High Accuracy Geochemical Map Generation Method by a Spatial Autocorrelation-Based Mixture Interpolation Using Remote Sensing Data

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Abstract: Generating a high-resolution whole-pixel geochemical contents map from a map with sparse distribution is a regression problem. Currently, multivariate prediction models like machine learning (ML) are constructed to raise the geoscience mapping resolution. Methods coupling the spatial autocorrelation into the ML model have been proposed for raising ML prediction accuracy. Previously proposed methods are needed for complicated modification in ML models. In this research, we propose a new algorithm called spatial autocorrelation-based mixture interpolation (SABAMIN), with which it is easier to merge spatial autocorrelation into a ML model only using a data augmentation strategy. To test the feasibility of this concept, remote sensing data including those from the advanced spaceborne thermal emission and reflection radiometer (ASTER), digital elevation model (DEM), and geophysics (geomagnetic) data were used for the feasibility study, along with copper geochemical and copper mine data from Arizona, USA. We explained why spatial information can be coupled into an ML model only by data augmentation, and introduced how to operate data augmentation in our case. Four tests—(i) cross-validation of measured data, (ii) the blind test, (iii) the temporal stability test, and (iv) the predictor importance test—were conducted to evaluate the model. As the results, the model’s accuracy was improved compared with a traditional ML model, and the reliability of the algorithm was confirmed. In summary, combining the univariate interpolation method with multivariate prediction with data augmentation proved effective for geological studies.

Keywords: geochemical mapping; remote sensing; machine learning; data augmentation; computational geometry

1. Introduction

1.1. Background

Geoscience data, which are dependent on the fieldwork of geologists, for example, lithological data, geochemical data, etc., are always sparse on distribution maps. In recent studies, besides conventional univariate geospatial interpolation methods, multivariate prediction models have been constructed to raise the geoscience mapping resolution by using various geoscience data, especially remote sensing data, as predictors. Pal et al. [1] have used fused multi-classifiers, which include multi-spectral data, to achieve high-resolution lithological classification. Kirkwood et al. [2] have applied random forests (RFs) machine learning (ML) methods to generate whole-pixel geochemical contents maps. The data in unknown locations are predicted by the geographical referenced input data containing co-located pixels specified by coordinates linked to a spatial reference frame, which is equivalent to processing the predictions in a geographic space where samples are only compared numerically [3].
Unlike lithological mapping, which is a classification work, a geochemical contents map is like an image with analogue values, thus generating a high-resolution whole-pixel geochemical contents map is a regression problem. It is suggested that considering the geological spatial dependencies, spatial location, and spatial autocorrelation due to geological continuity, it cannot be ignored [4]. Cracknell et al. [5] demonstrated the prediction accuracy raised by treating spatial coordinates as predictors to couple the spatial autocorrelation into the ML model. Sergeev et al. [6] proposed a method to include spatial autocorrelation into an artificial neural network (ANN) by applying a kriging model. However, in their models, ML models are needed for complicated modification.

Therefore, we propose a new algorithm called SABAMIN (spatial autocorrelation-based mixture interpolation) that can merge both spatial location and autocorrelation, which are generated from the univariate geospatial interpolation model into an ML model using a data augmentation strategy [7] to provide a high accuracy model that can generate a high-resolution geochemical map. The data augmentation strategy is currently used for solving small data machine learning, and has proved effective in raising the accuracy of the machine learning model [8,9]. It is only needed to contain pseudo training data generated from a reliable expert model into training datasets, which is an easy task. It has to be noted that the accuracy and reliability of the ML model is determined by the accuracy and reliability of the model for generating the pseudo training data.

In this research, to prove the effectiveness of this concept in geological study, we combine kriging interpolation [10] and RFs to construct our new algorithm as an example. Because kriging interpolation is a well-known method based on the spatial autocorrelation of data in Euclidian space, and RF is an interpretable ML method [11] that has the merit to determine which predictor is important in the model, here, we use kriging interpolation to create pseudo training datasets and RF to construct a prediction model. We explain why both spatial location and autocorrelation can be coupled into an ML model only by data augmentation, as well as how to create reliable pseudo training data in the case of using kriging interpolation. Moreover, we demonstrate the reliability of the algorithm using the example of generating a whole-pixel copper contents distribution map.

A portion of the findings in this report is based on the work [12] presented at the SPIE Remote Sensing 2019 conference. All the data used in this research are open data obtained from the United States Geological Survey (https://lpdaac.usgs.gov).

1.2. Why Spatial Information Can be Coupled into an ML Model Only by Data Augmentation

In this section, we will explain why both spatial location and autocorrelation can be coupled into an ML model only by data augmentation.

1.2.1. Spatial Information Calculated from Kriging Interpolation

The sampled target variable data points are defined as $S_i(x_{si}, y_{si}), si = 1, \ldots , N$, and vacant points are defined as $A_j(x_{aj}, y_{aj}), aj = 1, \ldots , M$. Here, $N$ and $M$ are the number of total sampled target variable points and vacant pixels, respectively.

In kriging interpolation, two steps are executed in the following order: (i) Construct a spatial distribution model of sampled points using variography; (ii) interpolate the vacant points. In the basic kriging model, the system of equations is shown in Equation (1):

$$\min_{\Lambda} E = \Lambda^T \cdot V \cdot \Lambda - C^T \cdot \Lambda - \Lambda^T \cdot C + V \quad \text{subject to} \quad 1^T \cdot \Lambda = 1 \quad (1)$$

where $E$ is the variance of prediction errors at vacant points, $\Lambda$ is the size of an $(M \times N)$ matrix whose elements $\lambda_{ij}$ are the weights to the measured value of $S_i$ for interpolating $A_j$, $V$ is the size of an $(N \times N)$ matrix whose elements $v_{a,b}$ is the variance (here, semivariance is used) within pairs of sampled points $S_a$ and $S_b$, and $C$ is the size of an $(N \times M)$ matrix whose elements $c_{a,b}$ are the covariance within pairs of sampled points $S_a$ and vacant pixel $A_b$. To ensure that the model is unbiased, the weights must be summed to one. To obtain the best prediction precision, $E$ should be minimized, which is equivalent to
optimization problem results in a kriging system. By using Lagrange multipliers, $\Lambda$ can be solved by Equations (2) and (3):

$$
\begin{bmatrix}
\Lambda \\
\mu
\end{bmatrix} = 
\begin{bmatrix}
V & 1 \\
1^T & 0
\end{bmatrix}^{-1}
\begin{bmatrix}
C \\
1
\end{bmatrix}
$$

(2)

$$
\Lambda = V^{-1}\cdot C + V^{-1}\left(-1^T\cdot V^{-1}\right)^{-1}\cdot 1^T\cdot V^{-1}\cdot C - V^{-1}\left(-1^T\cdot V^{-1}\right)^{-1}
$$

(3)

where $\mu$ is a Lagrange multiplier used in the minimization of the kriging error to honor the unbiasedness condition. Equation (3) is based on the analytic inversion formula for block-matrix inversion, and when the block matrix is not square, pseudo inversion is used. Then, the vacant pixels are interpolated through Equation (4), which is analogous to inverse distance weighting (IDW) interpolation [13]:

$$
P_A = \Lambda \cdot P_s
$$

(4)

where $P_s$ is the size of an $(N \times 1)$ vector whose elements $P_{si}$ are the measured target variable value at position $S_i$, and $P_A$ is the size of an $(M \times 1)$ vector whose elements $P_{Aj}$ are the interpolated target variable value at position $A_j$.

In the first step of kriging interpolation, a semivariogram of measured points that express the spatial autocorrelation of these points is formed to estimate the limitation of the autocorrelation range. For this estimation, the semivariance distribution in the semivariogram is regressed by a monotonically increasing function (a spherical model is used in this study) [14]. This regressed function can be expressed by Equation (5):

$$
V(x, y) = \beta(H(x, y), r, k)
$$

(5)

where $H$ is the size of an $(N \times N)$ matrix whose elements $h_{ab}$ are the Euclidean distances between pairs of sampled points $S_a$ and $S_b$, which are variables determined by coordinates $x$ and $y$. In accordance with geological continuity, near points are more similar; thus, the further the points are, the bigger the semivariance. Therefore, they should be saturated outside a specific distance, which means when the vacant pixels are further than this distance, the estimation is too bad to trust. Further, $r$ is defined as a specific distance that corresponds to the “range” of the regressed model of the semivariogram, and $k$ is defined as the semivariance at distance $r$, which corresponds to the “sill” of the regressed model of the semivariogram [15]. $\beta$ is the regressed function, and $C$ can be also constructed by this regressed function, which is expressed by Equation (6):

$$
C(x, y) = \beta(B(x, y), r, k)
$$

(6)

where $B$ is the size of an $(N \times M)$ matrix whose elements $b_{ab}$ are the Euclidean distances between pairs of sampled points $S_a$ and vacant pixel $A_b$, which are also variables determined by coordinates $x$ and $y$. It is noted that $h_{ab}$ and $b_{ab}$ cannot exceed $r$, thus after comparing $h_{ab}$ and $r$ in Equation (5) and comparing $b_{ab}$ and $r$ in Equation (6), the smaller one is used for calculation. As a result, $P_{Aj}$ can be expressed as a function, shown in Equation (7):

$$
P_{Aj} = f\left(A\left(x_{aj}, y_{aj}\right), x_{si}, y_{si}, P_s\right)
$$

(7)

1.2.2. Merge Spatial Information into RF Model

In a supervised ML process, a function or rule is created on the basis of example input–output pairs, for example, an RF. In the RF regression algorithm, decision trees are produced during the process. Given predictor vectors $t$, which will be listed in Table 2 as $t_d \in R^{53}$, $d = 1, \ldots, 63$, and given a target vector $\Pi$, the copper contents, $\Pi_u \in R^L$, $u = 1, \ldots, L$, where $L$ is the total number of training datasets, a decision tree recursively partitions the space $R$. Considering the kriging-interpolated target, variable points are included in the training datasets, $P_{Si} + P_{Aj} \in \Pi_u$, and $L = M + N$. 

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There is data $U$ at node $\gamma$, and for each candidate split $\theta = (t_d, g_\gamma)$ consisting of a feature $t_d$ and threshold $g_\gamma$, the data are partitioned into subsets $U_1(\theta)$ and $U_2(\theta)$. Sets $U_1(\theta)$ and $U_2(\theta)$, and their relationship with $U$ are expressed by Equations (8) and (9):

$$U_1(\theta) = (t, \Pi)|t_d \leq g_m$$

(8)

$$U_2(\theta) = U - U_1(\theta)$$

(9)

The effect of the split is evaluated by its impurity, which is composed of mean squared error (MSE) processing and a search for the minimum impurity to determine locations for future splits. The impurity $I(U, \theta)$ is computed using an impurity function $I$, shown in Equation (10):

$$I(U, \theta) = \frac{l_1}{L_1}I(U_1(\theta)) + \frac{l_2}{L_2}I(U_2(\theta))$$

$$= \frac{l_1}{L_1} \sum_{u_1 \in U_1} \left( \frac{L_1 - 1}{L_1} \sum_{u_j \in U_1} \Pi_{u_1} - \frac{1}{L_1} \sum_{u_j \in U_1} \Pi_{u_1} \right)^2 + \frac{l_2}{L_2} \sum_{u_2 \in U_2} \left( \frac{L_2 - 1}{L_2} \sum_{u_j \in U_2} \Pi_{u_2} - \frac{1}{L_2} \sum_{u_j \in U_2} \Pi_{u_2} \right)^2$$

(10)

where $L_1$ is the number of datasets in $U_1(\theta)$ and $L_2$ is the number of datasets in $U_2(\theta)$, respectively. Then, the optimized split $\theta_o$ is specified by minimizing $I(U, \theta)$, whose process can be expressed by Equation (11):

$$\theta_o = \text{argmin}_\theta I(U, \theta)$$

(11)

Finally, the RF model is constructed by repeating these processes. From Equation (10), we can also calculate the following:

$$\Pi_{u_1} = \frac{1}{L_1} \sum_{u_1 \in U_1} \Pi_{u_1}, \Pi_{u_2} = \frac{1}{L_2} \sum_{u_2 \in U_2} \Pi_{u_2}$$

(12)

where $\Pi_{u_1}$ and $\Pi_{u_2}$ are the average values of the target variables in $U_1(\theta)$ and $U_2(\theta)$, respectively. Then Equation (12) becomes:

$$\Pi_{u_1} \left( \lambda(A_{u_1}, y_{u_1}) \right) = \frac{1}{L_1} \sum_{u_1 \in U_1} \Pi_{A_{u_1} + \sum_{u_i \in U_{u1}} \Pi_{A_{u_1}}}$$

(13)

$$= \frac{1}{L_1} \sum_{u_1 \in U_1} \Pi_{A_{u_1}} + \sum_{u_1 \in U_{u1}} f(\lambda(A_{u_1}, y_{u_1}, x_{u_1}, y_{u_1}, P_{u_1}))$$

$$\Pi_{u_2} \left( \lambda(A_{u_2}, y_{u_2}) \right) = \frac{1}{L_2} \sum_{u_2 \in U_2} \Pi_{A_{u_2} + \sum_{u_i \in U_{u2}} \Pi_{A_{u_2}}}$$

(14)

$$= \frac{1}{L_2} \sum_{u_2 \in U_2} \Pi_{A_{u_2}} + \sum_{u_2 \in U_{u2}} f(\lambda(A_{u_2}, y_{u_2}, x_{u_2}, y_{u_2}, P_{u_2}))$$

where $L_{sq}$ is the number of sample points in $U_q(\theta)$, and $L_{aq}$ is the number of kriging interpolated points in $U_q(\theta)$ ($q = 1, 2$). As a result, $\Pi_{u_1}$ and $\Pi_{u_2}$ are determined by $\lambda$, and $\theta_o$, which determines the structure of the RF model and predicted value by this model, also becomes an $\lambda$-dependent variable.

Therefore, as long as we generate some pseudo training data by applying kriging interpolation and including them in training datasets, the spatial location and spatial autocorrelations can be combined into the ML model by following the processes above.
2. Materials and Methods

2.1. Study Area and Target Variable

Arizona, USA is a well-known copper-mining area. In this area, abundant remote sensing and geochemical databases have been created from previous geological studies [16–19]. Moreover, Arizona has satellite image data with less noise-like cloud. Therefore, it was considered as a good model case for a remote-sensing big data study, and copper was selected as our target element for testing the feasibility of our new algorithm for geochemical map generation.

The geochemical data including copper elements were obtained from the National Geochemical Database-Reformatted Data from the National Uranium Resource Evaluation (NURE) Hydrogeochemical and Stream Sediment Reconnaissance (HSSR) Program [20]. In this study, to constitute a big dataset for ML, mainly concerning the amount of sampled copper geochemical data, the data were obtained from Arizona and its outskirts as our region of interest (ROI), which included the boundary areas of California, Nevada, Utah, and New Mexico. The whole ROI was in a rectangle range from (W115.9040, N37.6758) to (W107.888, N30.9958) (not including the territory of Mexico). The copper contents data were obtained from the “cu_ppm” column of the table in the shape file ‘nuresed.shp’ (source: https://mrdata.usgs.gov/nure/sediment/nuresed.zip). Inside the ROI, a total of approximately 16,000 samples were obtained. In the negative data in the raw data table, for example, −5 ppm means less than 5 ppm [21]. To process the negative data properly, in this study, we assumed the real values of the negative values, for example, −5 ppm should be a random value between 0 and 5 ppm in accordance with the Gaussian distribution.

Figure 1a shows our ROI, and Figure 1b shows the boundaries of Arizona, the reference geological map, the study areas Ta, Tb, Tc, Pa, Pb, and Pc, and spatially-scattered sampled copper data points in this area. These study areas are all the detailed local areas inside the ROI. For the following validation test, Ta, Tb, Tc, Pa, Pb, and Pc were all selected to be near mining districts where mines are dense, which assured both geochemical anomalies and background content areas were included in the same map as the characteristic example model cases.

The geological map is arranged from the Geologic Map of the United States at a scale of 1:2,500,000 [22] (source: https://mrdata.usgs.gov/geology/us/kbgeology.zip). There are 50 geological/lithological (G/L) types included in the whole ROI referenced from “UNIT” in the shape file “kbge.shp”, which are numbered from 1 to 50 and are marked in different colors. A summary of G/L type names present within the ROI referenced from the “ROCK” column of the table in the shape file are listed in Table 1. We set the grid for the whole map (including the target variable mentioned above; the predictor variable is mentioned later) to 0.008 by 0.008 degrees in longitude and latitude, respectively (approximately 1 by 1 km in distance), resulting in an image dimension of 836 × 1003 pixels. By scattering the sampled copper data on the map in this resolution, over 50% of the map area is not covered by the sampled data, which leaves a huge blank area on the map. As a result, in our case, the sampled data compose three independent point groups (PGs) that are observed in the right side (PG1), left median position (PG2), and left upper position of the map (PG3). Figure 1c shows the statistics of sampled copper data amounts and average contents of all sampled copper data in each lithological type area.

The details of Ta, Tb, Tc, Pa, Pb, and Pc are shown in Figures 2 and 3, and Table 2. The sampled copper data are plotted by black circles on the reference geological map, with the size of a black circle proportional to the copper content. The coordinates of copper mines in Arizona referenced from the Mineral Resources Data System of the United States Geological Survey (USGS) [23,24] are also plotted by red diamonds (source: https://mrdata.usgs.gov/mrds/output/mrds-fUS04.zip). Only those copper mines whose “dev_stat” in the shape files are labeled “Producer” or “Past Producer” are marked. A number of important landmarks such as cities and mountains are also marked on the map to show as references of the approximate positions. A detailed introduction of the study areas is summarized in the supplementary material (Table S1).
Figure 1. Cont.
Figure 1. The region of interest (ROI) and the position study area in this research: (a) The approximate position in the USA; (b) the detailed study area, the boundaries of Arizona, the reference geological map of the study area, and spatially-scattered geochemical data points in this area. Different geological/lithological (G/L) types are numbered and marked in different colors. Each correspondent lithological type is referenced in Table A1 of Appendix A. Sampled copper points concentrated in different positions on the map are marked as PG1, PG2, and PG3. The study areas are marked as Ta, Tb, Tc, Pa, Pb, and Pc. (c) The statistics of sampled copper data amounts and average content of all sampled copper data in each lithological type area.

Figure 2. Cont.
Figure 2. Study area details of: (a) Ta; (b) Tb; (c) Tc. The sampled copper data are plotted by black circles on the reference geological map, and the circle radius is proportional to the copper content. The G/L types in this figure are marked the same number of those in Figure 1.

Figure 3. Cont.
Figure 3. Study areas of: (a) Pa; (b) Pb; (c) Pc. The sampled copper data are plotted by black circles, and the copper mine positions referenced in the United States Geological Survey (USGS) are plotted by red diamonds. The G/L types in this figure are marked the same number of those in Figure 1.

Table 1. Summary of geological/lithological (G/L) types present within the study areas referenced from “ROCK” column of the table in the shape file “kbge.shp”. The G/L type numbers correspond to the numbered colors in Figures 1–3.

| No. | G/L Type                      | No. | G/L Type                      | No. | G/L Type                      | No. | G/L Type                      |
|-----|--------------------------------|-----|--------------------------------|-----|--------------------------------|-----|--------------------------------|
| 1   | Water                          | 14  | Eocene                         | 27  | Upper Mesozoic eugeosynclinal  | 40  | Upper Paleozoic eugeosynclinal |
| 2   | Quaternary continental         | 15  | Paleocene continental          | 28  | Jurassic                        | 41  | Upper Paleozoic elastic wedge facies |
| 3   | Quaternary volcanic rocks      | 16  | Navarro Group                  | 29  | Lower Mesozoic volcanic rocks  | 42  | Lower Paleozoic                |
| 4   | Pliocene continental           | 17  | Taylor Group                   | 30  | Jurassic granitic rocks        | 43  | Cambrian                       |
| 5   | Pliocene volcanic rocks        | 18  | Latest Cretaceous granitic     | 31  | Lower Jurassic and upper Triassic | 44  | Z sedimentary rocks            |
| 6   | Pliocene felsic volcanic rocks | 19  | Austin and Eagle Ford Groups   | 32  | Lower Mesozoic                 | 45  | Y sedimentary rocks            |
Table 1. Cont.

| No. | G/L Type                  | No. | G/L Type               | No. | G/L Type                      |
|-----|--------------------------|-----|------------------------|-----|------------------------------|
| 7   | Miocene continental      | 20  | Upper Cretaceous       | 33  | Lower Mesozoic eugeosynclinal|
|     | Tertiary intrusive rocks |     |                        |     | Younger Y granitic rocks     |
| 8   | Tertiary intrusive rocks | 21  | Cretaceous continental | 34  | Triassic                     |
| 9   | Oligocene continental    | 22  | Cretaceous eugeosynclinal| 35  | Permian                      |
| 10  | Miocene volcanic rocks   | 23  | Uppermost Cretaceous   | 36  | X metasedimentary rocks      |
| 11  | Miocene felsic volcanic  | 24  | Cretaceous granite     | 37  | Lower part of Leonardian Series|
|     | rocks                    |     |                        |     | Lower part of Leonardian Series|
| 12  | Eocene continental       | 25  | Woodbine and Tuscalosa| 38  | Wolfcampian Series continental|
| 13  | Lower Tertiary           | 26  | Lower Cretaceous       | 39  | Upper Paleozoic              |

Table 2. Detailed information of $T_a$, $T_b$, $T_c$, $P_a$, $P_b$, and $P_c$.

| Area | Range | Resolution |
|------|-------|------------|
| $T_a$| (W111.7200, N33.5638)–(W110.9440, N32.7958) | 110 x 110 |
| $T_b$| (W111.7840, N32.5558)–(W110.9120, N31.6838) | 110 x 110 |
| $T_c$| (W110.7840, N32.7158)–(W109.9920, N31.9238) | 110 x 110 |
| $P_a$| (W114.3840, N35.4758)–(W113.9840, N35.0758) | 50 x 50   |
| $P_b$| (W109.6240, N33.2758)–(W109.2240, N32.8758) | 50 x 50   |
| $P_c$| (W112.2640, N34.5558)–(W111.8640, N34.1558) | 50 x 50   |

2.2. Predictor Variables—High-Resolution Remote Sensing Data

All of the available remote sensing data including that from the advanced spaceborne thermal emission and reflection radiometer (ASTER) data, digital elevation model (DEM), and geophysics (geomagnetic) data [25] are used to create an ML model to make predictions. Matlab (Mathworks, Natick, USA) is used for all of the data processing and calculations here and in the sections below.

All the ASTER data are extracted from “ASTER Level 1T” and DEM data are extracted from “ASTER Global Digital Elevation Model” in the USGS’s EarthExplorer search engine, respectively (source: https://lpdaac.usgs.gov/data_access/data_pool, courtesy of the NASA Land Processes Distributed Active Archive Center (LP DAAC), USGS/Earth Resources Observation and Science (EROS) Center, Sioux Falls, South Dakota). The ASTER data are part of the ASTER Level 1T group, and the search filter is set as follows: Cloud cover is “less than 10%”; correction achieved is “all”; and SWIR, TIR, and VNIR1 modes are all “on”. Geomagnetic data are obtained from magnetic anomaly maps and data for North America [25] (source: https://mrdata.usgs.gov/magnetic/USmag_origmrg.zip). The geomagnetic data are international geomagnetic reference field (IGRF) residual signals taken from a height of 305 m, the precision of which is approximately 1–10 nT.

One shot from an ASTER satellite can only cover $60 \times 60$ km, and a DEM can only cover an area approximately $100 \times 100$ km on Earth; therefore, to cover the whole ROI in this research, we retrieved a total of 637 ASTER images and 160 DEM images from the data pool and created a composite of all the images in accordance with their longitude and latitude to create a wide-range map. The detail of the source of ASTER and DEM data are listed in Tables S2–S4 of the supplementary document. All the ASTER band data are extracted from the “.hdf” file by the “hdftool” of Matlab, and DEM data are obtained by translating the “.tiff” DEM image to a digital number.

The resolutions of these predictors are all different (15 of ASTER visible and near-infrared (VNIR) band data, 30 of ASTER short wave infrared (SWIR) band data, 90 of ASTER thermal band data, 30 of DEM data, and 5 km of geomagnetic data). The VNIR and SWIR data and DEM data are 8-bit, and the thermal data are 12-bit. The geomagnetic data are processed into four types: Analytic signal-processed geomagnetic data, reduction to pole-processed geomagnetic data, residual IGRF- and vertical first
derivative-processed geomagnetic data. ASTER and DEM predictor variables and their derivatives are reprocessed from their original data grid to a regular 1-km grid to be studied using bicubic interpolation and geomagnetic predictor variables are reprocessed using the nearest interpolation, i.e., $5 \times 5$ pixels in a 1-km grid will be the same value.

The derivatives of ASTER as predictors include all the commonly used ASTER band ratios and their combinations that are reported as a significant lithological index for geological mapping [26,27]. For example, the carbonate index (No. 41 predictor in Table 3) map is acquired by calculating the image of Band 13, 14, 15 pixel-by-pixel. A total of 63 predictor variables used in this research are listed in Table 3. The predictors are categorized into five groups: Geomagnetic, DEM, ASTER band, ASTER lithological index, and Coordinates.

**Table 3. Predictor variables used in this research.**

| Group         | No. | Predictor | Predictor Description                                      |
|---------------|-----|-----------|------------------------------------------------------------|
| Geo-magnetic  | 1   | Mag_AS    | Analytic signal processed geomagnetic data                |
|               | 2   | Mag_RTP   | Reduction to pole processed geomagnetic data               |
|               | 3   | Mag_TMI   | The residual of international geomagnetic reference field (IGRF) |
|               | 4   | Mag_VD    | Vertical first derivative processed geomagnetic data       |
| DEM           | 5   | Altitude  | The altitude of the Earth surface, from 0–2500 m         |
|               | 6   | Slope     | The elevation of the Earth surface                        |
| ASTER Band    | 7   | Band_1    | The ASTER band 1 sensor data                               |
|               | 8   | Band_2    | The ASTER band 2 sensor data                               |
|               | 9   | Band_3N   | The ASTER band 3N sensor data                              |
|               | 10  | Band_4    | The ASTER band 4 sensor data                               |
|               | 11  | Band_5    | The ASTER band 5 sensor data                               |
|               | 12  | Band_6    | The ASTER band 6 sensor data                               |
|               | 13  | Band_7    | The ASTER band 7 sensor data                               |
|               | 14  | Band_8    | The ASTER band 8 sensor data                               |
|               | 15  | Band_9    | The ASTER band 9 sensor data                               |
|               | 16  | Band_10   | The ASTER band 10 sensor data                              |
|               | 17  | Band_11   | The ASTER band 11 sensor data                              |
|               | 18  | Band_12   | The ASTER band 12 sensor data                              |
|               | 19  | Band_13   | The ASTER band 13 sensor data                              |
|               | 20  | Band_14   | The ASTER band 14 sensor data                              |
|               | 21  | Band_1_R  | The reverse of 7                                           |
|               | 22  | Band_2_R  | The reverse of 8                                           |
|               | 23  | Band_3_R  | The reverse of 9                                           |
|               | 24  | Band_4_R  | The reverse of 10                                          |
|               | 25  | Band_5_R  | The reverse of 11                                          |
|               | 26  | Band_6_R  | The reverse of 12                                          |
|               | 27  | Band_7_R  | The reverse of 13                                          |
|               | 28  | Band_8_R  | The reverse of 14                                          |
|               | 29  | Band_9_R  | The reverse of 15                                          |
|               | 30  | Band_10_R | The reverse of 16                                          |
|               | 31  | Band_11_R | The reverse of 17                                          |
|               | 32  | Band_12_R | The reverse of 18                                          |
|               | 33  | Band_13_R | The reverse of 19                                          |
|               | 34  | Band_14_R | The reverse of 20                                          |
Table 3. Cont.

| Group No. | Predictor         | Predictor Description                                                                 |
|-----------|-------------------|---------------------------------------------------------------------------------------|
| 35        | R_Ferric_iron     | Feature index of Fe<sup>3+</sup>, =8/7                                                |
| 36        | R_Ferrous_iron    | Feature index of Fe<sup>2+</sup>, =11/9 + 7/8                                        |
| 37        | R_Laterite        | Feature index of laterite, =10/11                                                    |
| 38        | R_Gosan           | Feature index of gosan, =10/8                                                        |
| 39        | R_Ferrous_Silica  | Feature index of ferrous silicates, mainly Fe oxide Cu-Au alteration, =11/10          |
| 40        | R_Ferric_oxides   | Feature index of ferric oxides, =10/9                                                |
| 41        | R_Carbonate      | Feature index of carbonate/chlorite/epidote, =(13 + 15)/14                           |
| 42        | R_Epidote        | Feature index of epidote/chlorite/amphibole, =(12 + 15)/(13 + 14)                    |
| 43        | R_MgOH           | Feature index of Amphibole/MgOH, =(12 + 15)/14                                        |
| 44        | R_Amphibole      | Feature index of amphibole, =12/14                                                    |
| 45        | R_Carbonate2     | Feature index of carbonate, =19/20                                                   |
| 46        | R_Dolomite       | Feature index of dolomite, =(12 + 14)/13                                              |
| 47        | R_Sericite       | Feature index of sericite/muscovite/ilinite/smectite, =(11 + 13)/12                   |
| 48        | R_Alunite        | Feature index of alunite/kaolinite/pyrophyllite, =(10 + 1 2)/11                        |
| 49        | R_phengitic      | Feature index of phengitic, =11/12                                                    |
| 50        | R_Muscovite      | Feature index of muscovite, =13/12                                                    |
| 51        | R_Kaolinite      | Feature index of kaolinite, =13/11                                                    |
| 52        | R_Quartz         | Feature index of quartz rich rocks, =20/18                                             |
| 53        | R_Basic_deg      | Feature index of basic degree index of SiO<sub>2</sub>, =18/19                         |
| 54        | R_SiO2           | Feature index of SiO<sub>2</sub>, =19/18                                              |
| 55        | R_Siliceous_rock | Feature index of siliceous rocks, =17/16(16 × 18)                                      |
| 56        | R_Silica1        | Feature index of the first pattern of silica, =17/16                                 |
| 57        | R_Silica2        | Feature index of the second pattern of silica, =17/18                                |
| 58        | R_Silica3        | Feature index of the third pattern of silica, =19/16                                 |
| 59        | R_Vegetation     | Feature index of vegetation, =9/8                                                     |
| 60        | R_Clay           | Feature index of clay, =(11 × 13)/12<sup>2</sup>                                      |
| 61        | R_NDVI           | Feature index of NDVI, =(9 − 8)/(9 + 8)                                               |

2.3. Spatial Autocorrelation-Based Mixture Interpolation Algorithm

2.3.1. The Process of the SABAMIN Algorithm

Figure 4 shows the flow chart of the SABAMIN algorithm. First of all, the measured target variable data are input into the pseudo training data generation model by using kriging interpolation, and then the pseudo training data are generated. Both are applied for composing of training datasets. Meanwhile, both are input into a coordinate extraction function, then the coordinates of training datasets are extracted to construct a coordinate filter for selecting predictor variable data in correspondent pixel from the whole ROI. Furthermore, by this coordinate filter, the ROI is divided into two areas: “Areas for training” (T areas) and the remaining blank areas as “areas waiting to be predicted” (P areas), and the predictor variable data is divided into “data for training” and “data for prediction”, respectively. Through RF learning, a prediction model is generated and by inputting data for prediction into this model, the target variable data in P area are generated. Finally, using a mapping function, the measured data, pseudo training data, and predicted data are merged together to complete a whole-pixel map of the ROI. This is a process of mixing univariate interpolation and multivariate prediction; thus, we named our new algorithm spatial autocorrelation-based mixture interpolation.
2.3.2. How to Generate Pseudo Training Data in Our Case

The generation of pseudo training data is a crucial part of algorithm. It is important that pseudo training data are reliable. However, in our case, there are problems that need to be solved. Below, we introduce how to generate reliable pseudo training data in our case.

Figure 5 shows a conceptual drawing explaining these problems. As mentioned before, kriging interpolation is reliable in a range of \( r \). However, mineral contents have been reported to be highly correlated with lithological features [28–30]; thus, it is considered that the spatial autocorrelations within points of different geological areas are different and the correlation between two different areas is lower. Due to geological discriminations that exist at the area border, different G/L types are considered to have different \( r \). If the interpolation model is produced under the condition that a single \( r \) is set by fusing all sampled points in the study area, distribution crosstalk \( Cr_1 \) (Figure 5a) will occur in the regions near the borders of different geological areas because the neighboring sampled points inside the range of \( r \) may come from different geological areas. Therefore, the pseudo training data in \( Cr_1 \), which do not make sense in geoscience, are too unreliable to be used. To avoid \( Cr_1 \), it is necessary to determine reliable spatial sections that enable reliable pseudo training data points inside them. Therefore, kriging interpolation must be processed in each geological area independently.

Sampled points are categorized into different point groups in accordance with G/L types, which are defined as \( PG_{Geo} \), \( \psi = 1, \ldots, \Gamma \), where \( \Gamma \) is the total number of G/L types in the study area, and they need to be separated from each other spatially. Creating an envelope of the PGs enables them to be separated from each other. To achieve this, a computational geometry strategy is applied here.

Observing the distribution shape of \( PG_{Geo} \) domains, they are not usually geometrically convex, but always contain multiple concave parts. The alpha shape [31], an algorithm capable of generating an optimized envelope in this case, was chosen for an automatic enveloping generation process. An alpha
shape is a family of piecewise linear simple curves in the Euclidean plane associated with the shape of a finite set of points. The PG is split into multiple triangles by Delaunay triangulation [32], wherein each edge or triangle may be associated with a characteristic radius; the radius $\Phi$ of the smallest empty circle containing an edge or triangle. After the alpha shape has been formed, a closed spatial section is partitioned by the edge of the shape, and the points inside the shape that can be judged by a point-in-polygon algorithm [33] can be considered reliable to enable all vacant pixels to be interpolated inside the shape.

By applying the alpha shape algorithm without setting $\Phi$, Delaunay triangles (DTs) among the points are formed to connect all points in the PGs to each other. As shown in Figure 1, there are a number of G/L types that are included in $PG_1$, $PG_2$, and $PG_3$. The DTs cover the points in the intermediate blank parts even though they belong to different geological areas. This generates crosstalk $Cr_2$ (Figure 5b), which covers those blank areas. Compared with the relatively smaller DTs inside $PG_1$ or $PG_2$, the size of the DTs across two PGs are big and the distances from their internal points to most neighboring sampled points are far. Therefore, the reliability of the estimated points inside these big DTs is low for pseudo training data. To avoid $Cr_2$, it is necessary to correctly select the threshold of $\Phi$, to divide $PG_{Geo}$ into $PG_{Geo}\psi,\omega$, where $PG_{Geo}\psi,\omega$ belongs to $PG_\omega$, $\omega = 1, 2, 3$, and then to obtain the optimized shape of the envelope of each $PG_{Geo}\psi,\omega$. Therefore, for all data points, a threshold $\Phi_1$ that forms three independent parts in the alpha shape process is needed to be specified first.

When the points are concentrated at different places on the plane that consist of different groups, for all points, the value of $\Phi$ within the PGs is small but the amount of small $\Phi$ is large, while the value of $\Phi$ between the PGs is large but the amount of such large $\Phi$ is small. If the threshold of $\Phi$ is set to cover most of $\Phi$ within the PGs and discard the remaining ones, the DTs between the PGs will disappear and the envelope of each PG will be generated. To achieve this, determining the turning point between the histogram of $\Phi$ from the maximum position to infinite is considered effective to find $\Phi_1$, which can be actualized by, for example, a triangle thresholding algorithm [34]. The histogram of $\Phi$ of all the datasets used in this research is shown in Figure 6a. According to the triangle thresholding algorithm, $\Phi_1$ is specified by the following process: Connect the peak value and the value at infinity by a line, search for the furthest point $I_1$ from the line on the histogram envelope, and specify the corresponding $\Phi$ of $I_1$. The alpha shapes are calculated under the conditions of setting $\Phi$ to infinity (Figure 6b), $\Phi_i < \Phi < +\infty$ (Figure 6c), and $\Phi = \Phi_i$ (Figure 6d). The calculated $\Phi_1$ is 0.0848 degree in longitude and latitude, defined as $\Phi_1$ here, and the length is approximately 10 pixels in our case. For each $PG_{Geo}\psi$, it is divided into $PG_{Geo}\psi,\omega$.

In a number of cases, it is clear that there are multiple PGs, $PG_{Geo}\psi,\omega,\chi$, $\chi = 1, \ldots, \Omega$, not connecting with each other that belong to the same $PG_{Geo}\omega$. The distances $D_\chi$ between these areas may be smaller than $\Phi_1$, which indicates that this alpha shape algorithm is not effective to divide all observed independent PGs, leaving crosstalk parts with adhering geological area $Cr_3$ (Figure 5c). Here, $\Omega$ is the total number of these independent PGs in $PG_{Geo}\psi,\omega$. We first directly performed kriging interpolation while ignoring the effect of $Cr_3$. Then, to handle $Cr_3$, G/L type labels were plotted for every pixel on the map in accordance with the data, as shown in Figure 1, to create image masks, $M_{Geo}\psi$ for excluding those points not covered by $M_{Geo}\psi$. After the filtering process, $Cr_3$ was also eliminated, and kriging interpolation on a single geological area could be successfully performed.

It should be noted that all vacant pixels in a DT must correlate with at least three neighboring points (the apaxes of the correspondent DTs and three points determine a plane) in accordance with kriging interpolation, where $\Phi$ of a DT must be smaller than $r_\psi$. If not, several vacant pixels located at the center of a DT will not be subject to kriging interpolation. Therefore, to form an alpha shape of each $PG_{Geo}\psi$, $\Phi_1$ should be set as $\Phi_1 = \min(\Phi_1, r_\psi)$.

If we did not apply $M_{Geo}\psi$ after the DTs were formed and before $\Phi_1$ was specified, a number of badly estimated pseudo training data points would be included, even though part of the process could be omitted. In cases when the distance between $PG_1$ and $PG_2$ and $D_\psi$ were smaller than $r_1$ (the example in Figure 5b), vacant pixels in DTs included in $Cr_2$ would be also interpolated. Even
if $Cr_2$ was eliminated by $M_{\text{Geo}_{\psi}}$ cropping, those badly estimated points (inside the green dashed line surrounding the region in Figure 5b) remained in the candidate training region. If we followed the process mentioned above, although there were still a number of less reliable areas remaining, the surface would be greatly suppressed (green dashed line surrounding the region in Figure 5d).

![Diagram](image)

**Figure 5.** Conceptual drawings of Figure 1 to explain how to spatially divide kriging interpolation regions and machine learning (ML) regions on a geochemical map: (a) The case of only setting a single $r$ by fusing all sampled points in the study area to kriging interpolation. Distribution crosstalk $Cr_1$ occurs in the regions near the border of different geological areas; (b) Delaunay triangles (DTs) generated by applying the alpha shape algorithm without setting $\Phi$. Crosstalk $Cr_2$ occurs at the intermediate parts between different PGs; (c) DTs generated by applying the alpha shape algorithm with setting threshold at $\Phi_t$. Remaining crosstalk $Cr_3$ occurs at the intermediate parts between different $PG_{\text{Geo}_{\psi,\omega}}$; (d) a geological mask $M_{\text{Geo}_{\psi}}$ is applied; $Cr_3$ is also eliminated.

According to Equation (3), an inverse computation of an $\left(N \times N\right)$ matrix is included in kriging interpolation. To ensure the computation speed of the computer, we compromised by generating pseudo training data using only three neighboring points. Thanks to the alpha shape algorithm, interpolation was processed independently in a single DT unit by using the apexes’ values.

The result after kriging interpolation in all $PG_{\text{Geo}_{\psi,\omega}}$ is shown in Figure 7. Here, T areas is the colored area and the remaining white areas are P areas, respectively. However, not every interpolated data point in the T areas can be treated as pseudo training data. As mentioned above, because the
further the distance from the interpolated points to the measured points is, the lower the reliability of the interpolated value. A large amount of relatively low reliability data will reduce the reliability of the constructed prediction model. To ensure reliability, a penalty on selection probability \( Q \) going into the training dataset is given to each data point. The selection probability is proportional to the minimum distance \( b_{\min} \) from the candidate interpolated point \( A_j \) in a DT to the three apexes’ measured data \( S_{\alpha} \), where \( \alpha = 1, 2, 3 \). \( b_{\min} \) can be expressed by Equation (15), and \( Q \) can be expressed by Equation (16):

\[
b_{\min} = \min_{A_j \in DT} \| A_j - S_{\alpha} \| \quad \alpha = 1, 2, 3
\]

\[
Q_\psi = \begin{cases} 
\left( \frac{\Phi_\psi - b_{\min}}{\Phi_\psi} \right)^2, & b_{\min,\psi} < \Phi_\psi \\
0, & b_{\min,\psi} \geq \Phi_\psi \text{ or } A_j \notin DT
\end{cases}
\]

The equations indicate that the apexes (\( b_{\min} = 0 \)), which are also the measured points, will be absolutely selected, and the points outside these DT areas will not be selected. Whether the interpolated data in the T areas will be selected as pseudo training data is determined by their \( Qs \). Finally, the flow chart of pseudo training datasets in our case is shown in Figure 8.

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**Figure 6.** Applying the alpha shape algorithm to our case: (a) Specifying \( \Phi_t \) by triangle thresholding algorithm; (b) alpha shape under the condition of setting \( \Phi \) to infinity; (c) alpha shape under the condition of setting \( \Phi \) to \( \Phi_1 < \Phi < +\infty \); (d) alpha shape under the condition of setting \( \Phi \) to \( \Phi_1 \).
Figure 7. Completed kriging interpolation copper content distribution map. Colored parts are defined as T areas and white parts are defined as P areas.

Figure 8. Flow chart of pseudo training dataset generation processing in our case.

2.4. Algorithm Validity Test

The validity test of the proposed algorithm includes four parts: (i) Cross-validation of measured data, (ii) blind test in T areas, (iii) temporal stability test in P areas and (iv) predictor importance test. The validity of the constructed model by SABAMIN is compared with that by only RF, which does not merge the spatial autocorrelation into ML.

First, the cross-validation test of measured data was done to test the precision of the constructed prediction model. The measured data were the most reliable values in the datasets: If the prediction values in the cross-validation were near to the real value, the precision of the model was good.
The precision was evaluated by the root mean squared error (RMSE) between the predicted value and real value through 10-fold cross-validation. In the cross-validation, all training datasets were randomly divided into 10 groups, then 9 groups were selected for training, and the remaining 1 group was selected for validation. To compare the two models fairly, only real and predicted values in actual sampled points were picked up for calculating RMSE.

Second, blind testing in T areas was done to test the accuracy of the T area. If the accuracy is good, spatial continuity should still be reserved well by the model, that is, the predicted map should have a similar distribution to the original one. To confirm this, a blind test was conducted in the T area. During one test, data in the selected area were excluded from the training data, and the remaining data were used to construct the prediction model. The distribution of the excluded area was then predicted by the constructed model and compared with the real distribution. Only data from actual sampled points were selected in the evaluation. The RMSE and Pearson product–moment correlation coefficient ($R_t$) were used to evaluate the similarity between the predicted and real distributions. The study areas, $T_a$, $T_b$, and $T_c$ were selected for three different blind tests. To confirm whether the model could handle the prediction of geochemical anomalies, zones around the mining districts were considered as candidate areas for the test.

Third, the temporal stability test in P areas was done to test the accuracy of P area. Since there were no geochemical data in a P area, it was difficult to validate our algorithm by comparing the predicted value with the real value. We considered that, as shown in Figure 2, the copper content in the rocks around a copper mine should be high, so checking whether high copper content areas were located around copper mining districts was considered a compromised method for the reliability test. Furthermore, it was considered that if the reliability of the algorithm was high, the similarity between the different predicted distributions using ASTER images of different time series should be high. In all cases, the predicted high content areas should be located around the copper mines. The similarity of all pairs of predicted maps in different time series were calculated, and an average similarity of them was used to compare the reliabilities of SABAMIN and RF. The Pearson product–moment correlation coefficient ($R_p$) was used to evaluate the similarity. Three study areas, $P_a$, $P_b$, and $P_c$ in the P areas, which are all located around copper mining districts, were selected for these tests.

Fourth, the predictor importance of different groups $ImG$ was also calculated to check whether the constructed model was geologically reasonable and to determine whether the constructed model was biased to any predictor. If the predictors in the high importance ranking were the factors which were already known to be highly correlative to copper content-based conventional geological knowledge, for example, high copper contents tend to distribute around copper mine restricts which are always located in the mountain, and if the topographic factors were in the high importance ranking, the model was considered geologically reasonable. To check predictor bias, first, the normalized importance of all predictor variables $Im_{w}$, $w = 1, \ldots , 63$, were calculated by the RF, and then, in accordance with Table 2, they were allotted to different groups $ImG_{mag}$, $ImG_{dem}$, $ImG_{band}$, $ImG_{lith}$, $ImG_{cor}$, which represent geomagnetic variables, DEM variables, ASTER band variables, ASTER lithological index variables, and coordinate variables groups, respectively. $ImG$ was obtained by summing all of the members’ importance in the group. As a good selection of predictor variable, the constructed model should not be biased to any predictor.

Random processes are seen in both the RF and SABAMIN. In the RF, the random creating node of trees by randomly selecting training variables induces a random process. That random selection process for creating a prediction model generates a random factor in SABAMIN. As a result, the calculated RMSE, $R_t$, and $R_p$ are not constant values. Therefore, their means and standard deviations are calculated from 30 repeated tests and used for comparison in tests (i) and (ii), and the average predicted distributions of the repeated tests are used in test (iii). Moreover, the mean and standard deviation of the predictor’s importance are calculated from 30 repeated tests. Two-sided $t$-tests are used to examine the differences between groups. For all analyses, the statistical significance is set to $p < 0.05$. 
3. Results

3.1. Cross-Validation Test of Measured Data

The 10-fold cross-validation results of the RF and SABAMIN are shown together in Figure 9. The test was repeated 30 times. As a result, the RMSE of the RF and SABAMIN was 329.10 ± 0.53 and 281.54 ± 1.04 ppm, respectively. A significant difference is seen between the precision test results, which means the constructed prediction model by SABAMIN was more accurate than that of RF.

![Figure 9. Ten-fold cross-validation results of random forests (RFs) (dots) and SABAMIN (triangles). The asterisk means there is a significant difference (p < 0.05) between them.](image)

3.2. Blind Test in T Area

The blind test results of T_a, T_b, and T_c are shown in Figure 10a–c, respectively. In each panel, the real distribution, RF-predicted distribution, and SABAMIN-predicted distribution are compared. The results of the RMSE and $R_t$ are shown in Table 4. In both RF and SABAMIN prediction, all $R_t$s exceeded 0.6, the distribution of high and low copper contents were almost correctly predicted, and their predicted distribution was highly similar to the real distribution, which means that, like the RF, SABAMIN can effectively use sampled points to predict other completely vacant areas. Nevertheless, compared with the RF, the RMSE and $R_t$ were improved by SABAMIN, and the RF-predicted distribution was noisier than that of the SABAMIN-predicted one. In the local areas marked by black arrows, the SABAMIN predicted areas are closer to the real ones, while those of the RF deviated. These results suggest that the spatial autocorrelation is reserved well in the prediction model.

![Figure 10. Cont.](image)
Table 4. Root mean squared error (RMSE) and $R_t$ results of the RF and SABAMIN. Tests are repeated 30 times, with the (mean ± standard deviation) listed.

| Area | RMSE RF | RMSE SABAMIN | $p$-Value | $R_t$ RF | $R_t$ SABAMIN | $p$-Value |
|------|---------|--------------|-----------|---------|--------------|-----------|
| $T_a$ | 172.5 ± 5.5 | 133.9 ± 3.9 | 0.000 ** | 0.546 ± 0.052 | 0.629 ± 0.050 | 0.000 ** |
| $T_b$ | 419.0 ± 10.5 | 410.6 ± 9.0 | 0.001 * | 0.757 ± 0.032 | 0.779 ± 0.030 | 0.005 * |
| $T_c$ | 431.2 ± 8.8 | 398.6 ± 7.3 | 0.000 ** | 0.737 ± 0.038 | 0.756 ± 0.020 | 0.021 * |

* $p < 0.05$; ** $p < 0.001$.

3.3. Temporal Stability Test in P Areas

The results of the temporal stability test in the P areas by $P_a$, $P_b$, and $P_c$ are shown in Figure 11a–c, respectively. Three predicted distributions by the remote sensing data (see Table S5 in the supplementary material) in three different temporal series ($tem_1$, $tem_2$, $tem_3$) approximately covering the same region are shown together in temporal order. For the reference, reconstructed images by ASTER band 1, 2, and 3N images are shown together. In all temporal series of all test areas, the copper content around a mining district was predicted to have a high value, which is considered to be a reasonable result. Even if the ASTER images were captured at different timings, the predicted distributions would look very similar. Table 5 shows the similarity between the pairs of $tem_1$, $tem_2$, and $tem_3$, which was evaluated by $R_p$. All $R_p$s almost exceeded 0.5, which indicates that the prediction was steady temporally. The results above suggest a high reliability of our algorithm.
Figure 11. Cont.
Figure 11. Temporal stability tests in P area results of (a) Pa, (b) Pb, and (c) Pc. Red diamonds show the mine position. The crowd of the mine positions corresponds with the mining district shown in Figure 3.

Table 5. Rp results between pairs of tem1, tem2, and tem3. Tests are repeated 30 times, with the (average ± standard deviation) listed.

| Area | Rp tem1 vs. tem2 | Rp tem1 vs. tem3 | Rp tem2 vs. tem3 |
|------|----------------|----------------|----------------|
| Pa   | 0.681 **         | 0.347 *         | 0.548 *         |
| Pb   | 0.756 **         | 0.710 **        | 0.834 **        |
| Pc   | 0.743 **         | 0.581 **        | 0.562 **        |

*p < 0.05; ** p < 0.001.

3.4. The Predictor Importance Test

The results of the predictor’s importance analysis by RF are shown in Figure 12 and Table 6. From the results that show the importance of coordinates are ranked at first and third, the spatial information is the most important factor in our constructed model, which corresponds with the knowledge that spatial information is crucial for geochemical mapping, (see previous study [5,6]). Next, the altitude and slope are ranked at second and sixth, which corresponds with the fact that copper mines (high copper content areas) are always located in the mountains in the Arizona area [35], and mining changes the topographic surface. Of the four geomagnetic predictors, one was ranked in the top 10, and the remaining three were ranked in the top 20, which corresponds with the fact that metal elements influence the geomagnetics. In this copper prediction case, the ASTER lithological index obtained by cross band calculation, such as Mg-OH, muscovite, dolomite, kaolinite, silica, quartz, and alunite are in high ranks, and are commonly seen alteration minerals exposed around active mines. All the above information demonstrates that the constructed model is reasonable for geochemical contents mapping.
A comprehensive look at Figure 12 and Table 6 does not show any significance that any factors were more overwhelming than all other factors. All predictors were fairly treated to construct the model.

![Figure 12. Importance of all predictors.](image)

**Table 6.** Total predictor’s importance in each variable group.

| Group                  | Predictor Number | ImG (Mean ± SD) |
|------------------------|------------------|-----------------|
| Geomagnetic            | 4                | 0.088 ± 0.012   |
| DEM                    | 2                | 0.088 ± 0.018   |
| ASTER Lithological Index | 28            | 0.210 ± 0.023   |
| ASTER Band             | 28               | 0.457 ± 0.025   |
| Coordinates            | 2                | 0.158 ± 0.021   |

4. Discussion

By applying the data augmentation strategy, pseudo training datasets were created using the kriging interpolation model. The accuracy of the ML model was improved with augmented data, which agree with applications in different fields [36,37]. The geochemical content in the majority positions were only at the background level, and the high regions’ performances were considered to be anomalies on the map. In a univariate interpolation, it is difficult to describe all the geological anomalies perfectly by maximum likelihood estimation using such data with high skewness; thus, usually the geological anomalies sometimes contradict the distant geospatial association. In this study, to avoid that, although we still calculated the weight by kriging, in the interpolation operation, the vacant pixels in a triangular area were only interpolated by the three most neighboring points and sacrificing the contribution by some low weight measured points. This might be the one reason that an only slight improvement of RMSE was observed in our study. Recently, Kim et al. [38] proposed a curvature interpolation method which is suggested to be a more accurate univariate interpolation.
method for geospatial data than IDW, and is considered a candidate method to improve the accuracy of pseudo training data generation. ML usually only constructs a model to fit the majority in the datasets, where the minority tends to be ignored by the model. As a result, the high values are underestimated. This might be another reason for slight RMSE improvement. By the hint of mineral prospectivity mapping studies [39–41], the geochemical distribution can be treated as classification labels. It is observed from Figure 10 that, although predicted distribution does not perfectly correspond with the real one, a high accuracy of classifying high and low contents should be achieved. Therefore, the prediction accuracy can be further improved using the method that firstly classifies the study area into different classes, and then constructs different regression models in the different region with different class to ensure all data can be correctly estimated in specific ranges.

In the temporal stability test, ASTER images in different temporal series were used for the test. We hypothesized that there was no or only slightly topography alteration in the short term, and by ignoring some human-made changes, ASTER images in different temporal series should be the same. Therefore, the predicted geochemical distribution should be similar in this case. However, the ASTER images show the solar optical reflectance information from the Earth’s surface, and despite the effect of the surface vegetation coverage condition, the reflected luminance may change due to different solar heights and angles at different times in the day or during different seasons. This unstable component was regarded as signal noise in temporal series. Although it is possible to achieve a higher similarity in $P_a$, $P_b$, and $P_c$ by correcting them in accordance with the solar zenith angle [42], a reasonable result that predicted high content regions are located at or very near to actual mining districts in all temporal series indicated that our prediction model was robust to this fluctuation induced by solar condition changing. Thus, the signal of the predictor variables seem to be more determinative than the noise in the prediction process.

5. Conclusions

In this study, to provide a high accuracy model for generating a high-resolution geochemical map, we proposed a new algorithm, SABAMIN, which merged both spatial location and autocorrelation into an ML model by using data augmentation and computational geometry strategies. We applied kriging interpolation to generate pseudo training datasets and applied an alpha shape method to build these pseudo training datasets to become geologically reliable.

In the blind test results, a higher similarity meant the spatial autocorrelation was reserved well in the prediction model. Furthermore, coordinate variables were ranked at the top level in the predictor’s importance test. These results suggest that we successfully merged spatial autocorrelation into the ML model by only co-training with the sampled datasets and pseudo training datasets.

Compared with the RF constructed model, SABAMIN achieved a lower RMSE in the cross-validation test. The model also performed a steady prediction in a temporal stability test using multiple ASTER images in different temporal series. The effect and reliability were confirmed by these results.

According to the ranking of predictor importance, the machine learning suggested that the major factor related to copper content distribution corresponded with common geological knowledge, which suggests that the constructed model is reasonable for geochemical contents mapping.

In summary, SABAMIN is effective for generating high-resolution geochemical maps with high accuracy. Combining the univariate interpolation method with multivariate prediction with data augmentation also proved effective for geological studies.

In the future, we will further improve the pseudo training data generation method or try to provide different prediction model to different region with contents in different levels to improve the accuracy of the prediction model.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2072-4292/12/12/1991/s1, Figure S1: The detail of remote sensing data: (a) ASTER band 1 data; (b) DEM altitude data; (c) Analytic signal processed geomagnetic data, Table S1: Detailed introduction of the study areas, Table S2: The default filename of source files of ASTER data, Table S3: The default filename of source files of DEM altitude data, Table S4: The
default filename of source files of DEM elevation data, Table S5: The default filename of source files of ASTER
data used in temporal stability test,

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