High-performance solutions of geographically weighted regression in R

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ABSTRACT

As an established spatial analytical tool, Geographically Weighted Regression (GWR) has been applied across a variety of disciplines. However, its usage can be challenging for large datasets, which are increasingly prevalent in today's digital world. In this study, we propose two high-performance R solutions for GWR via Multi-core Parallel (MP) and Compute Unified Device Architecture (CUDA) techniques, respectively GWR-MP and GWR-CUDA. We compared GWR-MP and GWR-CUDA with three existing solutions available in Geographically Weighted Models (GWmodel), Multi-scale GWR (MGWR) and Fast GWR (FastGWR). Results showed that all five solutions perform differently across varying sample sizes, with no single solution a clear winner in terms of computational efficiency. Specifically, solutions given in GWmodel and MGWR provided acceptable computational costs for GWR studies with a relatively small sample size. For a large sample size, GWR-MP and FastGWR provided coherent solutions on a Personal Computer (PC) with a common multi-core configuration, GWR-MP provided more efficient computing capacity for each core than thread FastGWR. For cases when the sample size was very large, and for these cases only, GWR-CUDA provided the most efficient solution, but should note its I/O cost with small samples. In summary, GWR-MP and GWR-CUDA provided complementary high-performance R solutions to existing ones, where for certain data-rich GWR studies, they should be preferred.

1. Introduction

Geographically Weighted Regression (GWR) (Brunsdon, Fotheringham, and Charlton 1996, 1998; Fotheringham, Charlton, and Brunsdon 1998; Fotheringham, Brunsdon, and Charlton 2002) is a technique specifically developed to explore spatial heterogeneities in a regression’s “response to predictor variable” relationships. Unlike a fixed coefficient regression, such as an Ordinary Least Squares (OLS) regression, GWR allows regression coefficients to vary spatially; the resultant coefficient maps allow an investigation into their change (if any) across space. The GWR methodology has been extensively developed in terms of its usage and extensions (Comber et al. 2022), but where inference in GWR is not always as stable as that found with say, an OLS regression and as such, GWR adaptations exist to counter this (da Silva and Fotheringham 2016; Harris et al. 2017). GWR has been widely applied in many scientific domains, including regional economics (e.g. Jin, Xu, and Huang 2019), urban planning (e.g. Cao et al. 2019b), sociology (e.g. Yin et al. 2018), ecology (e.g. Liu et al. 2019), public health (e.g. Wang et al. 2019; Xu et al. 2021), agriculture (e.g. Harris et al. 2017), and environmental science (e.g. Cao et al. 2019a; Huang and Wang 2020).

Our increasingly digital world continues to generate huge volumes of data – many of which are spatially indexed (Lee and Kang 2015; Ivan et al. 2017). However, in order to attribute process understanding to such “Big Spatial Data” almost all spatial models require adaptation so they can be efficiently calibrated and validated within tolerable time frames. GWR is one such model that is computationally demanding and in this respect has benefitted from high-performance computing solutions (Harris et al. 2010; Murakami et al. 2020; Li et al. 2019b). Commonly, such solutions only exist for the conventional forms of GWR, where many extended GWR models are more computationally demanding still – for example, multiscale GWR (Lu et al. 2018; Li and Fotheringham 2020) which requires a complex iterative solution to its calibration. Similarly, Geographically and Temporally Weighted Regression (GTWR) (Huang, Wu, and Barry 2010; Fotheringham, Crespo, and Yao 2015) for space-time processes has a higher computational demand than that found with conventional GWR.

Unsurprisingly, there are an increasing number of (conventional) GWR applications exploring “Big Data” (e.g. Cao, Diao, and Wu 2019). Here, we conducted a bibliometric study, searching the keyword “Geographically Weighted Regression” via Web of
Science (WoS), where in total, 2014 articles were found from 1999 to 2019, and their keywords are visualized in a word cloud form (Figure 1). Observe the frequency (size) of “Big Data”, which appears second only to “GWR”. Thus, the demand for high-performance solutions for GWR is clear, where its application in “Big Data” problems can be limited (Murakami et al. 2020), even with the employment of the existing solutions listed above (section 1.2).

1.1. Existing implementations of GWR

There are a number of standalone implementations with GWR enabled, such as GWR3 (Charlton, Fotheringham, and Brunsdon 2003), GWR4 (Nakaya et al. 2009), the GWR tool in ESRI ArcGIS (ESRI Corp 2011), and Multi-scale GWR (MGWR) (Li et al. 2019b). GWR is also available through scripting platforms with: the mgwr module of the PySal package in Python (Oshan et al. 2019); as part of the econometrics toolbox in MATLAB (LeSage and Pace 2009); and five R packages – spgwr (Bivand and Yu 2006), Geographically Weighted Models (GWmodel) (Lu et al. 2014b; Gollini et al. 2015), gwr (Wheeler 2013), McSpatial (McMillen 2015) and lctools (Kalogirou 2016). The five R packages considered as a whole provide the richest suite of GWR forms (e.g. conventional, robust, heteroskedastic, multiscale, space-time and more) and therefore development here is most appropriate. However, all suffer computationally, particularly given the strict memory limit for specific operation systems (R Core Team 2020). Workarounds to exceeding computational limits exist, such as coarse-scaling the observations, or the use of aggregations via upscaling (e.g. Yang et al. 2019) – all prior to a GWR fit, but none are ideal given important sources of information, fine scale detail and variability are lost.

1.2. Existing high-performance solutions

Efforts to improve the computational efficiency of GWR exist. Firstly, through Harris et al. (2010) who implemented a grid-based (parallelization) approach to conventional GWR. More recently, Li et al. (2019a) developed a python implementation (FastGWR) that optimizes the conventional GWR algorithm together with embedding multi-core parallel computing technology. This computational scheme has also been transplanted for use with multiscale GWR (Li and Fotheringham 2020). Wang et al. (2020) proposed a high-performance solution of GWR with the Compute Unified Device Architecture (CUDA), namely Fast-Parallel-GWR (FPGWR) which was developed with Microsoft Visual Studio 2015 and CUDA development kit. Finally, a mathematical approach was taken by Murakami et al. (2020) who proposed Scalable GWR (ScaGWR) that saves on computational overheads via the pre-compressing of large matrices and vectors with polynomial kernels. For ScaGWR, the computational cost presents a linear relationship with the sample size, while a quadratic order appears for the usual un-adapted GWR form. The ScaGWR routine can be found in the R package sggwr (Murakami et al. 2019) and GWmodel. ScaGWR provides approximate coefficient estimates in comparison with conventional GWR, where the results from ScaGWR might vary slightly when different parameters are specified – for example, the chosen degree or order of the polynomials (Murakami et al. 2020).

Figure 1. Word cloud of keywords from 2014 articles on GWR queried via WoS from 1999 to 2019.
1.3. Our study’s approach in R

In this sense, the computational bottleneck is still problematic for GWR (and its extensions) in the R environment, particularly for the geographically weighted functions in GWmodel. However, generic high-performance computing options have been incorporated in many packages since R release 2.14.0 (Eddelbuettel 2020), where grid computing, cloud computing, multi-core and Graphic Processing Unit (GPU) are commonly invoked. In this respect, this study investigates high-performance solutions for (conventional) GWR within the R package GWmodel, where our workflow consists of three hierarchies: 1) optimize the algorithm for a GWR calibration for accepting out-of-memory issues with “Big Data”; 2) adopt multi-thread parallel computing for a GWR calibration (GWR-MP), which enables analysis on a standard Personal Computer (PC) with a multi-core processor; 3) apply parallel computing on the GPU devices via CUDA (GWR-CUDA).

For performance evaluation, we compare the performances of the new solutions proposed (i.e. GWR-MP and GWR-CUDA) with existing solutions found in GWmodel, MGWR and FastGWR using varying sample sizes, where the latter two are outside of the R environment. We haven’t included FPGWR where CUDA was also adopted as: 1) the source code or tool is not available; and 2) key aspects of FPGWR are not clear, such as distance calculation, kernel function implementation, making FPGWR difficult to fully reproduce. This study is organized as follows. Firstly, we provide a description of conventional GWR methodology and the new high-performance techniques proposed. Secondly, competing high-performance solutions to GWR are objectively compared through a designed experiment. Thirdly, we summarize and suggest future research.

2. The GWR methodology and high-performance solutions

2.1. Basics of GWR

The conventional GWR model characterizes spatially varying relationships via location-specific regressions whose coefficients are estimated by (geographically) weighted least squares. The model can be expressed as (Brunsdon, Fotheringham, and Charlton 1996; Fotheringham, Brunsdon, and Charlton 2002):

$$ y_i = \hat{\beta}_0(u_i, v_i) + \sum_{k=1}^{l} \hat{\beta}_k(u_i, v_i)x_{ik} + \epsilon_i $$

(1)

where $y_i$ is the dependent variable at location $i$ on a two-dimensional space; $x_{ik}$ is the value of the $k$th independent variable at location $i$; $l$ is the number of independent variables; $\hat{\beta}_0(u_i, v_i)$ is the intercept parameter at location $i$; $\hat{\beta}_k(u_i, v_i)$ is the local regression coefficient for the $k$th independent variable at location $i$; $(u_i, v_i)$ are the spatial coordinates of location $i$; and $\epsilon_i$ is the independent random error at location $i$.

In line with Tobler’s first law of geography (Tobler 1970), extended to consider situations in which nearby regression relationships are more similar than distant ones, GWR consists of a series of local regressions where observations are weighted (i.e. given decreasing influence) via a distance-decay kernel function (Lu et al. 2014a). The estimator of the coefficients at location $i$ has the following matrix expression:

$$ \mathbf{\hat{\beta}}(u_i, v_i) = (X^T W(u_i, v_i) X)^{-1} X^T W(u_i, v_i) y $$

(2)

where $X$ is the matrix of the independent variables with a column of 1s for the intercept; $y$ is the dependent variable vector; $\mathbf{\hat{\beta}}(u_i, v_i) = \left( \hat{\beta}_0(u_i, v_i), \hat{\beta}_1(u_i, v_i), \ldots, \hat{\beta}_m(u_i, v_i) \right)^T$ is the vector of $m + 1$ local regression coefficients; $W(u_i, v_i)$ is a $n \times n$ diagonal matrix denoting geographical weights of each observation for calibrating the local regression at location $i$, and is defined as:

$$ W(u_i, v_i) = \begin{bmatrix} w_{i1} & 0 & \ldots & 0 \\ 0 & w_{i2} & \ldots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \ldots & w_{in} \end{bmatrix} $$

(3)

where $w_{ij}(j = 1, \ldots, n)$ is calculated via a kernel function decaying with respect to Euclidean distance, or some other distance metric (Lu et al. 2014a), between locations $i$ and $j$, and $n$ represents the number of observations. Gaussian, exponential, bi-square, box-car, tri-cube are among the many kernel functions that can be specified (Gollini et al. 2015), where an optimal kernel bandwidth is commonly found by leave-one-out cross-validation or by a corrected Akaike Information Criterion (AICc) procedure. The kernel bandwidth relays the chosen spatial scale of the regression relationships.

Diagnostics for a GWR model’s fit are essential, where $R$-squared, adjusted $R$-squared and AICc are commonly reported. These can be expressed as (Fotheringham, Brunsdon, and Charlton 2002):

$$ R^2 = \frac{\sum_i \left( \hat{y}_i - \bar{\hat{y}} \right)^2}{\sum_i \left( y_i - \bar{y} \right)^2} $$

(4)

$$ R_{\text{adjusted}}^2 = 1 - \left( 1 - R^2 \right) \frac{n - 1}{n - 2 \text{tr}(S) + \text{tr}(S^2) - 1} $$

(5)
\[
\text{AIC_c} = 2n \ln(\hat{\sigma}) + n \ln(2\pi) + n \left\{ \frac{n + \text{tr}(S)}{n - 2 - \text{tr}(S)} \right\}
\]

where \(\hat{y}_i\) is the fitted value at location \(i\); \(\hat{y}\) is the mean value of \(y\); \(\hat{\sigma}\) is the estimated standard deviation of the error term:

\[
\frac{2}{\sigma^2} = \frac{\sum_i (\hat{y}_i - \hat{y})^2}{n - 2\text{tr}(S) + \text{tr}(S^T S)} = \frac{y^T(I - S)^T(I - S)Y}{n - 2\text{tr}(S) + \text{tr}(S^T S)}
\]

where \(I\) is an \(n \times n\) identity matrix and \(\text{tr}(S)\) and \(\text{tr}(S^T S)\) denote the traces of the hat matrix \(S\) and \(S^TS\). For GWR, each row \(S_i\) of the hat matrix can be found as follows:

\[
S_i = X_i(X^TW(u_i; v_i)X)^{-1}X^TW(u_i; v_i)
\]

where \(X_i\) is its \(i^{\text{th}}\) row of the matrix \(X\) of independent variables.

Furthermore, \(t\) statistics at each individual regression point can be produced along with the coefficient estimates. For each estimated regression coefficient at location \(i\), \(\beta_k(u_i; v_i)\), the \(t\) statistic can be calculated by:

\[
t_{k,i} = \frac{\hat{\beta}_k(u_i; v_i)}{\text{SE}(\hat{\beta}_k(u_i; v_i))}
\]

where \(\text{SE}(\hat{\beta}_k(u_i; v_i))\) is the localised standard error of \(\beta_k(u_i; v_i)\). For each location-specific calibration, the standard errors are obtained from:

\[
\text{SE}(\hat{\beta}_i) = \hat{\sigma} \sqrt{\text{diag}(C_iC_i^T)}
\]

where

\[
C_i = (X^TW(u_i; v_i)X)^{-1}X^TW(u_i; v_i)
\]

The given calculations are commonly reported in most GWR software tools, where the algebraic matrix operations are programmed in a straightforward manner. However, their computational cost is expensive, particularly when dataset size (number of observations \(n\)) is large. Computational burden is primarily a consequence of: 1) complex matrix operations, particularly the \(n \times n\) matrices involved, like the hat matrix \(S\); and 2) a large number of matrix operations are repeated in the location-wise calibrations, kernel bandwidth optimization and when calculating the model fit diagnostics.

### 2.2 Reducing memory cost for GWR

A variety of GWR forms and extensions are present in GWmodel, making it the most comprehensive GWR R package (Comber et al. 2022). In early versions of GWmodel, all GWR functions were developed directly from the algebraic formulations in Section 2.1 above. This requires a number of \(n \times n\) matrices to be calculated and stored, specifically for calculating diagnostic information and enabling statistical inference (Leung, Mei, and Zhang 2000). Note here, that it is almost impossible to allocate as much as 2 GB to a single vector in a 32-bit or 64-bit build of R due to predefined allocations of address space on Windows (R Core Team 2020). Allocating memory for a \(16,000 \times 16,000\) numeric matrix in R will normally be an upper limit. This means the maximum \(n\) for any of the conventional GWR functions in R is around 16,000. However, in practice, the maximum number of observations a conventional GWR tool can handle is likely to be much smaller (i.e. \(n \ll 16,000\)).

It is therefore necessary to first relieve these memory constraints when developing high-performance solutions for GWR, and to support GWR analyses of very large datasets. In this respect, Li et al. (2019a) optimized the calculations of AICc and localised standard errors by avoiding the storage of the entire hat matrix, which reduced the memory storage size from \(O(n^2)\) to \(O(nm)\). This strategy of avoiding any \(n \times n\) matrix operation or storage is effective and makes it workable when dealing with a large dataset on any basic PC. Therefore, as a potential approach for reducing memory costs, we re-formulated the algebraic operations of a GWR calibration, as follows, which are essentially the same as the optimizations proposed by Li et al. (2019a).

First observe that Equation (2) can be divided into the following two parts:

\[
X^TW(u_i; v_i)X = \begin{pmatrix}
X_i^T & \cdots & X_n^T
\end{pmatrix}
\begin{pmatrix}
w_{i1} & \cdots & 0 \\
\vdots & \ddots & \vdots \\
0 & \cdots & w_{in}
\end{pmatrix}
\begin{pmatrix}
X_i \\
\vdots \\
X_n
\end{pmatrix}
\]

\[
= \sum_{j=1}^{n} w_{ij}X_j^TX_j
\]

\[
X^TW(u_i; v_i)y = \begin{pmatrix}
X_i^T & \cdots & X_n^T
\end{pmatrix}
\begin{pmatrix}
w_{i1} & \cdots & 0 \\
\vdots & \ddots & \vdots \\
0 & \cdots & w_{in}
\end{pmatrix}
\begin{pmatrix}
y_i \\
\vdots \\
y_n
\end{pmatrix}
\]

\[
= \sum_{j=1}^{n} w_{ij}y_jX_j^T
\]

where \(X_j\) is the \(j^{\text{th}}\) row of the \(X\) matrix. In this sense, the point-wise estimator of GWR can be regarded as a cross manipulation between the inverse of a \((m+1) \times (m+1)\) matrix and a \((m+1) \times 1\) vector. Accordingly, the weight matrix \(W(u_i; v_i)\) can only be stored as a vector with its diagonal elements.
For diagnostics of GWR, more complicated computations are involved, particularly for the $n \times n$ matrices, including hat matrix $S$, its square $S^2S$ and the matrix $Q = (I - S)^T(I - S)$. As shown in Equations (5–7), the traces of $S$ and $S^2S$ are needed, but where they could be found in these two steps:

$$\text{tr}(S) = \sum_{i=1}^{n} S_{ii} = \sum_{i=1}^{n} X_i C_i^T$$

$$\text{tr}(S^2S) = \sum_{i=1}^{n} S_i S_i^T$$

where $C_i$ means the $i^{th}$ column of matrix $C$. Moreover, the matrix $Q$ can also be expressed as follows:

$$Q = (I - S)^T(I - S) = \sum_{j=1}^{n} (e_i - S_i)^T(e_i - S_i)$$

where $e_i$ is the $i^{th}$ row of the identity matrix $I$. Observe the matrix $Q$ is also required in many statistical tests for GWR (i.e. for spatial non-stationarity), such as the $F$-tests proposed by Leung, Mei, and Zhang (2000) and Fotheringham, Brunsdon, and Charlton (2002), which are similarly included in most GWR software tools (GWR3, GWR4, as well as GWmodel). In this sense, it is natural for these $F$-tests to benefit from the high-performance solutions proposed.

According to the above equations, the storage of all $n \times n$ matrices required for a GWR calibration and associated diagnostics can be avoided, by storing only vectors of length $n$ and matrices of size $n \times (m + 1)$ in the location-wise computations. Thus, the memory cost of GWR can be similarly reduced to $O(nm)$, essential for working with “Big Spatial Data” in R. In the current release of GWmodel, the GWR functions have already been optimized in this respect. Therefore, for this study, the next steps are an assessment of high-performance solutions embedded in paralleling computing techniques.

### 2.3. Parallelization solutions for GWR

The two parallelization solutions adopted were: (a) multcore Central Processing Unit (CPU) and GPU accelerator via multithreading parallel (GWR-MP) and (b) CUDA (GWR-CUDA), respectively.

As illustrated in Figure 2, the procedure of GWR-MP was carried out in the following steps:

1. Create the coefficient matrix $\beta_{p \times (m+1)}$ and the vectors of $n$ dimensions ($S_i$) for recording the diagonal elements of $S^2S$ if diagnostic information is calculated;
2. Create $c^2$ threads, and divide the $n$ point-wise operations into them, i.e. $n_i$ operations are conducted on the $i^{th}$ thread, where $\sum_{i=1}^{n_i} n_i = n$;
3. For each thread, create a vector of $n$ dimensions ($S_i$) and a vector $Q_i$;
4. Carry out the following operations for each location $i$:
   a. Calculate the weight vector $w_i = (w_{i1}, \ldots, w_{im})$ from the corresponding distances of the observations from the location $i$;
   b. For estimating $\beta$, calculate Equation (2) in two parts, i.e. Equation (12);

![Figure 2. Parallel computing flowchart of the GWR-MP algorithm.](image-url)
c. Calculate the $i^{th}$ row of the hat matrix and assign it to $S_i$ in memory; then renew the $i^{th}$ element of $S_i$ as $S_i = S_i S_i^T$;

d. Renew the vector $Q_i = (e_i - S_i)^2$.

(5) Repeat Step 4 until all the location-wise operations are finished, and the coefficient estimates $\beta_{n \times m}$, $\text{tr}(S)$, $\text{tr}(S^T S)$ and $\hat{\sigma}^2$ are ready for the final output.

By contrast, the parallelizing strategy for GWR-CUDA is designed specifically to fit GPU devices. As illustrated in Figure 3, the detailed procedure of GWR-CUDA includes the following steps:

(1) Read the data matrices or vectors (i.e. $X, Y$ and coordinates) from memory into GPU;

(2) Divide the $n$ point-wise operations into groups, and within each group, $g$ or fewer point-wise calculations are conducted in parallel, where $g$ should meet the following condition:

$$bgkn + 2bgkk + bgk \leq M - \alpha$$

where $b$ is the number of bytes needed for each element in the matrix or vector (commonly set as 8), $M$ is the GPU memory size, $\alpha$ is the memory reserved for intermediate calculations. In practice, the number of variables $k$ is far less than the number of observations, $n$ (i.e. $k \ll n$), the number of $g$ is more dependent on the term, $bgkn$.

(3) Create arrays $A_{g \times k \times m}$, $\Omega_{g \times k \times k}$ and a matrix $\xi_{g \times k}$, and conduct the following location-wise calibrations ($i = 1, \ldots, g$) in parallel within the current group:

a. Calculate the weight vector $w_i = (w_{i1}, \ldots, w_{in})$ from the corresponding distances of the observations from the location $i$;

b. Calculate $X^T W_i = \sum_{j=1}^{n} w_{ij} X_j^T$ and assign it to the $i^{th}$ $k \times n$ component of $\Delta_i$;

c. Calculate $X^T W(u_i, v_i) X = \sum_{j=1}^{n} w_{ij} y_j X_j^T$ and assign it to the $i^{th}$ $k \times k$ component of $\Omega$, and then calculate its inversion;

d. Calculate $X^T W(u_i, v_i) y = \sum_{j=1}^{n} w_{ij} y_j X_j^T$ and assign it to the $i^{th}$ row of $\xi_i$;

e. Calculate the location-wise coefficient estimate $\beta_i$, $S_i$, $S_i S_i^T$ and $(e_i - S_i)^2$.

(4) Repeat Step 3 until parallel computations for all the groups are finished, and the coefficient estimates $\beta_{n \times m}$, $\text{tr}(S)$, $\text{tr}(S^T S)$ and $\hat{\sigma}^2$ will be ready for the final output.

It is important to note that both GWR-MP and GWR-CUDA procedures are designed to include the GWR model’s diagnostic information calculations, with the calibration points the same as the observations and an assessment of model fit is required (which includes kernel bandwidth optimisation). Otherwise, the above procedures would be greatly simplified with only coefficient estimates returned. We implemented GWR-MP and GWR-CUDA in R, coded the parallel part via C++ and wrapped them via the Rcpp package (Eddelbuettel 2013).

3. Experimental design

3.1. General information

For this study, we compared the computational performances of GWmodel (version 2.1–4), FastGWR (updated on 12 August 2019), MGWR (version 2.1.1), GWR-MP and GWR-CUDA for

![Figure 3. Parallel computing flowchart of the GWR-CUDA algorithm.](image-url)
implementations of conventional GWR only to ensure the same results. As shown in Table 1, we adopted two devices for running GWR where both devices had a GPU specification for running GWR-CUDA.

In terms of experimental data, we produced a series of simulated datasets of size \( n \) ranging from 1000 to 10,000 with increments of 1000, from 10,000 to 100,000 with increments of 10,000, and from 100,000 to 1,000,000 with increments of 100,000. For MGWR, FastGWR and GWR-MP, we specified a different number of cores ranging from 2 to 48 to run them in parallel. We didn’t test them with all the combinations, but selectively adopted 2, 3, 4, 5, 6, 7, 8, 12, 24, 36 and 48 cores for typical tests. Note that the number of physical cores on the experimental device is 24, but the number of logic cores could be up to 48 through the hyper-threading technology. Notably, the current GWR routine in GWmodel is a serial program, so that the setting of multicores will not work differently for it. As shown in Figure 3, we adopted a different computing strategy for GWR-CUDA, where all samples are divided into groups, and location-wise calibrations within each group are conducted in parallel on the GPU device. Thus, the number \( g \) is the parallel computation counts for executing GWR-CUDA, and we took \( g = 384 \) (or less if samples were insufficient for the final group) according to Equation (16). For each sample size \( n \) and GWR implementation, 10 experiments were conducted independently on the two devices, respectively. Moreover, samples sizes ranging from 100,000 to 1,000,000 are adopted only for the extreme performance tests of GWR-CUDA, not for tests on the other four GWR solutions, as relatively inefficient solutions result due to unacceptably long time frames.

### 3.2. Performance indicators

Two indicators were used to evaluate performance – the average time cost and the “speedup” of the parallel computations. For each sample size \( n \) and GWR implementation, the average time cost was calculated as follows:

\[
\bar{T}_{n,GWR} = \frac{\sum_{i=1}^{m} T_{i,GWR}}{m}
\]

where \( T_{i,GWR} \) represents the time cost of running the \( j \)th GWR implementation with a sample size \( n \), \( m \) (in this case, taken as 10) is the number of individual experiments and \( \bar{T}_{n,GWR} \) refers to the average time cost. Note that in all cases, the time costs include both the (automated) kernel bandwidth optimization (by AICc) and the GWR model calibration.

Speedup is an important indicator to evaluate the performance of parallel computations (Hill and Marty 2008). According to its original definition, we take a simple expression for its calculation, as:

\[
k = \frac{T_S}{T_M}
\]

where \( k \) is the speedup, \( T_S \) is the time cost of serial computing and \( T_M \) is the time cost of parallel computing with multi-cores. For this study, we repeated 100 independent experiments for each scenario, meaning speedup could be calculated using average time costs from \( m \) individual runs, i.e.:

\[
k = \frac{\bar{T}_S}{\bar{T}_M}
\]

where \( \bar{T}_S \) and \( \bar{T}_M \) are the average time costs of serial and parallel computations, respectively. We can verify that the estimation of speedup is significantly valid and reliable, by assuming the time cost for each individual test is a random variable subject to a normal distribution.

### 4. Results and discussion

In Figure 4, we present the averaged time costs of GWmodel, FastGWR, MGWR, GWR-MP and GWR-CUDA with a different number of cores with samples of sizes ranging from 1000 to 100,000. As the GWR implementation in GWmodel is a serial program, the time cost will not be affected by increasing cores, but averaged time costs grow exponentially as sample size increases. This indicates that the basic GWR function in the latest release of GWmodel is not working efficiently with a large dataset, even though the function
benefits from algorithmic optimization and code implementation with C++. It can handle a relatively large dataset, say greater than 100,000, but its running time will be incredibly long, particularly when bandwidth optimization is additionally conducted. MGWR is applicable for running on multi-cores, but it failed

Figure 4. Average time costs of GWmodel, FastGWR, MGWR, GWR-MP and GWR-CUDA with a different number of cores.
to calibrate the GWR models with sample sizes greater than 60,000. In the limited number of visible tests, it performed similarly to FastGWR, which is also developed via Python by the same research team (Oshan et al. 2019; Li et al. 2019a). Thus, results for MGWR can be represented by those for FastGWR and are not discussed further.

Both FastGWR and GWR-MP are naturally designed for multi-core parallelism. From Figure 4, FastGWR and GWR-MP always outperform the serial routine in GWmodel, and these advantages grow as the number of cores increase and as sample size increases. From Figure 4, GWR-MP performs similarly to FastGWR in most cases, but where GWR-MP tends to outperform FastGWR for samples greater than 60,000. Relative to GWR-MP, the time costs of FastGWR become exponentially large when the number of cores exceeds 24 for sample sizes greater than 60,000, which means that all the physical cores will be employed and the logic cores will be used via the hyper-threading technology. On this condition, the performance and stability of each physical core could worsen due to frequent switches between two logic cores on each physical core. In addition, FastGWR was developed with the Message Passing Interface (MPI), a standard and portable message-passing system for parallel programming (Dalcín et al. 2008). The MPI was originally designed for distributed memory systems, then extended to shared memory parallel computing for effectively utilizing node-level architecture (i.e. stand-alone machine with multi-cores). Its communication efficiency could be more or less affected by the memory capacity pressure, particularly when all the cores are fully occupied (Brinskiy, Lubin, and Dinan 2015). That could be the main reason of the relatively weak performance of FastGWR when the number of cores exceeds 24. From Figure 4, GWR-CUDA consistently performs the best of all across all scenarios.

For critically testing GWR-CUDA, we extend the size of samples up to one million with two different versions of GPU devices. In Figure 5, we present the average time costs of GWmodel and GWR-CUDA. The time consumption of GWmodel increases exponentially, particularly when the sample size is larger than 8000; in contrast, the time cost of GWR-CUDA grows much more slowly as sample size gets larger, but a dramatic increase occurs for sample sizes of around 800,000 or more. The two versions of GPU devices present different performances for running GWR-CUDA, dealing with samples of one million for around 3.5 h (12,766 s) on GPU-1, and around 5 h (17,828 s) on GPU-2. As one of the world’s most advanced GPU, NVIDIA® Tesla® V100 (GPU-1) renders a great advantage over the GeForce RTX 2060 Mobile (GPU-2), a mobile graphics chip embedded in a laptop. Given that a laptop cannot run stably with full capacity for a long period, we only tested GWR-CUDA on GPU-2 with samples of sizes ranging from 1000 to 100,000, and

Figure 5. Average time costs of GWmodel and GWR-CUDA with different sample sizes.
1,000,000 only. Moreover, results indicate that the physical parameters of the CPU and the GPU device will affect the performances of the chosen high-performance solutions. Equipment (laptop or PC) with high-end CPU or GPU devices will provide better performances, and where High-

Figure 6. Speedup indicators of GWmodel, FastGWR, MGWR, GWR-MP and GWR-CUDA with a different number of cores.
Performance Computing (HPC) infrastructure should provide a better scope for potential improvement in this aspect.

The average time costs could be specific for the devices adopted, and almost impossible to be reproduced with a different device. From an objective assessment, we used the speedup indicator to evaluate how improvements benefited from the parallel strategies implemented in FastGWR, MGWR, GWR-MP and GWR-CUDA. A larger speedup means the parallel solution for a specific GWR implementation makes a greater optimization in computational efficiency than running it serially. As shown in Figure 6, the performance of GWR-MP and FastGWR (and MGWR) improves as more cores are used for running them in parallel. Again GWR-MP demonstrates better usage of multi-core equipment for samples of sizes ranging from 3000 to 10,000, while performance does not show an improvement when the number of cores exceed 24, i.e. all the physical cores are fully occupied. For GWR-CUDA, its superiority in parallel performance is apparent when the sample size is greater than 10,000, but note that the speedup falls sharply to 40 when the sample size reaches 90,000. In Figure 5, we can see that the average time costs of GWR-CUDA also increase exponentially as the sample size becomes large, where a sharper increase occurs around a sample size of 80,000.

Note in the inset figure of Figure 5, GWR-CUDA is not always the best performer in comparison with the serial solution in GWmodel. GWR-CUDA takes more time than the serial solution when the sample size is less than 3000. To implement GWR-CUDA, all the pre-defined data matrices or vectors (i.e. $X$, $Y$ and coordinates) are transferred from memory into the GPU, and the results, including hat matrix $S$ and coefficient estimates $\hat{\beta}$ are transferred from GPU back to memory – widely known as I/O issue important for GPU performance (Fuji et al. 2013). In other words, the I/O cost is predominant when the sample size is less than 3000, and the computational advantage of GWR-CUDA starts to emerge when the size is getting larger than 3000. The I/O cost could be affected by the physical parameters of GPU, CPU and protocol type, so GWR-CUDA will perform differently with different devices. Thus, the critical value (i.e. 3000 in this study) could fluctuate marginally width different computational configurations.

The results also reveal a fact that the high-performance solutions would be not be recommended for samples with relatively small sizes, say less than 5000, the most common data volume in the previous GWR applications. On the flip side, GWR applications with a relatively large data set (e.g. large than 20,000) were rarely found due to the lack of and universal access to high-performance tools. The findings in the context of rich scenarios are beneficial to both development and optimization of the high-performance solutions.

5. Summary

In this study, we have proposed two high-performance solutions for GWR via multi-core parallel and CUDA techniques: GWR-MP and GWR-CUDA, respectively. We objectively compared them with existing GWR implementations found in GWmodel, MGWR and another high-performance solution FastGWR. Results indicate that no solution was always the best in terms of computational efficiency, as summarized in Figure 7 by their relative speeds for four sample size intervals (less than 2000; greater than 2000, but less than 10,000; greater than 10,000 but less than 100,000; greater than 10,000). As (effectively) serial solutions, both GWmodel and MGWR provide adequate GWR implementations for (small) sample sizes $< 10,000$, as computational costs were considered acceptable.

For multi-core parallel solutions, GWR-MP provided a commensurate solution with GWR-CUDA for dealing with (large) sample sizes between 10,000 to 100,000 on a computer of common multi-core configuration, where GWR-MP demonstrated more efficient computing capacity.
for each core or thread than FastGWR, whose design is more suited to non-shared memory clusters. For example, Li et al. (2019a) adopted FastGWR with a dataset of 1.28 million points on a 512-core computing cluster. However, high-performance computing clusters are usually too expensive and too few in number to be accessed by many researchers.

Conspicuously, GWR-CUDA provided a relatively cheap but highly efficient solution for analyzing a very large dataset, of which the size could be much larger than 1,000,000, the upper number in this study. The study GPU (NVIDIA GeForce RTX 2060 (Mobile)) only cost around $350, but we found we could implement a GWR model (including bandwidth optimization and model calibration) with one million data points in around 5 h. A better configuration of the GPU, like with NVIDIA Tesla V100 could reduce this time to 3.5 h, but at a cost of around $9000. Note however, GWR-CUDA should only be preferred when sample size is very large in terms of balancing cost with speed (as clearly seen in Figure 7). Note, however the Figure 7 roughly show the comparative performances of these solutions, and could more or less vary when different devices adopted.

Both GWR-MP and GWR-CUDA were implemented in R with wrappers on the C++ code, which has been incorporated into the latest release of GWRmodel (say version GWRmodel_2.2–8). Note that, nowadays it is straightforward to execute R from Python, and vice versa. Therefore, this is not a black-or-white type of choice to run these solutions in R or Python. Moreover, all the C++ code could be easily transferred to a standalone application, which we are currently working on. Inspired by Figure 7, an important feature of this, is to adaptively set a computational strategy according to sample size and the computing environment, and this study provides a direct support for such a strategic optimization. An ultimate solution could be an application developed under the service-oriented architecture with powerful computers or clusters, and the algorithms proposed here would provide fundamental support. Moreover, the solutions proposed here are directly applicable to extended GWR forms beyond the conventional GWR form, such as GTWR; and also directly applicable to other geographically weighted models (Lu et al. 2014b) outside of those for regression (e.g. GW PCA). Further, more pertinent issues, such as robust statistical inference in GWR with a massive data set (Griffith 2015) would also be worthy of investigation.

Notes
1. Note that the diagnostic information cannot be calculated when an individual set of regression locations are adopted.

2. Theoretically, the number of threads $c$ could be larger than the number of cores available, but we would suggest creating no more than the number of cores for ensuring the performance of each thread.

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Data availability statement
The data that support the findings of this study are available with the identifier(s) at the link (https://figshare.com/s/13f325af1e37c3bc15fc).

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