

Analyzing Spatial Patterns and Stochastic Interpolation of Cobalt Values from Vancouver Island Using Simple Kriging Model

Introduction

This following study is intended to gain further perspectives on the statistical methods of geochemical prospecting. Analyzing the spatial distribution of element concentration values and recognizing unusual features in the geographical context for potential mineralization is a major concern in geochemical prospecting (Howarth, 111). The study is based on sample data of cobalt values from Vancouver Island in British Columbia collected by the Geological Survey of Canada. This investigation attempts to analyze the pattern of spatial dependencies among the cobalt values and carry out stochastic interpolation of cobalt values in terms of a simple kriging model in order to identify potential cobalt deposits.

The Vancouver sample data extends over the area at the northern tip of the island shown in the Figure 1.1a and the area outlined in red denotes the full extent of the data site. For purposes of analysis, a smaller set of 286 sample sites was selected, as shown by the dots in Figure 1.1b.



Figure 1.1a. Vancouver Sample Area

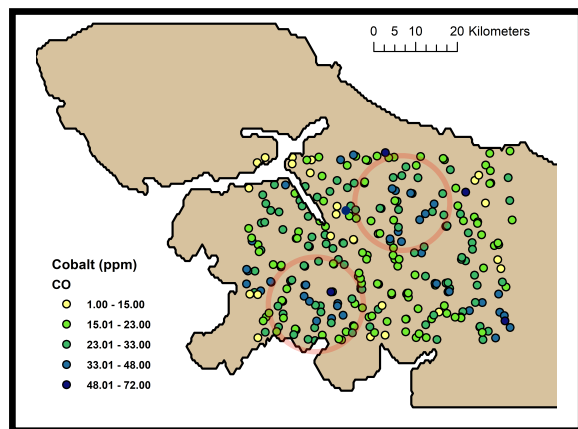


Figure 1.1b. Vancouver Sample Area

We explore distributions of patterns of cobalt elements here because cobalt (Co) is a potentially critical mineral increasingly used in magnets and rechargeable batteries. The disposition of the cobalt sample points in this study varies based on the nature of investigation and stream sediment or lake sediment sampling is dependent on drainage pattern (Howarth, 136). We can find the curvilinear patterns of the sample points. As with many geochemical surveys, samples are here taken mainly along stream beds and lake shores, where minerals deposits are more likely to be found (Smith, 33). In this sense, we want to use kriging, which is a statistical method of interpolation, to prospect cobalt values across our sample area. According to the general principles of geochemical prospecting, mineral deposits represent anomalous concentration of specific elements and mineral deposits has a central zone where valuable elements surrounding the zone decrease in concentration and we measure this concentration in parts per million (ppm)

or parts per billion (ppb) (Horsnail, 2). As shown by the red circles in Figure 1.1b, we can see that there are potential “central zones” of cobalt deposits that need further investigation.

Geochemical mapping have been executed in British Columbia to display and investigate concentrations of chemical elements in the past. One such study is R.E.Lett’s paper that describes a reconnaissance scale regional geochemical survey carried out over the Bowser Lake in Northwestern British Columbia (Lett, 61). The survey produced new multi-element geochemical data and found abundant gold grains in two creeks, which reflected results of the Iskut-Palmiere prospect. The survey also found spatial dependencies of gold grain counts as gold grain counts are higher at sites close to the Eskay Creek mine site and lower at areas far from the mine site. Drawing on this paper, our study will explore the spatial dependencies of cobalt values at the Vancouver Sample Area and perform geochemical prospecting on the cobalt values.

Methodologies and Results

Since the mapped cobalt values exhibits strong similarities between neighboring values (as shown in Figure 1.1b), we can expect to find a substantial range of spatial dependence in this cobalt data. However, we can also see that the covariance-stationarity assumption of Isotropy, which states that similarities between concentration levels at different locations depend only on the distance between them, is questionable for this data. We can see diagonal “waves” of high and low cobalt values rippling through the sample area and we want to explore the spatial patterns of cobalt values in the variogram estimation procedure to follow.

Analysis of spatial dependencies is first carried out using empirical variogram estimation in Matlab. The key difference between continuous spatial data of cobalt values and point patterns is that there is a meaningful value at every location in the study area. Since our spatial variable in this case tend to exhibit some degree of continuity over space, we expect these variables to exhibit similar values at locations close together in space. In order to identify spatial trends in the spatial stochastic process, we expect that for sites that are sufficiently close together, the associated spatial residuals not captured by global trend will tend to exhibit statistical dependence, which is measured by the covariance of these spatial residuals. In particular, positive dependencies among spatial residuals will tend to be reflected by positive covariance among these residuals.

Before proceeding, it is important to emphasize the notion of spatial stationarity to model covariances among spatial random effects in cobalt values. According to the Spatial Random Effects Theorem, for any random vector of multi-location effects comprised of a sum of individual random factors with zero means and covariance matrices, if the distributions of these random factors are “not too different”, and the dependencies among these random factors are “not too strong”, then the distribution of multi-location effects is approximately multi-normal. In that sense, our aim is to specify the unknown covariance matrix for the random effects in cobalt values and model these unobserved dependencies with general spatial dependencies that should be common to all these covariance structures.

A spatial stochastic process, $\{Y(s): s \in R\}$, is said to be covariance stationary if and only if the following two conditions hold for all $s_1, s_2, v_1, v_2 \in R$:

- (i) $E[Y(s_1)] = E[Y(s_2)]$, ii) $\|s_1 - s_2\| = \|v_1 - v_2\| \Rightarrow \text{cov}[Y(s_1), Y(s_2)] = \text{cov}[Y(v_1), Y(v_2)]$

The covariogram and its normalized form, the correlogram, are by far the most intuitive methods for summarizing the structure of spatial dependencies in a covariance stationary process (Smith, 1). Here, we will use $C(h)$ to represent the common covariance value such that $\text{cov}[Y(s), Y(v)] = C(h)$ and h represents distances. Since the covariance values, $C(h)$ are unique for each distance value, h , the function, C , of these distances is designated as the covariogram for a given covariance stationary process. However, as covariograms have particular units and are difficult to interpret, we analyze dependencies between random variables in terms of (dimensionless) correlation coefficients, which is a normalized form of the covariogram called the correlogram for a covariance stationary process. The correlation between any $Y(s)$ and $Y(v)$ is estimated by:

$$\rho[Y(s), Y(v)] = \frac{\text{cov}[Y(s), Y(v)]}{\sqrt{\text{var}[Y(s)]}\sqrt{\text{var}[Y(v)]}} = \frac{C(h)}{\sqrt{C(0)}\sqrt{C(0)}} = \frac{C(h)}{C(0)}$$

In our study, estimation of correlogram presents certain difficulties so we use variogram to display the variability between data points as a function of distance. The first step of our empirical variogram estimation is to aggregate point pairs of cobalt values with similar distances and hence estimate the estimator of the variogram value at only a small number of representative distances for each aggregate. We first partition distances into intervals, called bins, and take the average distance in each bin to be the appropriate representative distances, called lag distances. If N_k denotes the set of distance pairs, (s_i, s_j) , in bin k , [with the size (number of pairs) in N_k denoted by $|N_k|$], and if the distance between each such pair is denoted by h_k , then the lag distance h_k , for bin k is defined to be:

$$h_k = \frac{1}{|N_k|} \sum_{(s_i, s_j) \in N_k} h_{ij}$$

The empirical variogram is usually informative in terms of the possible shapes of the true variogram. To examine possible maximum distances for the desired variogram, we first observe (by an application of the measurement tool in ARCMAP) that a neighborhood of 20,000 meters around typical sites appears to be large enough to contain most positive dependencies with other sites. Then we choose to fit a spherical variogram, which is a widely used variogram model. It is defined for all $h \geq 0$ by:

$$\gamma(h; r, s, a) = \begin{cases} 0 & , h = 0 \\ a + (s - a) \left(\frac{3h}{2r} - \frac{h^3}{2r^3} \right) & , 0 < h \leq r \\ s & , h > r \end{cases}$$

where r denotes the maximum range of positive spatial dependencies (designated simply as the range of the variogram), s corresponds to the sill of the variogram, and a corresponds to the nugget.

The choice of the maximum lag distance also involves some implicit restrictions as it cannot be greater than half of the maximum pairwise distance. We observed (again by using the measurement tool) that half of the maximum pairwise distance in this case is about 40,000 meters. Therefore, we first used a reasonable maximum distance of 20,000 meters and then used a maximum distance of 40,000 meters to construct two spherical variograms. As shown in Figure 1.2a, the blue dots are the empirical variogram points maximum distance of 20,000 meters and the estimated spherical variogram is shown in red, which rise toward a “sill” at about 17,010

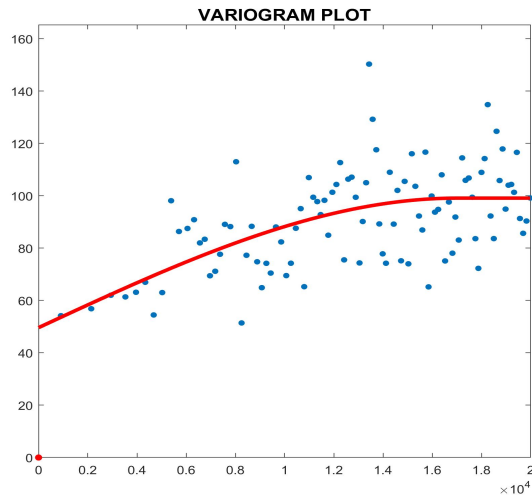


Figure 1.2a: Fitted Spherical Variogram

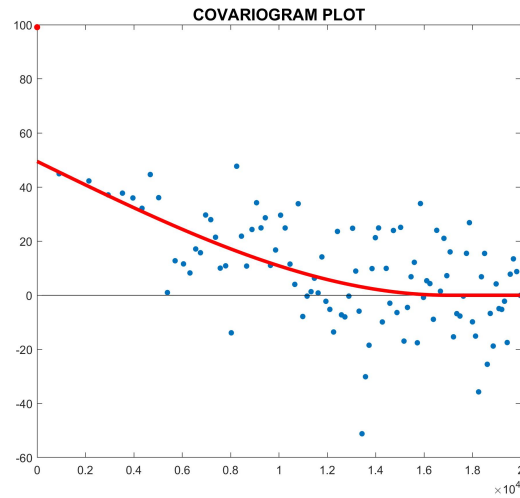


Figure 1.2b: Derived Spherical Covariogram

meters according to parameter estimate results. Figure 1.2b is the derived spherical covariogram corresponding to the empirical variogram in Figure 1.2a. It is clear that 17,010 meters is about the distance at which covariance (and hence correlation) first falls to zero. It denotes the distance beyond which there is estimated to be no statistical correlation between cobalt values. Turning to the other estimated parameters, note first from Figure 1.2a that the sill is about 99, which is seen to be the estimated variance of individual cobalt values (i.e., the estimated covariance at “zero distance”). Similarly, as shown in Figure 1.2b, the nugget is about 50, which is seen to be that part of the individual variance that not related to spatial dependence among neighbors. Since in this case the relative nugget effect is 0.51 ($=50/99$), the underlying process exhibits some degree spatial dependence.

Then we fit a spherical variogram with maximum distance of 40,000 meters as shown in Figure 1.3a, which still rise toward a “sill” at about 13,767 meters. In this case, the sill is about 92 according to the parameter estimate and the nugget is about 49. Therefore, the relative nugget effect is 0.53 ($=49/92$), so the underlying process exhibits some degree spatial dependence.

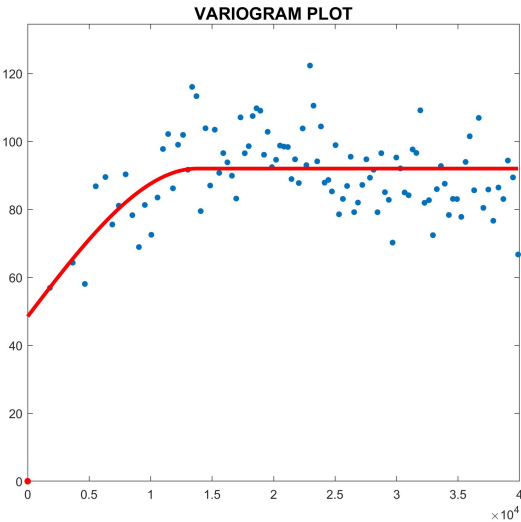


Figure 1.3a: Fitted Spherical Variogram

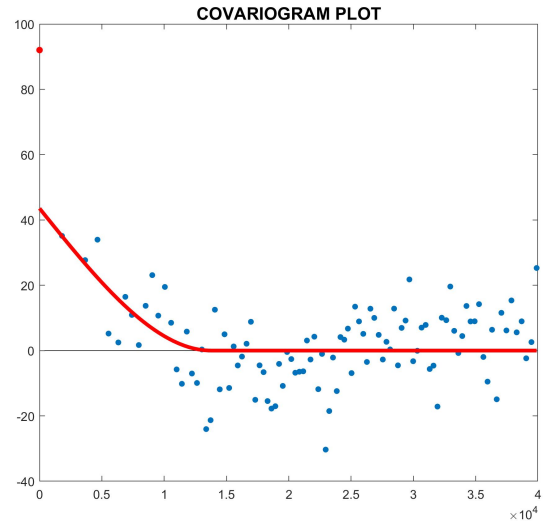


Figure 1.3b: Derived Spherical Covariogram

While we compare these two derived covariograms in Figure 1.2b and Figure 1.3b, it is important to note that the vertical (squared difference) scales for these two figures are the same, but the horizontal distance scales are now different. While the segment of Figure 1.3b up to 20,000 meters is qualitatively similar to Figure 1.2b, the bins and corresponding lag distances are not the same as in Figure 1.2b. Given this scale difference, we can see that the covariogram in Figure 1.3b shows a rise starting at about 20,000 meters, which can be interpreted to mean that pairs of y-values (cobalt measurements) separated by less than 20,000 meters tend to be more similar (positively correlated) than those separated by slightly larger distances. By again using the measurement tool in ARCMAP, it can be seen that the spacing of successive waves is about 20,000 meters. So it does appear that this effect is being reflected in the empirical variogram and derived spherical covariogram.

In the next step, we will use the variogram estimate obtained for the 20,000-meter max distance and simple kriging model to carry out a stochastic interpolation of cobalt values. Kriging model is a spatial prediction models that predict values based on local information. Simple kriging assumes that underlying stochastic process itself is entirely known and that the spatial trend is constant. We treat the observed data as a finite sample from a spatial stochastic process $\{Y(s) : s \in R\}$ and we assume that some appropriate subset of sample locations, $S(s_0) \subseteq \{s_i : i = 1, \dots, n\}$ has been chosen for prediction. To determine a prediction $\hat{Y}(s_0)$ based on sample data, we determine the prediction as a function of the random variables, $\{Y(s_1), \dots, Y(s_{n_0})\}$ associated with the observed data. We hypothesize that $\hat{Y}(s_0)$ can be represented as some linear combination of these random variables:

$$\hat{Y}(s_0) = \sum_{i=1}^{n_0} \lambda_{0i} Y(s_i)$$

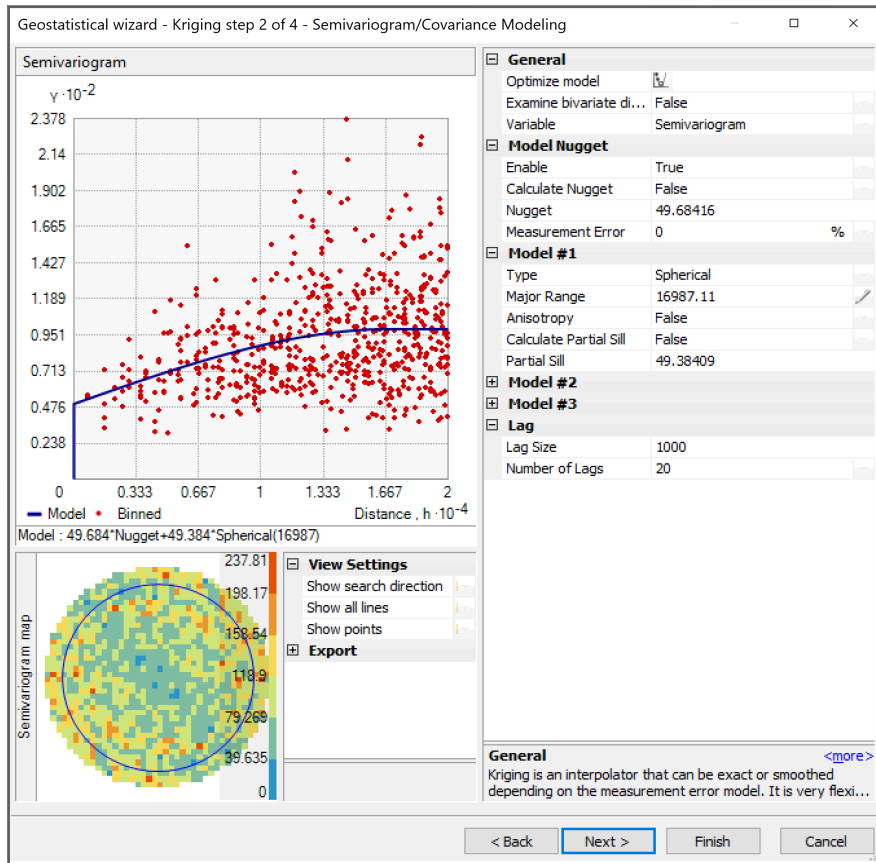
where the weights λ_{0i} are yet to be determined and this hypothesis is referred to as the linear prediction hypothesis. We will use simple kriging model by derive optimal prediction weights in our analysis.

If we want to predict the cobalt value, $Y(s_0)$, at some location, then since $\mu(s_0) = \mu$ is already known, we see from the identity $Y(s_0) = \mu + \varepsilon(s_0)$ that it suffices to predict the associated error with optimal weights:

$$\hat{\varepsilon}(s_0) = \sum_{i=1}^{n_0} \lambda_{0i} \varepsilon(s_i)$$

We first krige this data in MATLAB and use the estimated parameter values to predict cobalt value at a point location (615000, 586500) using a kriging bandwidth of 5000 meters. According to the results of the variogram estimate obtained for the 20,000-meter max distance, the estimated range is 17,009, the estimated sill is 99, and the estimated nugget is 50. After we run the prediction using simple kriging, we get the predicted cobalt value as 17.73 and the standard error of prediction at the point is 8.04. Then we construct a 95% prediction interval for Cobalt value at this location, which is (4.44, 31.02). This means that there is a 95% probability that the cobalt value will be contained within this prediction interval.

Then we will use the spherical variogram parameters for 20000-meter max distance case to krige the cobalt data using the Geostatistical Analyst extension in ARCMAP. Before Kriging the Cobalt Data, we first calculate the mean Cobalt value using the attribute table and the mean value is 25.27. Then we use Geostatistical Wizard and get a mean value of 25.27 as well.

**Figure 1.4 Spherical Variogram Fit in Geostatistical Analyst**

As shown in Figure 1.4, by setting the number of lags to 1,000 and choosing a constant bin size of 20 meters (as seen in the Lag window in the lower right of Figure 1.4), we will obtain a maximum distance of exactly 20,000 meters (as seen on the distance axis of the variogram plot). The fitted spherical variogram is shown by the blue curve in Figure 1.4, and the empirical variogram is shown by red dots. Now we compare the Major Range and Nugget values in Geostatistical Analyst with the Range and Nugget computed in MATLAB. A comparison of the parameter estimates using both MATLAB and GA in this example (Figure 1.5 below) show that in spite of the differences above, they are qualitatively very similar.

	MATLAB	GA
Range	17,009.8	16,987.1
Sill	99.1	99.1
Nugget	49.6	49.7

Figure 1.5 Parameter Estimates

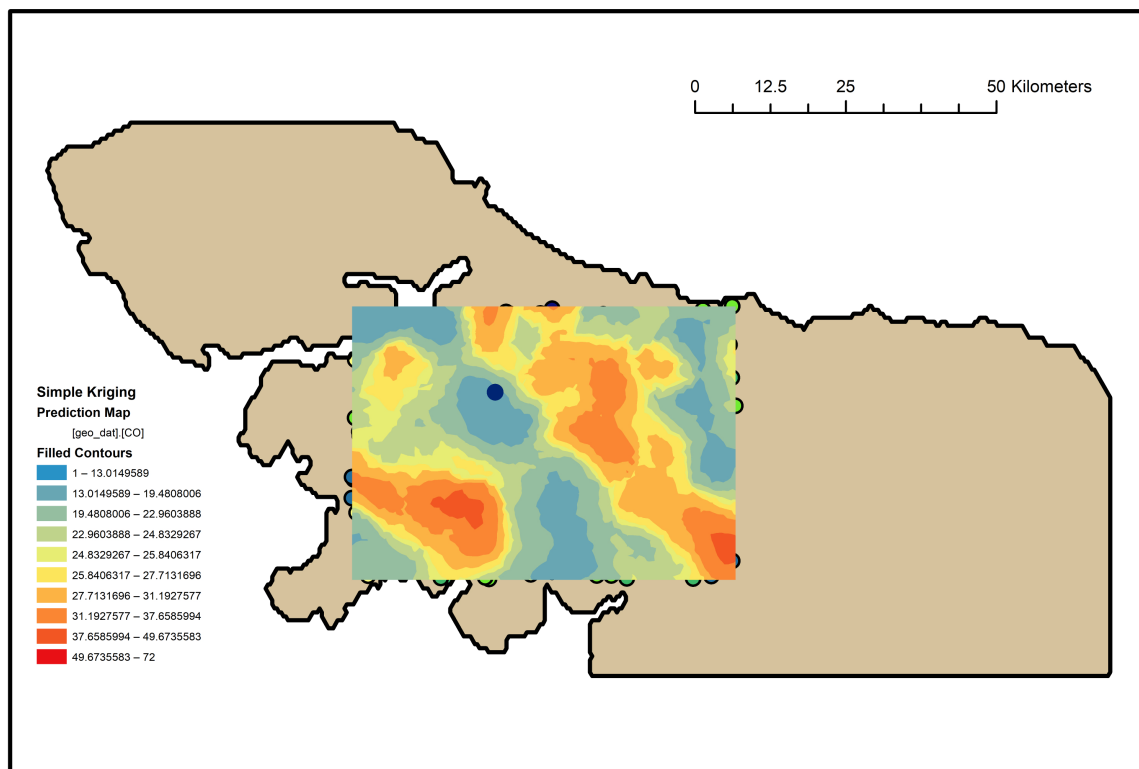


Figure 1.6 Simple Kriging

As shown in Figure 1.6, spatial interpolation of cobalt values is executed using simple kriging model in Geostatistical Analyst. The deep blue point is the point at (615000, 586500) and we want to examine the cobalt values near this point and get the mean of these cobalt values. We first choose a sample of five neighboring kriged cobalt value, which are 17.6, 17.9, 18.5, 18.7, 16.2, and calculate the average of these values, which is $17.78 = (17.6 + 17.9 + 18.5 + 18.7 + 16.2) / 5$. Then we compare it with the our predicted cobalt value (17.73) in Matlab and find that they are very similar. This means that the cobalt value predicted by Geostatistical Analyst using simple kriging is almost the same as the cobalt value predicted in MATLAB.

Then we want to select six sites surrounding the point at (615000, 586500) and calculate the mean value of these six cobalt values as shown in Figure 1.7. From the results in ARCMAP, the mean cobalt value from these six sites is 16.5, which is very similar to the predicted value we calculated in MATLAB and the mean value of our previous sample of five neighboring kriged cobalt values. Therefore, the kriged value seems reasonable in relation to these selected data points.

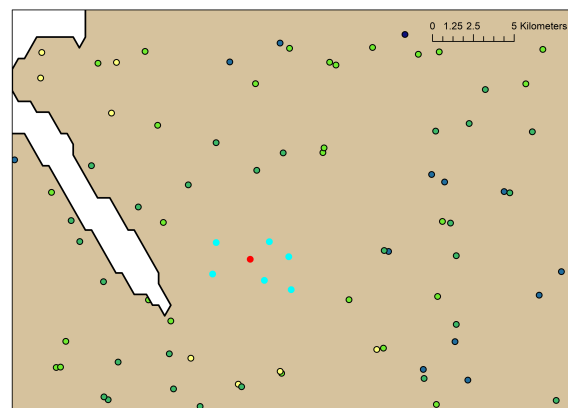


Figure 1.7 Six Selected Sites

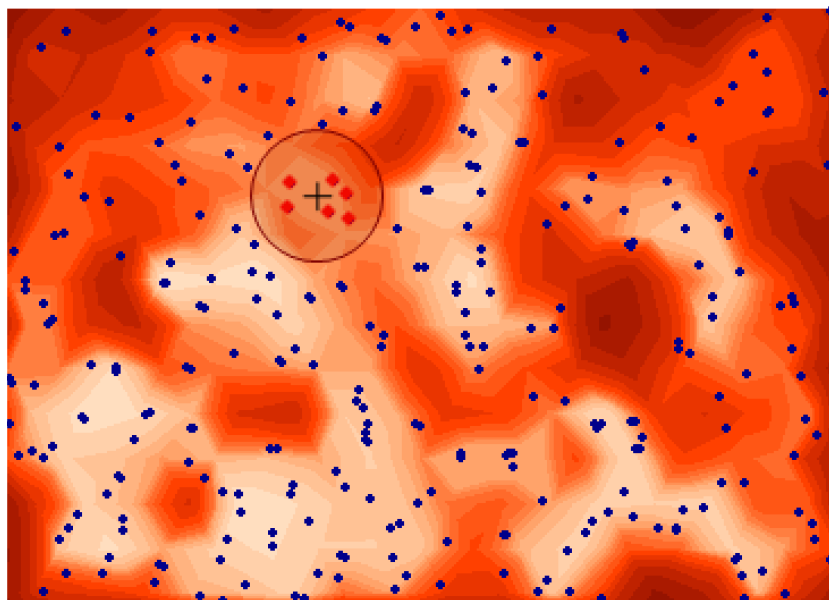


Figure 1.8 Standard Error of Prediction Value

Then we will again examine the prediction-error value at point (615000, 586500) using Geostatistical Analyst. The six neighbors of the point that have been used in the Kriging prediction are shown in the red circle in Figure 1.8. The standard error of predicted value at point (615000, 586500) is 8.04, which is exact the same as the prediction-error value we get in MATLAB. Areas in red have higher standard errors of predicted values than areas in orange.

Finally, we investigate the simple kriging prediction map in Figure 1.6 and see that it shows spatial dependencies of predicted cobalt values. As there are three spots that have very high predicted cobalt values, predicted cobalt values close to these three spots are higher and predicted cobalt values far from these three spots are lower. We also compare the simple kriging prediction map in Figure 1.6 and the standard error of predicted value in Figure 1.8 and see that they both show diagonal “waves” of high and low predicted cobalt values and standard errors of predicted values. This corresponds with our expectation that an assumption of covariance stationarity might be an over-simplification of this spatial data pattern. These waves in standard error of predicted value are roughly parallel to the Pacific coastline, and would seem to reflect the history of continental drift in this region.

Discussion

We first analyzed the spatial dependencies of cobalt values from Vancouver Island using fitted spherical variograms and derived spherical covariograms. We arrived at results that pairs of cobalt values separated by less than 20,000 meters tend to be more similar (positively correlated) than those separated by slightly larger distances. We then run simple kriging model in both MATLAB and ARCMAP to predict cobalt values across the sample area and arrived at very similar prediction results at a specific point. Our findings in the spatial interpolation of cobalt values correspond with R.E.Lett’s paper that sites close to the “central zone” of cobalt values

have higher predicted values and sites far from the “central zone” have lower predicted cobalt values (Lett, 65).

Although the calculated relative nugget effect shows the underlying process exhibits relatively little spatial dependence, the relative nugget effect at a maximum distance of 20,000 meters is smaller than at a maximum distance of 40,000 meters. This means that there is relatively more spatial dependencies when pairs of cobalt values were separated by less than 20,000 meters. From the simple kriging prediction map, we can also conclude that while the “wave” effect is being reflected in the spatial interpolation and an assumption of covariance stationarity might be an over-simplification of this spatial data pattern, our predicted cobalt values do show “central zones” where valuable elements surrounding the zone decrease in concentration.

References

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