Evaluating Geographically Weighted Regression Models for Environmental Chemical Risk Analysis

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ABSTRACT: In the evaluation of cancer risk related to environmental chemical exposures, the effect of many correlated chemicals on disease is often of interest. The relationship between correlated environmental chemicals and health effects is not always constant across a study area, as exposure levels may change spatially due to various environmental factors. Geographically weighted regression (GWR) has been proposed to model spatially varying effects. However, concerns about collinearity effects, including regression coefficient sign reversal (ie, reversal paradox), may limit the applicability of GWR for environmental chemical risk analysis. A penalized version of GWR, the geographically weighted lasso, has been proposed to remediate the collinearity effects in GWR models. Our focus in this study was on assessing through a simulation study the ability of GWR and GWL to correctly identify spatially varying chemical effects for a mixture of correlated chemicals within a study area. Our results showed that GWR suffered from the reversal paradox, while GWL overpenalized the effects for the chemical most strongly related to the outcome.

KEYWORDS: environment, GWR, GWL, lasso, chemical mixtures, reversal paradox, cancer risk

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INTRODUCTION

Humans are exposed to mixtures of chemicals that may be influential for cancer risk. For example, risk of non-Hodgkin lymphoma (NHL) is suspected to be associated with several chemicals through environmental or occupational routes of exposure, and geographic variation in NHL rates suggests the importance of environmental risk factors. Positive associations have been found with persistent organochlorine chemicals, including polychlorinated biphenyls (PCBs), particularly PCB congener 180, and dichlorodiphenyldichloroethylene.

Environmental exposure patterns are typically complex with inherent correlations among co-occurring chemicals or their metabolites. For example, many PCB congeners exhibit a high degree of correlation. Important questions in the analysis of mixtures include whether and how the health effect of one chemical should be adjusted for other chemicals present, even when those chemicals are highly correlated. Furthermore, the relationship between environmental chemicals and health effects (eg, cancer risk) is not always constant across a study area. Exposure levels may be different spatially due to environmental factors. For example, pesticide levels measured in house dust may be higher in agricultural communities (eg, in Iowa) or those in temperate climates where more pesticides are applied throughout the year (eg, Los Angeles) compared to the levels in urban locations (eg, Detroit). Acknowledging the principle that “the dose makes the poison,” the risk of adverse health effects such as NHL is greater in regions where exposure is higher. Thus, environmental health models that account for these spatially changing exposure/risk regions can be informative.

Models with spatially varying coefficients include geographically weighted regression (GWR), which is similar to local linear regression (eg, references 8–10) in that both methods use a kernel function to calculate weights that are applied to observations in a series of local weighted regression models. One issue with GWR is that GWR models have been found to be affected by local collinearity. Local collinearity in weighted explanatory variables can lead to GWR coefficient estimates that are correlated locally and across space, have inflated variances, and are at times counterintuitive and contradictory in sign to the global regression estimates, ie, evidence of the reversal paradox.
To illustrate, Wheeler and Tiefelsdorf\(^1\) highlighted the issue of collinearity in GWR in a simple model to explain white male bladder cancer mortality rates (1970–1994) in the 508 State Economic Areas of the US. Their model consisted of two explanatory variables: population density, a proxy for environmental and behavioral differences in urban/rural life, and lung cancer mortality rates, a proxy for the risk factor smoking, a known risk factor for bladder cancer. These two variables had a global correlation estimate of \(-0.59\); however, local correlation estimates were generally more extreme (ie, more strongly negative; median = \(-0.63\); Q3 = \(-0.71\) as approximated from their Fig. 4), with strongest inverse association in parts of Northeastern and Midwestern US (their Fig. 3). The resulting maps of GWR coefficients for population density and the smoking proxy showed a clear inverse map pattern. When the local smoking proxy parameter was high (primarily in the West and Northeast), the local population density parameter was negative. When the local smoking proxy parameter was negligible, the population density parameter was large and positive (primarily in the Midwest and Southeast). As noted by Wheeler and Tiefelsdorf,\(^1\) the important question is whether this complementary relationship in the parameters is real, meaningful, and interpretable, or whether it is an artifact of the statistical method. The natural research question is whether such inverse patterning in regression coefficients is an example of the reversal paradox\(^16\) due to strong local correlations between the two variables.

According to the reversal paradox, the association between two variables can be reversed, diminished, or enhanced when another variable is statistically controlled for.\(^16\) For example, consider two explanatory variables, \(x_1\) and \(x_2\), where the bivariate correlation between \(x_1\) and \(y\) is 0.2, and between \(x_2\) and \(y\) is 0.1. Figure 1 presents the standardized beta coefficients in the multiple regression model \(\mu = \beta_0 + \beta_1 x_1 + \beta_2 x_2\). As the correlation between the variables increases, the regression coefficient associated with \(x_1\) increases and the coefficient associated with \(x_2\) becomes large and negative – which could lead to a misleading interpretation of the association between \(x_2\) and \(y\). Use of statistical models with correlated data may produce consistent, replicable, yet erroneous results.\(^16\)

To address the issue of collinearity with GWR and to limit its effects, the geographically weighted lasso (GWL) adds a constraint on the magnitude of the estimated regression coefficients.\(^14\) The GWL also performs local model selection by potentially shrinking some of the estimated regression coefficients to zero in some locations of the study area, thereby diminishing the adverse effects of the correlation pattern. However, when accurate variable selection is the focus of the analysis, such a strategy makes it difficult to determine whether a variable was excluded from the model due to a lack of association with the outcome or due to its correlation with variables in the model.

Our objective in this study is to evaluate the impact of collinearity of the geographically weighted regression models GWR and GWL in a chemical exposure and risk assessment context. We use a simulated data set for which the truth is known and further assess the ability of GWL to control collinearity effects, such as the reversal paradox, when the effects of correlated environmental chemicals are of interest. We begin by describing the process used to simulate data that we propose are environmentally relevant – ie, regions with low exposure and regions with higher exposures and where different chemicals may have related exposure patterns but not necessarily the same association with a health effect of interest. We conduct GWR and GWL analyses in a scenario with independent chemicals and a scenario with correlated chemicals.

**Methods**

Simulating spatially varying exposure- and dose-dependent association with an outcome. Consider the scenario in which there are three predictor variables (eg, environmental chemicals) that vary over space in a study area (Fig. 2). We assumed that the first predictor variable, \(x_1\), was present at high enough levels to be associated with an increase in the mean response in the upper region of the study area, while being present only at background levels where there is no increase in mean response in the lower region of the study area. Furthermore, we assumed that both \(x_2\) and \(x_3\) were present at uniform levels across the study area and that \(x_2\) was not related to the response variable, while the relationship between \(x_3\) and the response was moderate. Additionally, we considered two cases for the relationships among the predictor variables: Case 1, where the predictor variables were independent (ie, multivariate normal with zero correlation), and Case 2, where the...
predictor variables were correlated (ie, $\rho_{12} = 0.7$, $\rho_{13} = 0.3$, and $\rho_{23} = 0$). We used a unit grid as the study area and divided the grid into three equal-sized rows. A total of 500 locations were randomly generated inside the study area, and locations falling in the upper one-third of the study area were defined as belonging to Region 1 ($n_1 = 160$), while locations falling in the lower two-thirds of the study area were defined as belonging to Region 2 ($n_2 = 340$).

For each case, multivariate normal data were simulated separately for Region 1 and Region 2. We assumed that the levels of $x_1$ were the highest in Region 1 (mean of 3.0) and negligible in Region 2 (mean of 0.1). We further assumed that the mean of both $x_2$ and $x_3$ was constant (mean of 0.1) across the entire study area. In the case of independence, an identity matrix was used for the covariance, while for the correlated case, the aforementioned correlation pattern was imposed. To simulate the corresponding mean related to the three predictor variables, we used the following nonlinear threshold model:

$$
\mu = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3
$$

with parameters defined as $\beta_0 = \beta_3 = 0$, $\beta_1 = 2$, $\beta_2 = 1$, and $\delta = 2$. The response variable, $y_i$, was generated by adding a standard normal error term to the mean. Using this model, we imposed that $x_1$ was active in Region 1 and inactive in Region 2. More specifically, we allowed $x_1$ to be present at high enough levels to be associated with an increase in mean response in Region 1, while being present only at background levels (ie, less than the threshold) and not associated with the mean response in Region 2. This specification effectively removed $\beta_1$ from the model in Region 2, with $\beta_1 = 0$ for almost all of the locations in Region 2 for both the correlated and uncorrelated cases. The parameter $\beta_1$ was equal to 2.0 in the majority of locations in Region 1. Hence, there was a simple spatially varying relationship for $x_1$ and the outcome variable. Finally, we imposed that $x_2$ was not related to the response variable, while the relationship between $x_3$ and the response was moderate and uniform across the study area. A total of 100 data sets of size $N = 500$ were generated for each case, and the results are later presented aggregated over the 100 simulated data sets.

GWR model. In GWR, the spatial coordinates of data are used in the calculation of distances that are input into a kernel function to determine weights for spatial dependence among observations. Local regression models are related through shared data, but the dependence between regression coefficients at different locations is not specified. For example, consider $n$ observations measured at different locations. The GWR model at location $i$ is represented as follows:

$$
y_i = X_i \hat{\beta} + \epsilon_i
$$

where $y_i$ is the dependent variable at location $i$, $X_i$ is the column vector of explanatory variables at location $i$, $\hat{\beta}$ is the row vector of regression coefficients at location $i$, and $\epsilon_i$ is the random error at location $i$. The vector of estimated regression coefficients at location $i$ is

![Figure 2. Plots of average simulated concentration values across 100 simulated data sets over a square study area for two scenarios: independent chemicals (Case 1) and correlated chemicals (Case 2).](image)
\( \hat{\beta}_i = \left( X^T W X \right)^{-1} X^T W y \)  \hspace{1cm} (3)

where \( X \) is the design matrix of explanatory variables; \( W_i \) is the diagonal weights matrix that is calculated for each location \( i \) and applies weights to observations \( j = 1, \ldots, n \); and \( y \) is the vector of dependent variable values. Examples of kernel functions for defining the weight matrix include the Gaussian function, the bi-square nearest-neighbor function, and the exponential function, used herein. The weight from the exponential kernel function between any location \( j \) and the model location \( i \) is calculated as

\[
W_{ij} = \exp \left( - \frac{d_{ij}}{\phi} \right) \hspace{1cm} (4)
\]

where \( d_{ij} \) is the distance between locations \( i \) and \( j \), and \( \phi \) is the kernel bandwidth parameter.

**GWL model.** The lasso is defined\(^{17} \) as,

\[
\hat{\beta}_L = \arg \min_\beta \sum_{i=1}^n \left( y_i - \beta_0 - \sum_{k=1}^p \beta_k x_{ik} \right)^2 + \lambda \sum_j |\beta_j| \hspace{1cm} (5)
\]

where \( \lambda \) controls the amount of shrinkage of the regression coefficients, the value of which is chosen through algorithms such as least-angle regression (LARS)\(^{18} \) to find the lowest root-mean-square prediction error (RMSPE). Wheeler\(^{15} \) extended lasso to a geographically weighted version by defining a weighted \( X \) matrix as

\[
X_{W} = W^{1/2} X \hspace{1cm} (6)
\]

and estimating a lasso model with the LARS algorithm corresponding to each of the \( i \)th locations, \( i = 1, \ldots, n \).

**Evaluation of models.** The focus of the study was to determine whether the methods were able to correctly detect a strong relationship between \( x_1 \) and the mean response (\( \hat{\beta}_1 = 2 \)) in the upper third of the study grid and a moderate but uniform relationship between \( x_2 \) and the mean response (\( \hat{\beta}_2 = 1 \)) over the entire study area. Additionally, we were also interested in whether or not the methods can correctly discern that there is 1) no relationship between \( x_1 \) and the mean response in the lower two-thirds of the study grid and 2) no relationship between \( x_2 \) and the mean response over the entire study area. To evaluate the performance of GWR and GWL in identifying the spatially varying patterns in the coefficients, we started by mapping the average of the coefficient estimates at each location over the study area for both methods.

For each model, we calculated the root-mean-square error (RMSE) from estimation, the RMSPE, and the \( R^2 \) value. The RMSE is defined as

\[
RMSE = \sqrt{\frac{1}{n} \sum_{i=1}^n \left( y_i - \hat{y}_i \right)^2} \hspace{1cm} (7)
\]

while RMSPE is defined as

\[
RMSPE = \sqrt{\frac{1}{n} \sum_{i=1}^n \left( y_i - \hat{y}_i \right)^2} \hspace{1cm} (8)
\]

where \( \hat{y}_i \) is the predicted value of observation \( i \) with location \( i \) left out of the estimation data set. We then described these summary statistics using the median and interquartile range (IQR) over the 100 simulations.

To evaluate the performance of GWL in terms of variable selection, the percentages of coefficient estimates that were positive, negative, or zero were calculated by region for each simulated data set. We summarized the results across the simulated examples using medians and IQRs. Because GWR does not perform variable selection, we calculated the percentage of coefficient estimates that were positive and negative within each region. Additionally, in an effort to further evaluate the performance of GWR, we approximated the variance of the estimated GWR regression coefficients and created confidence intervals for the estimates at each location based on one and two standard errors (SEs) (ie, \( \hat{\beta}_k \pm \text{SE}(\hat{\beta}_k) \)) and \( \hat{\beta}_k \pm 2 \text{SE}(\hat{\beta}_k) \), for the \( i = 1, \ldots, n \) locations, and \( k = 0, \ldots, p \) parameters. The estimates were then classified as positive if the confidence interval was above zero, negative if the confidence interval was below zero, and zero (negligible) if the confidence interval contained zero. The covariance of the estimated regression coefficients was approximated\(^{19} \) as

\[
Var[\hat{\beta}_i] = (X^T W X)^{-1} X^T W \left( (X^T W X)^{-1} X^T W \right)^T \sigma^2 \hspace{1cm} (9)
\]

where the estimated error variance, \( \hat{\sigma}^2 \), is given as

\[
\hat{\sigma}^2 = \frac{\sum_{i=1}^n \left( y_i - \hat{y}_i \right)^2}{n - (2 \text{trace}(H) - \text{trace}(H^T H))} \hspace{1cm} (10)
\]

with the \( i \)th row of the hat matrix defined as

\[
H_i = X_i (X^T W X)^{-1} X^T W_i \hspace{1cm} (11)
\]

**Results**

The average observed concentration levels across the 100 simulated examples are plotted over the study area for each case in Figure 2, wherein we see that the average levels of \( x_2 \) and \( x_3 \) are uniform over the study area, while the mean level for \( x_1 \) is higher in Region 1 (the upper one-third of the grid space) as desired. The observed means for both the predictor and response variables are consistent with the study design and are summarized by case and region in Table 1. The observed correlation patterns were also consistent with the study design (results not shown).

The summary statistics across the 100 data sets are listed in Table 2. GWL outperformed GWR in terms of RMSPE
Geographically-weighted regression models

Table 1. Average predictor and response values across the 100 simulated data sets for the cases of independent chemicals (Case 1) and correlated chemicals (Case 2).

| Case 1 | Region 1 | Region 2 |
|---|---|---|
| $X_1$ | 3.00 | 0.10 |
| $X_2$ | 0.11 | 0.11 |
| $X_3$ | 0.09 | 0.10 |
| $Y$ | 5.63 | 0.23 |

| Case 2 | Region 1 | Region 2 |
|---|---|---|
| $X_1$ | 3.00 | 0.10 |
| $X_2$ | 0.11 | 0.10 |
| $X_3$ | 0.11 | 0.11 |
| $Y$ | 5.62 | 0.25 |

Table 2. Median (interquartile range) of summary statistics for GWR and GWL models across the 100 simulated data sets for the cases of independent chemicals (Case 1) and correlated chemicals (Case 2).

| Case 1 | GWR | 1.4 (1.4, 1.5) |
|---|---|---|
| RMSE | 1.0 (1.0, 1.2) |
| $R^2$ | 0.9 (0.9, 0.9) |
| GWL | 1.2 (1.1, 1.2) |

| Case 2 | GWR | 1.4 (1.4, 1.5) |
|---|---|---|
| RMSE | 1.2 (1.0, 1.2) |
| $R^2$ | 0.9 (0.9, 0.9) |
| GWL | 1.2 (1.1, 1.2) |

in the uncorrelated case, while in the correlated case, GWL outperformed GWR in terms of both RMSPE and RMSE, with a greater improvement for prediction of the outcome (RMSPE) than for estimation of the outcome (RMSE).

Pairwise plots of the average regression coefficients are shown in Figure 3. Correlation in the parameter estimates is evident for both GWR and GWL in the cases of both independent and correlated chemicals. In the uncorrelated case, the relationship is most pronounced between the intercept and $\beta_1$ parameters (denoted by $b_0$ and $b_1$, respectively). In the correlated case, there is a noticeable pattern among all of the parameter estimates, with a strong linear relationship evident between the estimates for $\beta_2$ and $\beta_3$ (denoted by $b_2$ and $b_3$, respectively). While GWL breaks up some of the strong correlation among the parameter estimates that is evident in GWR, strong relationships are still present between many of the regression coefficients.

As demonstrated in the box plots of the averaged regression coefficients from the models for the 100 simulated data sets (Fig. 4), GWR appears to accurately capture the importance

![Case 1: Independence (GWR, left panel; GWL, right panel)](image1)

![Case 2: Correlation (GWR, left panel; GWL, right panel)](image2)

Figure 3. Pairwise plots of average regression coefficients across the 100 simulated data sets for the cases of independent chemicals (Case 1) and correlated chemicals (Case 2) for GWR and GWL.
of $x_1$ in Region 1, with distributions centered around 2.0 for the $\beta_1$ estimates in both the independent and the correlated cases. However, we also see that GWR overstates the importance of $x_1$ in Region 2, with distributions centered above zero for the $\beta_1$ estimates regardless of the relationship among the predictor variables. Furthermore, GWL performs shrinkage as expected, demonstrated by the frequent reduction in the magnitude of the parameter estimates when comparing GWR to GWL. However, in both the independent and the correlated cases, GWL often understates the importance of $x_1$ in Region 1 and overstates its importance Region 2, with distributions for the $\beta_1$ estimates centered below 2.0 in Region 1 and around 1.0 in Region 2.

The GWR and GWL regression coefficient estimates from the 100 simulated data sets were averaged at each location and are plotted in Figure 5. The coefficient maps reveal a high degree of correlation between the GWR estimates of $\beta_1$ and $\beta_3$ in both the independent and the correlated cases. This strong negative relationship is also evident in the pairwise scatter plots of the regression coefficients (Fig. 3). Similarly, correlation in the intercept and $\beta_1$ is also apparent in the GWL models, although the correlation between the estimates is not as strong and is largely positive. When examining the coefficient maps for $\beta_1$ in both the independent and the correlated cases, GWR and GWL correctly identified Region 1 as the area of highest activity for $x_1$, but tended to oversmooth the effect into the upper part of Region 2 (i.e., the second row of the grid space). GWL also tended to overshrink the parameter estimates for $\beta_1$ in Region 1, thereby understimating the effect of $x_1$ in the region of activity.

When considering the estimated $\beta_1$ coefficients, GWR appears to identify several clusters of positive and negative associations in the independent case, while in the case of correlation, it finds a positive association in Region 1 and a negative association in Region 2, probably a reflection of the high degree of correlation between the predictors $x_1$ and $x_2$. Although the estimates are small in magnitude, it is clear that GWR is identifying artificial patterns in the $\beta_2$ regression coefficients. In contrast, the maps of the GWL estimates for $\beta_1$ demonstrate little-to-no systematic patterning in both the correlated and uncorrelated cases, suggesting that GWL is able to break up some of the artificial patterning seen in the GWR estimates of $\beta_1$.

Finally, with respect to $\beta_3$, GWR appears to incorrectly identify several clusters of a stronger positive relationship between $x_1$ and the response variable in the uncorrelated case. Furthermore, in the correlated case, GWR incorrectly identifies a strong spatial pattern in the $\beta_3$ estimates, with Region 1 appearing to be an area of high activity. The similarity in the GWR coefficient maps of $\beta_2$ and $\beta_3$ (Fig. 5) reflects the strong linear positive relationship demonstrated in the pairwise plots of the estimated GWR regression coefficients in the correlated case (Fig. 3). The artificial spatial pattern in the GWR estimates of $\beta_3$ parallels the true spatial variation in $\beta_3$ and is probably induced by the correlation between $x_1$ and $x_3$. In contrast, GWL is able to reduce the correlation between the $\beta_2$ and $\beta_3$ estimates and appears to correctly identify the uniform moderate relationship between $x_3$ and the response, regardless of the relationship among the predictor variables.

The percentages of positive and negative GWR coefficient estimates are summarized by region for each correlation case in the left side of Table 3. We see that across the simulated data sets, the GWR estimates of $\beta_1$ were positive nearly 100% of the time in Regions 1 and 2 for both the independent and the correlated cases. This is further evidence that GWR overstates the importance of $\beta_1$ in Region 2. When considering $\beta_2$, 53% of the GWR estimates in Regions 1 and 2 were negative in
the case of independence for at least half of the simulated data sets, while 28% and 77% of the GWR estimates in Regions 1 and 2, respectively, were negative in the case of correlation. Given that $x_2$ has no relationship with the outcome in the simulated data, we suspect the presence of the reversal paradox, which could lead to incorrect inference about the impact of this predictor.

Similarly, as shown in the right side of Table 3, the GWL estimates of $\beta_1$ were positive nearly 100% of the time in Regions 1 and 2 for both the independent and the correlated cases. This indicates that GWL failed to appropriately perform variable selection for $x_1$ in Region 2, the region of inactivity. Furthermore, in at least half of the simulated examples, 17% and 35% of the GWL estimates of $\beta_1$ in Regions 1 and 2, respectively, were zero in the case of independence, and 15% and 35% of the estimates of $\beta_1$ in Regions 1 and 2, respectively, were zero in the case of correlation. Given that $x_1$ is moderately positively associated with the outcome across the study area, these results could lead to the incorrect conclusion that this predictor is not positively associated with the adverse outcome.

The results of applying one and two SEs to the GWR estimated coefficients to classify them as positive, negative, or zero are listed in Table 4. Using the one-SE criteria, GWR incorrectly classified 83% of $\beta_1$ estimates in Region 2 as positive at least half of the time for the independent case and incorrectly classified 84% of $\beta_1$ estimates in Region 2 as positive at least half of the time when the predictors were correlated. Similarly, when applying the two-SE criteria, GWR incorrectly classified 64% of $\beta_1$ estimates in Region 2 as positive at least half of the time for the independent case and incorrectly classified 66% of $\beta_1$ estimates in Region 2 as positive at least half of the time in the correlated case. This implies that GWR frequently yields nonnegligible positive estimates of $\beta_1$ in the region of inactivity.

Furthermore, when applying the one-SE rule, we see that in the case of independence, GWR correctly classified 64% and 72% of the $\beta_2$ estimates as zero in the upper and
Table 3. Median (interquartile range) percentage of GWR and GWL coefficient estimates that were positive, negative, and zero across the 100 simulated data sets for the cases of independent chemicals (Case 1) and correlated chemicals (Case 2).

| Region | Case 1 | Positive | Negative | Zero |
|--------|--------|----------|----------|------|
|        | GWR    | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ | GWR    | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ |
|        |        | 12 (6, 21) | 100 (100, 100) | 47 (39, 64) | 100 (100, 100) | 62 (16, 68) | 100 (100, 100) | 27 (0, 36) | 83 (72, 86) |
|        |        | 88 (79, 94) | 0 (0, 0) | 53 (36, 61) | 0 (0, 0) | 0 (0, 35) | 0 (0, 0) | 27 (4, 38) | 0 (0, 1) |
|        |        | 8 (0, 0) | 0 (0, 0) | 0 (0, 0) | 0 (0, 0) | 38 (33, 49) | 0 (0, 0) | 55 (44, 63) | 17 (14, 26) |

| Region | Case 2 | Positive | Negative | Zero |
|--------|--------|----------|----------|------|
|        | GWR    | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ | GWR    | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ |
|        |        | 76 (66, 86) | 96 (92, 99) | 47 (40, 61) | 100 (100, 100) | 7 (5, 27) | 100 (47, 100) | 15 (0, 23) | 65 (61, 81) |
|        |        | 24 (14, 34) | 4 (1, 8) | 53 (39, 60) | 0 (0, 0) | 19 (18, 21) | 0 (0, 12) | 16 (9, 21) | 0 (0, 0) |
|        |        | 74 (51, 76) | 0 (0, 37) | 81 (52, 85) | 35 (18, 39) | 42 (34, 51) | 0 (0, 0) | 44 (37, 55) | 15 (13, 23) |

Note: On average, the true $\beta_i$ parameter was nonzero (ie, $\beta_i \neq 0$) at 84% of locations in Region 1 and 3% of locations in Region 2 for both Case 1 and Case 2.

Table 4. Median (interquartile range) percentage of GWR coefficient estimates that were positive, negative, and zero across the 100 simulated data sets when considering ±1 and ±2 standard errors of regression coefficient estimates for the cases of independent chemicals (Case 1) and correlated chemicals (Case 2).

| Region | Case 1 | Positive | Negative | Zero |
|--------|--------|----------|----------|------|
|        | GWR (1 SE) | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ | GWR (2 SE) | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ |
|        |        | 12 (1, 10) | 100 (100, 100) | 16 (6, 27) | 100 (99, 100) | 0 (0, 3) | 100 (100, 100) | 1 (0, 6) | 98 (96, 100) |
|        |        | 73 (64, 82) | 0 (0, 0) | 15 (8, 28) | 0 (0, 0) | 53 (44, 63) | 0 (0, 0) | 2 (0, 6) | 0 (0, 0) |
|        |        | 21 (16, 28) | 0 (0, 0) | 64 (54, 73) | 0 (0, 1) | 45 (36, 54) | 0 (0, 0) | 94 (88, 98) | 2 (0, 4) |

| Region | Case 2 | Positive | Negative | Zero |
|--------|--------|----------|----------|------|
|        | GWR (1 SE) | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ | GWR (2 SE) | $\beta_0$ | $\beta_1$ | $\beta_2$ | $\beta_3$ |
|        |        | 35 (22, 44) | 83 (73, 92) | 12 (7, 18) | 100 (100, 100) | 8 (4, 14) | 64 (54, 78) | 1 (0, 3) | 99 (98, 100) |
|        |        | 3 (1, 6) | 0 (0, 1) | 14 (7, 22) | 0 (0, 0) | 0 (0, 0) | 0 (0, 0) | 1 (0, 3) | 0 (0, 0) |
|        |        | 60 (53, 70) | 17 (8, 24) | 72 (65, 77) | 0 (0, 0) | 91 (86, 96) | 36 (23, 46) | 96 (94, 99) | 1 (0, 2) |

Note: On average, the true $\beta_i$ parameter was nonzero (ie, $\beta_i \neq 0$) at 84% of locations in Region 1 and 3% of locations in Region 2 for both Case 1 and Case 2. A parameter estimate was counted as zero if its confidence interval based on one or two standard errors contained zero.
lower regions, respectively, at least half of the time. In the case of correlated predictors, only 51% and 44% of the $\beta_2$ estimates were correctly classified at least half of the time in the upper and lower regions, respectively. Finally, when using the two-SE criteria, 29% of the $\beta_2$ estimates in Region 1 were incorrectly classified as positive at least half of the time when the predictors were correlated. Thus, even when allowing "small" estimates to be considered as negligible, GWR results can still lead to erroneous inferences about the nature of a predictor variable that is not associated with the response.

As an illustrative example, we randomly chose one simulated data set for each correlation case and plotted the corresponding estimated regression coefficients from GWR and GWL, using open circles for the negligible estimates (i.e., GWR estimates with confidence intervals containing zero or GWL estimates of zero) (Fig. 6). This example visually
supports the aggregate results given in Tables 3 and 4. GWR accurately identified Region 1 as the region of high activity for \( \beta_x \) but overstated the effect of \( \beta_1 \), in Region 2, in which the predictor is inactive. In addition, in the correlated case, GWR produced a cluster of nonnegligible positive estimates for \( \beta_x \) in Region 1 and nonnegligible negative estimates for \( \beta_1 \) in Region 2. Finally, while GWL was able to correctly perform variable selection for \( \beta_x \) with some frequency (ie, estimate that \( \beta_x \) was zero, as shown by the open circles), we see that GWL was again unable to correctly identify the spatially varying pattern of \( \beta_x \) in the cases of both independence and correlation. More specifically, GWL estimated a nearly uniform effect for \( \beta_x \) across the study area, understating the effect of \( \beta_1 \) in Region 1 and failing to perform appropriate variable selection for \( \beta_x \) in Region 2 (implying that \( \beta_x \) is active in Region 2).

### Discussion and Conclusion

We have evaluated the ability of the geographically weighted regression methods of GWR and GWL to detect signal from noise in the context of modeling the associations of environmental chemicals and an adverse health effect using a simulation study with both independent and correlated chemicals. We found that GWR was able to identify regions of high activity for an important chemical when the predictors were independent and when they were highly correlated, but it demonstrated a tendency to overstate the importance of this chemical in its region of inactivity. Furthermore, GWR suffered from the reversal paradox for less-important chemicals when the chemicals were correlated, as the variable that was not associated with the outcome was largely positive in the upper study region and largely negative in the lower study region. We also found that with GWL, the signal of the most important chemical was diminished, with less distinction between the inactive and active study regions, regardless of the correlation among the chemicals.

Previous work has addressed the issue of collinearity in GWR. Wheeler and Tiefelsdorf\(^{12}\) first demonstrated the link between collinearity in GWR and correlation of estimated regression coefficients using simulation studies. These authors introduced systematic collinearity into the model by adding correlation to a pair of covariates and found consistent evidence of increasing correlation in GWR coefficients with increasing collinearity. Wheeler and Calder\(^{13}\) used two simulation studies to evaluate the coverage probability and accuracy of the regression coefficients from GWR. Results of the simulation studies include low coverage probabilities for the GWR coefficients and consistently increasing error in the coefficients when collinearity is increased. Wheeler\(^{12}\) conducted a simple experiment by systematically increasing collinearity in a data set to demonstrate that a penalized form of GWR, geographically weighted ridge regression, reduces the extreme effects of collinearity that afflict GWR. More recent simulation study work confirms that a nonnegligible amount of spatial variation of and correlation between GWR coefficient surfaces is inherently generated by the method.\(^{15}\) This work finds that the false-positive rates for GWR coefficients are typically much higher than convention would mandate, from \(< 10\%\) to \(> 50\%\) of the time (depending on the true correlation level between two covariates) when the true underlying process is stationary.

Wheeler\(^{14}\) expanded the simulation study of Wheeler and Calder\(^ {11}\) to contain four explanatory variables and 196 observations in a study of the performance of GWR and GWL. This work compared the coefficient accuracy and the predictive performance of the models in the presence of collinearity. In these experiments, 100 realizations of a data-generating process were used with the true local coefficients sampled from a multivariate normal distribution. These simulation studies show that the performance of GWR in terms of both prediction and coefficient accuracy can be improved by constraining the magnitude of its regression coefficients with techniques designed to remediate collinearity. However, the experiments reported in that study show that the correlation between local coefficients is reduced but not eliminated with GWL, and that although GWL can shrink some coefficients to zero to stabilize the model, the estimates still tend to be positively correlated with those from GWR.\(^ {15}\)

We have extended these results in the case of three environmental chemicals to identify evidence of the reversal paradox and evaluate the correct identification of local “hot spots” or regions of high activity for one chemical. Our results demonstrate that while GWR can correctly identify a region of high activity for one chemical, it has difficulty in identifying regions of inactivity or low exposure. Additionally, GWR artificially induces spatial patterning and suffers from the reversal paradox in the setting of highly correlated predictor variables. Finally, we have shown that while GWL reduces the correlation among the coefficient estimates and tempers the reversal paradox that is problematic with GWR, it suffers from an inability to adequately distinguish local regions of high activity regardless of the relationship among the predictor variables. The implications of our findings for environmental risk analysis is that GWR may incorrectly identify some chemicals as positively or negatively associated with disease risk, and GWL may not correctly estimate the magnitude of association for an important chemical in some regions of the study area. Given these findings, more methodological development is required to better estimate the effects of correlated environmental chemicals on diseases associated with environmental factors, such as many cancers.

### Author Contributions

Conceived and designed the experiments: JC, DCW, CG. Analyzed the data: JC, DCW, CG. Wrote the first draft of the manuscript: CG, JC, DCW. Contributed to the writing of the manuscript: JC, DCW, CG. Agree with manuscript results and conclusions: JC, DCW, CG. Jointly developed the structure and arguments for the paper: JC, DCW, CG. Made
critical revisions and approved final version: JC, DCW, CG.
All authors reviewed and approved of the final manuscript.

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