real world (Lam & Quattrochi 1992, Montello 2001). Its counterpart is analysis scale (or methodological scale), which denotes the unit size used for aggregation (Lam & Quattrochi 1992, Montello 2001). The concurrent term “resolution” basically describes the same concept in remote sensing, where it is used to specify the width of equally sized grid cells. Another more general description of the concept of resolution/analysis scale has been given by Waldo Tobler. He describes this concept as the representation of the smallest distinguishable parts (Tobler 1988).

Throughout the remainder of this paper, we use the term “scale” to refer either to phenomenon or analysis scale. Both are interrelated. If one is analyzing a spatial phenomenon at a wrongly adjusted analysis scale, the analyst misses out the essential information (i.e., spatial variation) (Goodchild 2001). Thus, it is crucial to harmonize the phenomenon scale (or the “real-world” scale) and the analysis scale.

3. Literature review

A broad range of indicators for measuring spatial autocorrelation has been developed over the last decades. Many of them are of global nature and describe the average spatial autocorrelation across a given region. Popular examples include the autocovariance-based Moran’s I (Moran 1950), the semivariance-based Geary’s C (Geary 1954) or Tango’s C (Tango 1995) and Rogerson’s R (Rogerson 1998), the two latter being both related to the $\chi^2$-goodness-of-fit test. A statistic that moreover allows statements about the characteristics of the involved observations is Getis & Ord’s G (Getis & Ord 1992, Getis & Ord 1995). Zhang & Lin (2006) modified G for overcoming the problem whereby high and low values might cancel each other out. These authors also presented an alternative approach to G by decomposing Moran’s I into three separate statistics (Zhang & Lin 2007). These are respectively capable of finding either high-value, medium-value or low-value accumulation.

The indicators presented above are designed for dealing with numerical attribute values. However, more recently, some research has also taken place around indicating spatial autocorrelation in the context of categorical data. This kind of spatial association is indeed beyond the focus of this paper. However, some recent examples can be found in Boots (2003), Ruiz et al. (2010) and Leibovici et al. (2014). Most of these indicators are based on entropy measures.

Approaches to the treatment of scale and related issues can be distinguished into two general but complementary strategies: The use of local statistics and the design of spatial weight matrices. Local statistics are better suited for taking into account the local context than global ones (Fotheringham 2009). These measures assess the autocorrelation of a given local sub-region instead of subsuming the whole spatial autocorrelation structure by just one number. This category of statistics is relatively recent and is often designed to complement some corresponding and already available global measure. Examples of such statistic include $G_i$ and $G_i^*$ (Getis & Ord 1992), LISA statistics (the local versions of Moran’s I and Geary’s C) (Anselin 1995), U (Tango
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1995) or local R (Rogerson 1998). The general principle of these statistics is to compare a local neighborhood to some overall dataset. However, this is problematic when considering the potential heterogeneity of spatial regions with respect to underlying covariates. A recent approach that has been presented by Ord & Getis (2001) tries to overcome this issue by comparing contiguous regions instead.

The compilation of spatial weights matrices is another strategy for dealing with scale issues. Aldstadt & Getis (2004) revealed at least eleven different schemes for this purpose. Getis (2009) categorized them into three categories according to their respective nature. Following this, spatial weight matrices can be constructed by following a theoretical, empirical or topological point of view. Theoretical approaches are based on some underlying distance theory such as Zipf’s law (Zipf 1949). They assume the spatial weights to be exogenous to any system. The most frequently applied approach of this kind is using some sort of inverse distance. Scale is typically modelled by inducing an upper distance bound. The opposite of the theoretical approach to constructing weight matrices is constructing them in an empirical manner. Here, the analyst tries to estimate the neighborhood structure by extracting it from some reference region of a given dataset. However, this reference region is also the limiting factor for the explanatory power of such matrices. A third approach to matrix construction is trying to depict the topology as realistically as possible. These approaches are motivated by the well-known issue of topological invariance (Dacey 1965), which leads to similar matrices across different topological settings when using binary contiguity indicators.

An issue related to scale here is that differently sized spatial units are nevertheless treated similarly. Cliff & Ord (1969) suggested using suitable weighting schemes to overcome this problem. Examples of recent approaches for matrix construction include that of Getis & Aldstadt (2004) (utilization of a local statistic for assessing a proper matrix) or LeSage (2003) (Gaussian distance). Two interesting approaches with specific focus on scale are presented by Aldstadt & Getis (2006) and Rogerson & Kedron (2012). Both of them are based on successive expansions of the neighborhood size until a maximum value of a given local statistic (e.g., local Moran’s I) is reached. Another approach for finding a suitable scale is leveraging the range of local semivariograms (Lloyd 2011). However, this is more common with geostatistical scenarios such as kriging.

In summary, research on indicators for measuring spatial autocorrelation has a long-standing tradition. Indicators can be found for different types of data and originate from different domains. The same is true for scale problems, which have indeed always been important to geographic problems. However, dealing with scale remains a challenging and yet unsolved task (Getis 2006). It is interesting to note that even today, after decades of research, modeling scale remains one of the biggest challenges in spatial analysis (Fotheringham 2009). With the rise of mixed-scale datasets like those extracted from social media, this issue is becoming even more challenging. None of the available approaches focuses on this specific problem. Thus, this is the motivation for our research.
4. A scale-sensitive local G statistic

Before defining our scale-sensitive local G statistic, we first introduce the original method (Getis & Ord 1992, Ord & Getis 1995). This statistic aims to assess not only spatial autocorrelation but also the character of the observations that are involved. More specifically, it shows whether any local accumulation primarily consists of high, medium or low attribute values. Two slightly different versions of the local G statistics are available. One of them (called $G^*_i$) includes the current observation under investigation. Its counterpart (called $G_i$) neglects the observation being examined and only accounts for its neighbors. Equations 1 and 2 define both measures.

$$G^*_i = \frac{\sum_j \omega_{ij} \cdot x_j}{\sum_j x_j}$$  \hspace{1cm} (1) 

$$G_i = \frac{\sum_{j \neq i} \omega_{ij} \cdot x_j}{\sum_j x_j}$$  \hspace{1cm} (2)

The variable $x$ represents the attribute values. The matrix $\omega$ denotes a binary spatial weights matrix, where values of one indicate adjacency to observation $i$. However, non-binary matrices are also allowed. The index $j$ iterates over the adjacent observations.

4.1 Issues regarding scale

The problem that is addressed in this paper is the issue of inadequate scale treatment when it comes to multi-scale datasets. One issue that arises is related to the different scales involved in the nominator and denominator of the local G statistic. In equations (1) and (2), the nominators represent the sum of the accumulated attribute values contained in a given local neighborhood. That neighborhood may be defined by any given distance threshold. This sum is being compared against the overall sum of the attribute’s values throughout the entire dataset (represented by the denominators). Now, if one changes the distance threshold used to define the neighborhood, it will clearly result in a scale change in the nominators. However, there is no effect on the values they are being compared to, for the denominators remain unchanged. This fact causes a serious issue when it comes to multi-scale datasets, whereby phenomena occurring at different scales are compared with each other.

While the nominator represents spatial relations within a given distance range, the denominator comprises spatial relations across all scales that are present in the dataset. This is indeed not an issue with single-scale datasets, since only one scale is of interest under such circumstances. However, it becomes a problem when analyzing multi-scale datasets. In such cases, different scales are being mixed up, although they might represent different phenomena. Another problem is the way in which neighborhoods are typically defined. As mentioned in Section 3, many different approaches exist. However, they typically model the neighborhood as a fixed-size area around some observation. Furthermore, they assume to include single-scale observations. This is inappropriate for multi-scale datasets, since phenomena at different scales might be situated in close proximity to each other and overlap. Thus,
prior to redefining the original statistic, we need to introduce an alternative neighborhood definition.

### 4.2 Scale-adjusted neighborhoods

The first step of our proposed solution for overcoming the problems with multi-scale datasets is the use of scale-adjusted neighborhoods. Common approaches for neighborhood definition specify their shape, size or topological ordering (Getis 2009). The focal scale is usually modelled by choosing a sufficient neighborhood size. All instances being situated closer than a defined distance threshold are taken into account. The threshold's value is set based on the phenomenon being studied. However, in case of multi-scale datasets, one is implicitly dealing with observations at scales that are smaller (or even larger) than the intended one. Therefore, we suggest using an upper and a lower distance threshold. Moreover, these thresholds are then used for evaluating the pairwise distances between all features in the vicinity of the examined observation. If the distance between two of these features exceeds the upper bound or is shorter than the lower one, their relationship is neglected and excluded from the neighborhood. Figure 1 illustrates this approach.

![Figure 1. Schematic sketch of the proposed scale-adjusted neighbourhoods.](image)

\[ d = \text{distance}; j, k \in \mathbb{N} = \text{indices of observations}; \oplus = \text{“exclusive-or”}. \]

### 4.3 Development of the proposed GS statistic

In this sub-section, we define our approach to defining a local scale-sensitive high/low value autocorrelation statistic. This measure is derived by adapting the local G statistic, as stated above. We call our statistic “GS statistic”, where the added “S” reflects the emphasis on scale. It should be noted that our definition given below focuses on
pairwise relationships among observations. This kind of analysis is of broad interest for the analysis of data extracted from social media, where analysts are often interested in collective processes that occur within some geographic region. Thus, one might want to consider relationships among observations instead of focusing on single occurrences. Our tests, which are presented in Section 5, deal with one such example (where semantic similarities are used to establish relationships). However, it would also be of interest to generalize our basic principles to other geometric configurations. Since this is beyond the scope of this paper, we leave that open to future research.

It is necessary to introduce some preliminary definitions, which are presented in Table 1. These are used throughout the remainder of this paper. We define them at this early stage for the sake of readability of our equations. In addition, please note that we are using reduced designator notations (i.e., $G_{S}^i$ instead of $G_{S}^{d_{max}^i}$ and $f_{jk}$ instead of $f(x_j, x_k)$) for notational convenience.

### Table 1. Preliminary variable definitions

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
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<tbody>
<tr>
<td>$n$</td>
<td>Total number of point features</td>
</tr>
<tr>
<td>$\phi_{jk}$</td>
<td>Binary variable, indicating scale fit (1) or misfit (0)</td>
</tr>
<tr>
<td>$\omega_{jk}$</td>
<td>Spatial weights, indicating adjacency of $k$ to $j$</td>
</tr>
<tr>
<td>$f(x_j,x_k)$ $\triangleq f_{jk} := f : D \times D \to \mathbb{R}$</td>
<td>A function that maps two input attributes associated with points $j$ and $k$ to a real-valued variable</td>
</tr>
<tr>
<td>$\Gamma = \sum_{j} \sum_{k \neq j} \phi_{jk}f(x_j,x_k)$</td>
<td>The attribute value sum of all scale-fitting relationships shared by points $j$ and $k$</td>
</tr>
<tr>
<td>$\Phi = \sum_{j} \sum_{k \neq j} \phi_{jk}$</td>
<td>The total number of relationships fitting the analysis scale</td>
</tr>
<tr>
<td>$W = \sum_{m} \sum_{j} \sum_{k \neq j} \omega_{mj}\omega_{mk}\phi_{jk}$</td>
<td>The cumulative number of relationships across all neighborhoods fitting the analysis scale</td>
</tr>
<tr>
<td>$W_i = \sum_{j} \sum_{k \neq j} \omega_{ij}\omega_{ik}\phi_{jk}$</td>
<td>The number of scale-fitting relationships adjacent to observation $i$</td>
</tr>
<tr>
<td>$A = \sum_{m} \sum_{j} \sum_{k \neq j} \omega_{mj}\omega_{mk}\phi_{jk} f(x_j,x_k)$</td>
<td>The cumulative attribute value sum across all neighborhoods at the given analysis scale</td>
</tr>
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</table>

The definition of the proposed statistic is based on the original statistic as given by (1). Most formulas in the text are given without derivation. More detailed derivations can be found in appendices 1 to 5. Equation 3 shows our modified version of a scale-sensitive $G_{S}^i$ statistic:

$$G_{S}^i = \frac{\sum_{j}^{n} \sum_{k \neq j} \omega_{ij}\omega_{jk} \phi_{jk} f_{jk}}{\sum_{j}^{n} \sum_{k \neq j} \omega_{jk} \phi_{jk} f_{jk} m_{km}}$$
As we are operating on pairwise relationships between tuples of observations, the indices \( j \) and \( k \) represent the two observations being involved in that relationship. Thereby, the indices \( j \) and \( k \) have to be different. Otherwise, a single point would be set into a relationship to itself. An additional indicator variable denoted \( \phi_{jk} \) has also been included. Its value is 1 if the distance between two contiguous features \( j \) and \( k \) is within the interval \([d_{\text{min}}, d_{\text{max}}]\), and 0 otherwise. Furthermore, the spatial weights matrix \( \omega \) is evaluated twice. This is necessary because both observations \( j \) and \( k \) must be adjacent to observation \( i \). These modifications allow the inclusion of scale-adjusted neighborhoods as described in Section 4.2 and lead to a match between nominator and denominator scales.

Under the null hypothesis \((H_0)\) of spatial independence, each outcome of function \( f \) is supposed to be occurring equally likely (i.e., \( P(f_{jk}) = 1/n \)). Furthermore, we suppose pairwise independence between those outcomes. It follows that the expectation for \( f \) is estimated by:

\[
E[f] = \frac{\sum_j \sum_{k \neq j} \phi_{jk} f_{jk}}{\Phi} \quad (4)
\]

By using (4), we can define the empirical expectation of the GS\(_i^*\) statistic under \( H_0 \) (equation 5). The first factor and the denominator in (5) are constant across all neighborhoods. Therefore, these can be ignored and the equation reduces to \( W_i/W \). It follows that the statistic’s local expectation is supposed to be proportional to the respective neighborhood’s fraction among all neighborhoods at the given scale. This is analogous to the original method.

\[
E[GS_i^*] = \frac{E[f] \cdot W_i}{A} \quad \sim \frac{W_i}{W} \quad (5)
\]

In equations (6) and (7), we develop equations for the variance of the GS\(_i^*\) statistic. Therefore, we first need an equation for the estimate of the expectation of the squared test statistic (6). This is then used to estimate the empirical variance (7) by applying the so called one-pass algorithm (Chan et al. 1983).

\[
E[GS_i^{*2}] = \frac{W_i \sum_j \sum_{k \neq j} \phi_{jk} f_{jk}^2}{\Phi} + \frac{W_i(W_i - 1)}{A^2} \left( r^2 - \frac{\sum_j \sum_{k \neq j} \phi_{jk} f_{jk}^2}{\Phi - 1} \right) \quad (6)
\]

\[
\text{Var}_{GS_i^*} = \frac{W_i \sum_j \sum_{k \neq j} \phi_{jk} f_{jk}^2}{\Phi} + \frac{W_i^2 r^2}{A^2} + \frac{2W_i \left( r^2 - \frac{\sum_j \sum_{k \neq j} \phi_{jk} f_{jk}^2}{\Phi - 1} \right)}{A^2} \quad (7)
\]

As expected, the variance under \( H_0 \) becomes 0 if there are no neighbors in the vicinity of observation \( i \) (i.e., \( W_i = 0 \)). The same applies if no corresponding scale-fitting relationships are located in the neighborhood (\( \Phi = 0 \)) or if the total attribute value sum (\( A \)) of all those features equals zero. Similarly, the variance estimation also becomes
zero if the overall neighborhood sum equals zero. In contrast, the variance is greater than zero if all observations are contained in the neighborhood of the current feature. This is a difference from the original method. However, this becomes clear when recalling the statistic’s principle: One neighborhood is compared against all other neighborhoods. Thus, the denominator is always greater than the nominator, resulting in a nonzero variance.

The maximum value of our statistic is reached when all neighborhoods mutually contain each other. In such circumstances, the aggregation of all $\phi_{jk}$ for any tuple across the whole neighborhood forms an all-ones matrix. It follows that the maximum value of the $G^*_i$ statistic is given as:

$$\max G^*_i = \frac{1}{n} \quad (8)$$

Accordingly, the minimum value is reached if no values except the investigated observation itself are contained in some neighborhood. It follows that the minimum value is given by:

$$\min G^*_i = 0 \quad (9)$$

Equations (8) and (9) show that the range of the $G^*_i$ statistic is not fixed. This is a major difference compared to the original $G$ statistics, which range is the interval $[0,1]$. In contrast, the $G^*_i$ statistic depends on the number of input features. Thus, two $G^*_i$ values should not be compared with each other directly. A comparison is only meaningful after standardization. The standardized version of $G^*_i$ is given in (10). Applying this equation produces standard deviates (i.e., $z$-scores), which appear to be on the interval $[-\infty, \infty]$. Furthermore, following the well-known central limit theorem, these scores tend to be approximately normal, given a sufficiently large sample size. Therefore, these scores can be evaluated by means of normal theory.

$$Z_{G^*_i} = \frac{\sum \sum \phi_{ij}^2 \omega_{ij}^2 \phi_{jk}^2 - \frac{\sum \sum \phi_{ij}^2 \phi_{jk}}{\phi}}{\sqrt{\frac{W_i \sum \sum \phi_{ij}^2 \phi_{jk}}{\phi}} \left( \frac{\sum \sum \phi_{ij}^2 \phi_{jk} - \frac{\sum \sum \phi_{ij}^2 \phi_{jk}}{\phi}}{\phi - 1} \right) \left( \frac{\sum \sum \phi_{ij}^2 \phi_{jk}^2}{\phi^2} - \frac{\sum \sum \phi_{ij}^2 \phi_{jk}^2}{\phi^2} \right)} \quad (10)$$

5. **Empirical comparison between $G^*_i$ and $G^*_i$**

We now empirically illustrate the problems that occur when applying the original $G^*_i$ statistic to multi-scale datasets such as those extracted from social media. Furthermore, we also show that our approach overcomes these problems. Before this is done, we explain the datasets that we used and all the necessary preprocessing. Please note that we do not aim to analyze the regions that we sampled with respect to the qualitative properties of the underlying phenomena. All following steps are merely illustrative for testing our suggested approach with respect to the scale issues that are mentioned in Section 5.3.
5.1 Dataset description

The datasets we used were extracted from the social media service Twitter. They originate from an urban setting in the city of San Francisco, CA. We used two randomly chosen time slots. One of them covers the time period of January 30, 2014, from 8p.m. until 10p.m.; the second slot covers a whole week from the 20th of January until the 26th of January 2014.

Our automated crawler leveraged the public Twitter Streaming API. Since we are interested in applying methods from spatial statistics, we restricted our query to georeferenced tweets only. We crawled all tweets from a bounding box covering the city of San Francisco and its immediate surroundings. The bounding box had a size of approximately 15x15km. We did not restrict our data collection by using keywords or any other type of filter. The subsets of our dataset that we used for this paper sum up to a size of 1,291 tweets (for the two-hour slot) and 69,345 tweets (for the one-week slot). Figure 2 provides an overview of this subset and shows its distribution over the city.

Figure 2. Overview of our test datasets originating from San Francisco, CA. Blue = 20th of January until 26th of January; Red = 30th of January, 8pm until 10pm. More intense colours indicate higher numbers of superimposed Tweets. Base data: VMAP, National Geospatial-Intelligence Agency, US.
5.2 Preprocessing and data preparation

The crawled datasets consist of textual tweets. However, our approach as well as the original $G^*_i$ statistic are designed for dealing with numerical values. Thus, we first have to transform the textual tweets into some numerical representation. We have chosen to use similarities among the tweets for our test and comparative study. In a realistic scenario, high similarity scores might be interpreted as indicators of coherent social activity (i.e., people might be reporting about similar topics). In order to obtain meaningful similarities, several steps are conducted.

The first step is to split up the cohesive strings of words into single tokens. The tokenization process that we used follows some rules that have been adapted from the recent literature: The texts are split up at case changes, except if they occur at the beginning of a word (Metke-Jimenez et al. 2011); Twitter’s specific symbols (e.g., #, @) are kept (O’Connor et al. 2010), and short forms or contractions of English words (e.g., I’m) are retained (Pak & Paroubek 2010). Moreover, we split the tweets at whitespaces and punctuation marks. A large portion of the resulting tokens occurs frequently, but adds little meaning (e.g., “to”, “or”). Therefore, these so-called stop words are removed from the corpus in the second step. For this purpose, we relied on the English stop word list provided by the database system PostgreSQL.

The actual similarity assessment is based on the method of Latent Semantic Indexing (LSI) (Deerwester et al. 1990). The core principle of this method is based on a singular value decomposition (SVD). First of all, the tokens are transformed into normalized frequencies (called Term Frequency – Inverse Document Frequency (TF-IDF) scores). These are then used for extracting inherent components, based on word co-occurrence. LSI works in an unsupervised manner. Thus, no a priori knowledge about the text corpus is needed. However, a criterion for maintaining a reasonable number of components is required. In our experiments, we used a broken stick model for this purpose. This approach is usually used for modeling resource allocation in ecology. However, it has also proven to be useful for application of the SVD (Cangelosi & Goriely 2007).

Again, note that our approach for assessing similarities has been chosen for the sake of producing numerical tweet representations. Neither similarity assessment itself nor analyzing our test site is the focus of this paper. Thus, the chosen approach is appropriate for our experiments regarding the proposed statistic. We point out that more accurate semantic similarity approaches might be available (e.g., Latent Dirichlet Allocation (Blei et al. 2003) or probabilistic LSI (Hofmann 1999)). However, these are more sophisticated and require more detailed a priori knowledge about the composition of the text corpus. Whenever realistic conclusions are to be drawn from any dataset, careful consideration should be given to the choice of an appropriate semantic similarity approach.

5.3 Comparison between $GS^*_i$ and $G^*_i$

Our comparison focuses on three central problems that occur when the $G^*_i$ statistic is applied to multi-scale datasets. All these problems occur due to the issues highlighted in
Section 4.1. Moreover, we also demonstrate that these issues are solved by our proposed solution.

**Overemphasis of dominant scales**

Recall the property of scale mixing within social media data. Figure 3 illustrates the average composition of five differently scaled neighborhoods. These neighborhoods are heterogeneous. In most cases, the actual scale of interest contributes only approximately 30% of the total attribute value sum. This means that approximately 70% of all variation is contributed by scales other than the one of interest. Accordingly, when applying standard (i.e., single-scale) approaches for neighborhood definition, all these scales are considered together.

![Figure 3](image.png)

**Figure 3.** Average composition of the attribute value sum for five classes of neighbourhood sizes. The respective scales of interest are highlighted by displacement. Dataset: Twitter, 30th of January 2014, 8pm until 10pm.

However, if 70% of the total variation is contributed by phenomena beyond interest, it is likely to create some bias in autocorrelation results. This is particularly the case when one or more of these non-relevant scales are dominating a dataset. Figure 4 shows to what extent the respective scales are under- or overrated. It illustrates the ratio between the share in the attribute value sum and the share in the quantitative composition of the neighborhoods. It can be seen that the small scales (1-30m and 30-100m) are overrepresented in most neighborhoods. Thus, phenomena operating at such scales are excessively biasing the results at other scales.
Figure 4. Under-/ overestimation of various scales within neighbourhoods at different analysis scales. Dataset: Twitter, 30th of January 2014, 8pm until 10pm.

The problems described above affect the original $G_i$ statistic in two ways: On the one hand, scales are superimposed in the focal neighborhoods. On the other hand, these are then compared against an overall mixture of scales (i.e., the denominator of the statistic). The larger the scale, the more different scales are potentially being mixed up. Figure 5 shows one of the effects caused by that behavior. The mean of the z-values obtained through the $G_i^*$ statistic shows a strong trend with increasing scale. However, we are dealing with a standardized version of the statistic. Following the central limit theorem, the resulting standard variates are expected to be approximately normal. Thus, the mean is expected to be an unbiased estimator of the expectation, which should be close to zero in the present case. That is obviously not true for $G_i^*$ when it is applied to social media datasets. It is very likely that this effect is caused by the scale mixture described in the previous paragraph. That mixture implies different underlying populations, since different phenomena might be operating at the different scales. Thus, there are also different means present in the mixture. The mean of the z-values is influenced by that variety of means, which in turn leads to the observed bias.

Figure 5. Arithmetic means of $Z(G_i^*)$ and $Z(GS_i^*)$ across all tested scales. Dataset: Twitter, 20th of January until 26th of January.
These effects are diminished with our suggested scale-sensitive approach. Our method only extracts those scales from the vicinity of observations that are relevant for the current analysis scale. Thus, each diagram shown in Figure 3 would only consist of one pie slice, each representing the respective scale of interest. The composition of the attribute value sum of the neighborhoods is completely made up of observations fitting the scale of interest. Moreover, the same applies to the comparative size. The modified statistic only includes those observations in any calculation that are fitting the current scale of interest. Therefore, the estimated means obtained through our modified statistic (see Figure 5) remain close to zero across all investigated scales.

**Type I/Type II errors**

The problem of overemphasizing dominant scales leads to another closely related problem, which is the occurrence of type I/II errors. This is a well-known general issue of all local statistics (Nelson 2012). It is usually caused by missing strategies for facing multiple testing problems. However, when dealing with multi-scale datasets, this problem is further exacerbated by an additional problem. When some scales are dominating a dataset, they also hide weaker phenomena at less dominant scales. However, these less pronounced phenomena are not necessarily less important. Some analyst might indeed be interested in analyzing these weaker phenomena. Now, several different configurations are possible: Some weaker phenomenon might, for instance, consist of some high-value accumulation. These values might, however, only be high according to their own respective scale. Some contiguous and more dominant scale might comprise even higher values. In such situations, the dominance of the other scale with high values leads to type II errors. \( H_1 \) is rejected although high values are present at the adjusted scale of interest. These values just appear to be quite low in comparison to the more dominant adjacent scale that is present in the same neighborhood. The same situation occurs if a phenomenon of interest shows low-value accumulation. Higher values at another scale are again artificially raising the neighborhood score, leading to \( H_1 \) rejection. In contrast, type I errors occur whenever a scale of interest is actually not out of the ordinary, but gets interfered by a more dominant scale. This situation might occur in both directions, either toward low values (cold spots) or high values (hot spots). In such cases, the neighborhood score is artificially raised (or lowered) to a level that leads to a wrong acceptance of \( H_1 \).

One example from our dataset is depicted in Figure 6, which is showing two series of maps. Each of those series comprises four different scales of interest in ascending order. Those series illustrate both issues described so far. On the one hand, one can see the overemphasis of dominant scales. The results obtained through the original \( G_i^* \) at the two smallest scales show a large number of statistically significant high-value accumulations. In fact, 33.56% of all tweets of the dataset are identified to be statistically significant with \( G_i^* \) (scale = 1-30m; two-sided test; \( \alpha = 0.1 \) each). In other words, every third tweet is considered to be part of a neighborhood that comprises high-value accumulation higher than 90% of the other tweets. This is obviously an upwards biased value, due to the dominance of that scale compared to larger ones. In
comparison, the results obtained through our modified approach show a considerably lower number of extraordinary observations. Since the dominance of scales is not affecting the results, that method only evaluates 3.77% of the tweets to be somehow abnormal. Another issue that can be seen in Figure 6 is the existence of type I errors. Because of the dominance effect described above, H_0 is rejected too often. This does not only appear at the dominant scales, but is transferred onto all larger levels as well. Hot or cold spots occurring at small scales appear to be acting like “seeds” that are being enlarged at the next larger scale. Thus, the type I errors can be found with increasing frequency by enlarging the analysis scale. This effect also does not occur in the results obtained through our proposed statistic. Every scale is only analyzed against observations at the same scale. Thus, there is no dominance to be transferred, resulting into a lower number of type I errors. However, the effect of “seed” locations with G_i leads to another issue that is described in the following subsection.

**Loss of statistical independence between scales**

We already mentioned the spill-over effect of dominant scales that are transferred onto all larger ones. We can also observe that this effect results into “seed” locations that appear to be growing as the scale is getting enlarged. However, this phenomenon leads to another much more serious problem, which is the loss of independence between spatial autocorrelation results obtained for different scales. We assume all possible outcomes of spatial autocorrelation statistics to be equally likely. That is, we assume the probabilities to be \( P(x_a) = \frac{\sum x_a}{n} \). If different scales are being admixed, however, this assumption is no longer verified; this occurs, for instance, when assessing a non-zero spatial autocorrelation at some small scale. If the scale is adjusted to some larger value, these small-scale instances are again included. The problem is that now, the outcome of zero has become impossible. The effect of the non-zero spatial autocorrelation at the smaller scale might be blurred (due to mixing) or be changed in nature (from negative to positive or vice versa) because other observations are included in the neighborhood. However, the result of having no autocorrelation is no longer possible at any larger scale. In other words, the independence requirement \( P(x_b|x_a) = P(x_b) \) is no longer met. Since we are dealing with multi-scale datasets that reflect potentially unrelated phenomena, this is an inappropriate property.
Figure 6. Two series of analysis results. The left-hand side was obtained by applying the original $G_i^*$ statistic, the right-hand side originates from our proposed $G_{Si}^*$ statistic; Dataset: Twitter, 30th of January 2014, 8pm until 10pm; Base data: VMAP, National Geospatial-Intelligence Agency, US.
6. Conclusions

The arising interest in analyzing social media feeds and other kinds of human-generated datasets compels us to address the specific problems of such data. One of those problems is their multi-scale nature that is due to the uncontrolled data acquisition process. However, most spatial statistics are designed for single-scale datasets that result from controlled experiments. This paper introduced a scale-sensitive version of the popular $G_i^*$ statistic. The proposed approach comprises an alternative approach for neighborhood definition and a scale-adjustment of the statistic itself. Moreover, some scale-related issues that arise when dealing with multi-scale datasets are highlighted by comparing the results obtained through the original and the proposed statistics. These comparisons are carried out on a Twitter dataset for the city of San Francisco, CA. The results demonstrate that the suggested approach is better suited for dealing with multi-scale datasets, because it allows analyzing certain scales without cross-scale interferences. Thus, it can be used in real-world scenarios whenever social media or other human-generated datasets are analyzed.

However, scale-related effects affecting social media datasets are not yet fully understood. The list of issues mentioned in Section 5 is given without claiming completeness. There might be many more effects that are still to be discovered. Moreover, the effects we listed and observed have not yet been fully investigated. Thus, future research should focus on getting a better understanding of the multi-scale nature of user-generated datasets. In addition, there are many more methods from spatial statistics and other fields that are not yet sufficiently capable of dealing with multi-scale datasets. Our suggested approach might serve as a starting point for initiating methodological research towards multi-scale enablement.

With respect to local autocorrelation statistics in general, more emphasis should be put on the definition of the null hypothesis. Geographic space imposes uncontrolled variance, due to varying local environmental conditions (Goodchild 2009, Anselin 1989). Local statistics such as $G_i^*$ and our proposed solution already account for heterogeneity with respect to the spatial distribution of observations. In contrast, they usually include constant expectations of the observed variable. However, the outcomes of those variables might also be influenced by nonstationary environmental conditions. One way of overcoming this problem might be to use location-dependent expectation functions instead of constant values. Corresponding local values might be determined by methods such as Geographically Weighted Regression (Brunsdon et al. 1996). However, a specific problem to social media data is that the underlying driving forces are not yet fully understood.

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References


**Appendix 1: Derivation of the Empirical Expectation of GS$_i^*$**

\[
E[GS_i^*] = \frac{\sum_j \sum_{j<k} \omega_{ij} \phi_{jk} \hat{E}[f]}{\sum_j \sum_{j<k} \omega_{jk} \phi_{jm} \omega_{km} \phi_{km}} 
\]

\[
= \frac{E[f] \sum_j \sum_{j<k} \omega_{ij} \phi_{jk}}{\sum_j \sum_{j<k} \omega_{jk} \phi_{jm} \omega_{km} \phi_{km}} 
\]

\[
= \frac{E[f] \cdot W_i}{A} 
\]

Since \(\hat{E}[f]\) and \(A\) are constant, we can infer that the expectation is proportional to the share of the neighbourhood’s size among the overall sum of relationship outcomes:

\[
\sim \frac{W_i}{W} 
\]

**Appendix 2: Derivation of the Expectation of the Squared GS Statistic**

\[
GS_i^{*2} = \frac{\sum_j \sum_{j<k} \sum_{m} \omega_{ij} \phi_{jk} \omega_{km} \phi_{km} \phi_{mp} f_{jk} f_{mp}}{\sum_j \sum_{j<k} \sum_{m} \sum_{p} \omega_{ij} \phi_{jk} \omega_{jm} \phi_{jm} \omega_{pm} \phi_{pm} \phi_{km} \phi_{km} \phi_{mp} f_{jk} f_{mp}} 
\]

\[
E[f_1, f_2] = \frac{\sum_j \sum_{j<k} \sum_{m} \sum_{p} \omega_{ij} \phi_{jk} \phi_{mp} f_{jk} f_{mp} - \sum_j \sum_{j<k} (\phi_{jk} f_{jk})^2}{\Phi (\Phi - 1)} 
\]

\[
E[f^2] = \frac{\sum_j \sum_{j<k} \phi_{jk} f_{jk}^2}{\Phi} 
\]

Solving (12) leads to quadratic and non-quadratic terms. Thus, we need \(\hat{E}[f^2]\) and \(\hat{E}[f_1, f_2]\) for inferring the expectation of the squared GS statistic. Both of these values are constant. Therefore, we can extract them from the sums. Furthermore, \(\omega\) and \(\phi\) are binary and \(\omega_{ij}^2 = \omega, \phi_{j,k}^2 = \phi_{jk}\). Accordingly we can write:

\[
E[GS_i^{*2}] = \frac{W_i \cdot E[f^2] + W_i (W_i - 1) \cdot \hat{E}[f_1, f_2]}{A^2} 
\]

\[
= \frac{W_i \sum_j \sum_{j<k} \phi_{jk} f_{jk}^2}{\Phi} + \frac{W_i (W_i - 1) \left( r^2 - \sum_j \sum_{j<k} (\phi_{jk} f_{jk})^2 \right)}{\Phi (\Phi - 1)} 
\]

\[
= \frac{A^2}{A^2} 
\]
Appendix 3: Derivation of the Empirical Variance of the Local GS\textsubscript{i} Statistic

Applying the Steiner translation theorem leads to the variance of the statistic:

\[
\text{Var}_{GS_i} = E[GS_i^2] - (E[GS_i])^2
\]

\[
= \frac{W_i \sum \omega_i \Phi_{jk} f_{jk}^2}{\Phi} + \frac{W_i(W_i - 1) \left( \frac{1}{\Phi \sum \omega_i \Phi_{jk} f_{jk}^2} \right)}{A^2} - \frac{W_i^2 \left( \sum \omega_i \Phi_{jk} f_{jk}^2 \right)^2}{\Phi^2} - \frac{W_i^2 \left( \sum \omega_i \Phi_{jk} f_{jk}^2 \right)^2}{\Phi^2}
\]

Appendix 4: Derivation of the maximum of the GS\textsubscript{i} Statistic

\[
\text{max GS}_i = \frac{\sum \omega_i \Phi_{jk} f_{jk}^2}{N \cdot \sum \omega_i \Phi_{jk} f_{jk}^2}
\]

\[
= \frac{\sum \omega_i \Phi_{jk} f_{jk}^2}{n \cdot \sum \omega_i \Phi_{jk} f_{jk}^2}
\]

\[
= \frac{1}{n}
\]

Appendix 5: Derivation of the Standardised GS\textsubscript{i} Statistic

\[
Z_{GS_i} = \frac{GS_i^* - E[GS_i^*]}{\sqrt{\text{Var}_{GS_i}}}
\]

\[
= \frac{\sum \omega_i \Phi_{jk} f_{jk}^2}{A} \left( \frac{W_i \sum \omega_i \Phi_{jk} f_{jk}^2}{\Phi} + \frac{W_i(W_i - 1) \left( \frac{1}{\Phi \sum \omega_i \Phi_{jk} f_{jk}^2} \right)}{A^2} - \frac{W_i^2 \left( \sum \omega_i \Phi_{jk} f_{jk}^2 \right)^2}{\Phi^2} - \frac{W_i^2 \left( \sum \omega_i \Phi_{jk} f_{jk}^2 \right)^2}{\Phi^2} \right)
\]