

## **Geostatistical Simulation of Geochemical Compositions in the Presence of Multiple Geological Units: Application to Mineral Resource Evaluation**

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1 **Geostatistical Simulation of Geochemical Compositions in the Presence**  
2 **of Multiple Geological Units - Application to Mineral Resource**  
3 **Evaluation**

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10  
11  
12 **Abstract**

13  
14 An accurate prediction of benefit in ore deposits with heterogeneous spatial variations requires  
15 the definition of geological domains that differentiate the types of mineralogy, alteration, and  
16 lithology as well as the prediction of full mineral and geochemical compositions within each  
17 modelled domain and across boundaries between different domains. This paper proposes and  
18 compares various approaches (different combination of log-ratio transformation, gaussian and  
19 flow anamorphosis, and deterministic or probabilistic geological models) for geostatistical  
20 simulation of geochemical compositions in the presence of several geological domains.

21 Different approaches are illustrated through an application to a nickel-cobalt laterite deposit  
22 located in Western Australia. Four rock types (Ferruginous, Smectite, Saprolite, and  
23 Ultramafic) are considered to define compositionally homogeneous domains. Geochemical  
24 compositions are comprised of six different components of interest (Fe, Al, Mg, Ni, Co, Filler).  
25 Results suggest that the flow anamorphosis is a vital element for geostatistical modeling of  
26 geochemical composition due to its invariance properties and capability for reproducing  
27 complex patterns in input data including: presence of outliers, presence of several populations  
28 (due to the presence of several geological domains), nonlinearity, and heteroscedasticity.

29

30

31 Keywords: Compositional data, Log-ratio, Flow Anamorphosis, Geostatistical simulation,  
32 Geological domaining

33

## 34 **1 Introduction**

35 Ore deposits usually consist of ore materials with different characteristics. In order to maximize  
36 revenue in a mining project, a decision must be made regarding what processing plants are  
37 needed and the processing destinations of ore materials with different characteristics. For a  
38 better classification of ore materials, features such as rock type, alteration, microstructure,  
39 geochemical and mineral composition must be measured at sample locations and be predicted  
40 at the block model of the target deposit. These features normally have complex statistical and  
41 spatial relationships which should be reproduced in the predicted models (Boisvert et al. 2013;  
42 Maleki et al. 2016; Mery et al. 2017; Montoya et al. 2012; Mueller et al. 2014; Talebi et al.  
43 2017; Tolosana-Delgado et al. 2014; Tolosana-Delgado et al. 2015; van den Boogaart et al.

44 2014). Among the different features of ore materials, mineral and geochemical compositions  
45 have a great impact on the final destination of materials and performance of processing plants.  
46 However, the compositional nature of these data induces several challenges for multivariate  
47 geostatistical techniques to predict them at the block support (Tolosana-Delgado et al. 2014;  
48 Tolosana-Delgado et al. 2015). Compositional data are multivariate, non-negative values  
49 which represent the importance of some parts of a whole. In such data, the constant sum  
50 constraint forces at least one covariance to be negative and induces spurious correlations.  
51 Furthermore, they carry just relative information (Aitchison 1986). To transform compositional  
52 data into unbounded space and to increase mathematical tractability, different log-ratio  
53 transformations can be applied prior to using standard (geo)statistical techniques (McKinley et  
54 al. 2016; Pawlowsky-Glahn and Egozcue 2016; Pawlowsky-Glahn et al. 2015; Pawlowsky-  
55 Glahn and Olea 2004; Tolosana-Delgado and van den Boogaart 2013; van den Boogaart and  
56 Tolosana-Delgado 2013). Most of the multivariate geostatistical simulation techniques are  
57 based on the assumption of multivariate multigaussianity of the data. In real case studies, log-  
58 ratio transformed data alone do not ensure this assumption, so the log-ratio transform has to be  
59 combined with a normal score transform prior to using geostatistical simulation techniques in  
60 order to not violate the assumptions of multigaussianity (Chilès and Delfiner 2012; Mueller et  
61 al. 2014). However, compositional data do not have a unique, canonical representation and  
62 several log-ratio transformations are available, making invariance of the simulated results  
63 under the choice of log-ratio transform desirable. Normal score transformations based on  
64 quantile matching do not have the invariance property and in addition the transformed data  
65 might not be multivariate normal. Flow anamorphosis (FA) has been designed to address these  
66 challenges, which means, it is capable of transforming original multivariate data to multivariate  
67 normal space and at the same time being invariant under the choice of log-ratio transform  
68 (Mueller et al. 2017; van den Boogaart et al. 2017).

69 On the other hand, the heterogeneity of geological units in an ore deposit requires defining  
70 domains that differentiate various characteristics such as: types of mineralogy, alteration,  
71 lithology, and microstructures. Indeed, these geological domains control the ore characteristics.  
72 One can think of deposits in which the mean grades and patterns of spatial continuity depend  
73 upon the rock and/or alteration type. For example a porphyry copper deposit with high grades  
74 in potassic alterations surrounded by low grades in argillic and propylitic alterations (Talebi et  
75 al. 2013), presence of lately injected barren dykes of different sizes and orientations (Talebi et  
76 al. 2014), and spatial modelling of geological units in an uranium roll-front deposit (Renard  
77 and Beucher 2012). Currently, the most prevalent approach to model the uncertainty in the  
78 spatial distribution of the elements of interest is to divide the study area into subdomains  
79 (geological units) based on geological interpretation and to predict the variables of interest  
80 within each domain separately. This approach defines just one interpretation of the geological  
81 domains and does not offer any measure of the uncertainty in the position of the domain  
82 boundaries. This uncertainty can be evaluated by use of geostatistical simulation methods for  
83 categorical variables (Alabert 1987b; Armstrong et al. 2011; Mariethoz and Caers 2015).

84 The objective of this contribution is to compare different approaches for geostatistical  
85 simulation of geochemical compositions to assess mineral resources in a nickel-cobalt laterite  
86 deposit. To evaluate the effect of geological domaining on the accuracy of the predicted  
87 geochemical compositions, three approaches are analysed: geological controls are ignored, a  
88 deterministic geological model is applied, and a probabilistic geological model is used (Talebi  
89 et al. 2016). The probabilistic geological model is calculated based on a plurigaussian (PGS)  
90 model (Armstrong et al. 2011; Emery 2007). An isometric log-ratio transformation (ilr) is used  
91 to transform compositional data from the simplex to real space (Egozcue et al. 2003). The ilr-  
92 transformed data are transformed to normal scores and subsequently simulation is used to  
93 generate realisations at unsampled locations. Classical gaussian anamorphosis (GA) and flow

94 anamorphosis are compared based on the accuracy of the predicted compositions, as well as  
95 their capability to reproduce complex statistical and spatial patterns present in the input data.  
96 The various approaches considered are illustrated through an application to a nickel-cobalt  
97 laterite deposit and their performances are evaluated against a set of validation boreholes.

98 The paper is organised as follows: In Sect 2 the basics of the compositional data analysis are  
99 covered. Various log-ratio transformations used in this paper and transformation to multivariate  
100 normal space via flow anamorphosis are presented in this section. A new method for adjusting  
101 global proportion of geological domains is proposed in Sect 2.3. Various approaches for  
102 geostatistical simulation of regionalised compositions in the presence of several geological  
103 domains are presented in Sect 2.4. In Sect 3 the case study (Murrin Murrin nickel-cobalt laterite  
104 deposit) is introduced and the dataset is presented. A compositional contact analysis is  
105 implemented in Sect 3.3. Section 3.4 presents the process of generating deterministic and  
106 probabilistic geological models via a plurigaussian simulation approach. In Sect 4 the results  
107 are presented and proposed methods are compared to each other based on several criteria.  
108 Finally, some conclusions and final thoughts are presented in Sect 5.

109

## 110 **2 Methodology**

### 111 **2.1 Compositional Data Analysis**

112 Compositional data are multivariate data where the non-negative components are measured on  
113 the same scale and are constrained by a constant sum property, usually 100%. Geochemical  
114 and mineral compositions and proportions of various rock types or alteration types in a block  
115 are typical examples of compositional data in an ore deposit. The compositional space is a D-  
116 dimensional simplex

$$S^D = \left\{ \vec{Z}(u_\alpha) = [z_1(u_\alpha), z_2(u_\alpha), \dots, z_D(u_\alpha)] \mid z_i(u_\alpha) \geq 0; i = 1, 2, \dots, D; u_\alpha \in \mathbf{A}; \sum_{i=1}^D z_i(u_\alpha) = m \right\}, \quad (1)$$

117

118 where  $z_i(u_\alpha)$  represents the  $i^{th}$  component measured at location  $u_\alpha$  within the study area  $\mathbf{A}$ .  
 119 The number  $m$  is the constant sum and common values are 1 (proportions), 100 (percentages),  
 120  $10^6$  (ppm), and  $10^9$  (ppb). The constant sum constraint is known to induce problems of spurious  
 121 correlation (see Aitchison (1982), for a detailed report). Compositional data carry just relative  
 122 information, which is appropriately represented by taking log-ratio transformations (Aitchison,  
 123 1986). Pairwise log-ratio transformation (Aitchison 1986), centred log-ratio transformation  
 124 (Aitchison 1986), and isometric log-ratio transformation (Egozcue et al. 2003) are utilized in  
 125 this study. Independently of which transform used, the resulting log-ratio scores happen to be  
 126 free of the constraints of positivity and constant sum or of the spurious correlation problem,  
 127 which make log-ratio scores more amenable to (geo)statistical treatment. The pairwise log-  
 128 ratio transformation (pwlr) is defined as follows

129

$$pwlr(\vec{Z}(u_\alpha)) = \begin{bmatrix} 0 & \ln\left(\frac{z_1(u_\alpha)}{z_2(u_\alpha)}\right) & \dots & \ln\left(\frac{z_1(u_\alpha)}{z_D(u_\alpha)}\right) \\ \ln\left(\frac{z_2(u_\alpha)}{z_1(u_\alpha)}\right) & 0 & \dots & \ln\left(\frac{z_2(u_\alpha)}{z_D(u_\alpha)}\right) \\ \vdots & \vdots & \ddots & \vdots \\ \ln\left(\frac{z_D(u_\alpha)}{z_1(u_\alpha)}\right) & \ln\left(\frac{z_D(u_\alpha)}{z_2(u_\alpha)}\right) & \dots & 0 \end{bmatrix}. \quad (2)$$

130

131

132 The centred log-ratio transformation (clr), is calculated via the following formula

$$clr(\vec{Z}(u_\alpha)) = \ln\left(\frac{\vec{Z}(u_\alpha)}{\sqrt[ D]{\prod_{i=1}^D z_i(u_\alpha)}}\right). \quad (3)$$

133

134

135 Finally, ilr transformation is defined as follows

136

$$ilr(\vec{Z}(u_\alpha)) = V \cdot clr(\vec{Z}(u_\alpha)), \quad (4)$$

137

138

139 where  $V$  is a  $(D - 1) \times D$  matrix whose columns are pairwise orthogonal vectors and sums to  
140 zero. Each matrix  $V$  satisfying these conditions give rise to an ilr transformation. There are  
141 infinitely many ilr transformations and often it is recommended to select one that increases the  
142 interpretability of the ilr scores. However, the aim of this study is to predict the geochemical  
143 compositions and model the associated uncertainty at unsampled locations accurately. Hence,  
144 a default orthonormal basis is used (Egozcue et al. 2003). The predicted compositions do not  
145 depend on the actual log-ratio transformation implemented for the computations (Tolosana-  
146 Delgado 2006).

147

## 148 **2.2 Flow Anamorphosis**

149 The flow anamorphosis introduced by van den Boogaart et al. (2017) is a multivariate form of  
150 gaussian anamorphosis. In this method a kernel density estimate is deformed from the given  
151 multivariate density of the observations into the density of a standard multivariate normal  
152 distribution. Several statistical tests of multivariate normality can be applied to check the fit of  
153 the transformed data to normality (Korkmaz et al. 2014; Mardia 1970; Székely and Rizzo 2005;  
154 Székely and Rizzo 2013). The fit is highly dependent on the selection of the two parameters  
155 of FA,  $\sigma_0$  and  $\sigma_1$  (initial and final spreads of the smoothing kernels of the kernel density

156 estimates). Deformation of the underlying space is controlled by  $\sigma_0$ . The smaller the value, the  
157 closer the transformed data are to multivariate normality. Selecting a proper  $\sigma_0$  is dependent on  
158 the number of variables ( $D$ ), sample size, and complexity of the input data (Mueller et al. 2017).  
159 On the other hand,  $\sigma_1$  controls the ranges of the transformed distributions. In this study, in order  
160 to force the marginal distributions of transformed data to have standard deviations close to  
161 unity,  $\sigma_1$  is selected as  $\sigma_1 = \sigma_0 + 1$  (Mueller et al. 2017). Figure 1 shows the process of  
162 simulation via FA.

163 [\[Fig. 1 about here.\]](#)

164  
165 Figure 1(a) depicts the scatter plot and marginal kernel smoothing density estimates of the input  
166 data with two simulated variables whose relationship is complex. The characteristics include  
167 the presence of some outliers, nonlinear relationships, several populations, and  
168 heteroscedasticity. Figure 1(b) and Fig. 1(c) show the co-deformation of the underlying space  
169 and final distribution of the transformed data respectively. The transformed data in multivariate  
170 normal space can be simulated (Fig. 1(d)) via many geostatistical algorithms (Alabert 1987a;  
171 Deutsch and Journel 1998; Emery 2008; Emery et al. 2016; Emery and Lantuéjoul 2006).  
172 Several experiences have shown that the FA-transformed data are not only multivariate normal  
173 but often also exhibiting lack of spatial cross-correlation which make the geostatistical  
174 simulation of such orthogonal factors, straightforward (Mueller et al. 2017; van den Boogaart  
175 et al. 2017). Otherwise, multivariate simulation or spatial decorrelation technique followed by  
176 univariate simulation could be implemented. The simulated results are back-transformed to the  
177 original space via  $FA^{-1}$ . Figure 1(e) and Fig. 1(f) show the co-deformation back from simulated  
178 multivariate normal space to the original space and final distribution of the simulated data  
179 respectively. A visual comparison of these two plots shows that all the characteristics of the  
180 input the data are very well reproduced.

181

## 182 **2.3 Geological Domaining**

183 By constructing multiple spatial realisations of the geological domains, geostatistical  
184 simulation helps to improve the geological interpretation and to measure associated  
185 uncertainty. Several geostatistical methods for categorical variables can be used to this end. In  
186 the field of two-point geostatistics, PGS has gained popularity and proved to be suitable for  
187 reproducing complex configurations of geological domains without the need to define a  
188 training image. After simulating  $K$  geological domains at each location  $u_\alpha$  in the study area  $\mathbf{A}$ ,  
189 local proportions of the simulated domains,  $(q_1(u_\alpha), q_2(u_\alpha), \dots, q_K(u_\alpha))$ , can be calculated  
190 from the realisations. By assigning the geological domain  $k$ , ( $k = 1, 2, \dots, K$ ), with the highest  
191 proportion to each location  $u_\alpha$  the most probable map is obtained. In this study,  $K$  spatial  
192 proportion maps (associated with  $K$  geological domains) and the most probable map are used  
193 as probabilistic and deterministic geological models respectively. It is known that the global  
194 proportions  $\vec{Q} = (q_1, q_2, \dots, q_K)$  of simulated domains in the most probable map might be  
195 different from the global proportions  $\vec{P} = (p_1, p_2, \dots, p_K)$  of the domains in the input data.  
196 Some economically important domains might be under-represented. To reduce under-  
197 representation of geological domains in the simulated geological model, it is common to post-  
198 process the proportions. One such method is the Soares correction (Soares 1998) which restores  
199 the global proportions of the input data, but is known to lead to artefacts and does not take  
200 account of the location of the datum to be adjusted. The correction method proposed here is  
201 also based on the proportions of domains in the input data, but takes location into account by  
202 borrowing a technique from compositional data analysis. The local proportions at each location  
203  $u_\alpha$  in the simulated geological model are perturbed by putting

204

$$\vec{r}(u_\alpha) = (r_1(u_\alpha), r_2(u_\alpha), \dots, r_K(u_\alpha)) = \left( \frac{b_1 q_1(u_\alpha)}{R(u_\alpha)}, \frac{b_2 q_2(u_\alpha)}{R(u_\alpha)}, \dots, \frac{b_K q_K(u_\alpha)}{R(u_\alpha)} \right). \quad (5)$$

205

206 Here  $\vec{r}(u_\alpha)$  is the vector of perturbed proportions at location  $u_\alpha$ ,  $R(u_\alpha) = \sum_{k=1}^K b_k q_k(u_\alpha)$ ,  
 207 and finally  $(b_1, b_2, \dots, b_K) = \left( \frac{p_1}{q_1}, \frac{p_2}{q_2}, \dots, \frac{p_K}{q_K} \right)$ . The resulting new local probabilities  
 208  $(r_1(u_\alpha), r_2(u_\alpha), \dots, r_K(u_\alpha))$ , form the new probabilistic model of geology and can be used to  
 209 determine the adjusted most probable map of geological domains.

210

#### 211 **2.4 Approaches to Geostatistical Simulation of Compositional Data**

212 In this study several scenarios are investigated to assess the effects of geological models and  
 213 transformation to normal space on the geostatistical simulation of geochemical compositions.  
 214 Selected approaches to incorporate geological information are as follows: prediction without  
 215 geological control, prediction by using a deterministic geological model, and prediction by  
 216 using a probabilistic geological model. GA (Wackernagel 2003) and FA algorithms are  
 217 compared to assess the effect of transformation to normal space and subsequent back-  
 218 transformation on the spatial simulation of regionalised compositions.

219 In the first ( $\mathbf{M}_0$ ) and second ( $\mathbf{M}_1$ ) proposed methods there is no geological control (Table 1).  
 220 In these two scenarios all input compositions (without considering a geological domain  
 221 partition of the deposit) are transformed to real space via an ilr transformation and subsequently  
 222 to normal space via GA and FA, respectively. If the normal scores are spatially correlated, they  
 223 are transformed to spatial orthogonal factors via Min/Max autocorrelation factors (MAF),  
 224 (Bandarian et al. 2008; Desbarats and Dimitrakopoulos 2000; Rondon 2012; Switzer and Green  
 225 1984). This orthogonalization makes the simulation step more straightforward. However,  
 226 normal scores obtained via FA are normally spatially orthogonal and independent (Mueller et

227 al. 2017; van den Boogaart et al. 2017). The normal scores (or MAFs) are simulated  
228 independently at the simulation grid (in this case all the locations of the validation data) via  
229 turning bands (TB) technique (Emery and Lantuéjoul 2006). The simulated results are back-  
230 transformed to the original space afterwards.

231 In the simulation via a deterministic geological model ( $\mathbf{M}_2$ ) input compositions are divided into  
232 several subsets based on their associated geological domains. Compositions in each subset are  
233 transformed to ilr space and subsequently transformed to multivariate normal space via FA.  
234 The simulation grid is also divided into mutually exclusive and exhaustive domains based on  
235 the most probable rock types (deterministic geological model) achieved by a PGS model.  
236 Normal scores from each subset are simulated at the associated part of the simulation grid.  
237 Finally simulated results at each part of the simulation grid are back-transformed to the original  
238 space independently.

239 The last proposed method ( $\mathbf{M}_3$ ) is geostatistical simulation of geochemical compositions using  
240 a probabilistic geological model. In this case input compositions are divided into subsets based  
241 on their associated geological domain. Compositions in each subset are transformed to ilr space  
242 and subsequently transformed to multivariate normal space via FA. Normal scores from each  
243 subset are simulated on the entire simulation grid and back-transformed to the original space  
244 independently. This process provides several sets of simulated geochemical compositions  
245 associated with the geological domains. Final simulated compositions can be obtained via  
246 weighting the simulated compositions associated with the different domains by the local  
247 probabilities of occurrence of each domain (Emery and González 2007a; Emery and González  
248 2007b; Talebi et al. 2015)

$$\vec{Z}(u_\alpha) = \sum_{k=1}^K q_k(u_\alpha) \vec{Z}^k(u_\alpha), \quad (6)$$

249

250 where  $K$  is the number of classes (geological domains),  $q_k(u_\alpha)$  is the probability of geological  
251 domain  $k$  at the location  $u_\alpha$  calculated by a PGS model, and  $\vec{Z}^k$  is the simulated composition  
252 associated with geological domain  $k$ . The final simulated vector  $\vec{Z}(u_\alpha)$  is still a composition  
253 because

$$\sum_{i=1}^D z_i(u_\alpha) = \sum_{i=1}^D \sum_{k=1}^K q_k(u_\alpha) z_i^k(u_\alpha) = \sum_{k=1}^K q_k(u_\alpha) \sum_{i=1}^D z_i^k(u_\alpha) = \sum_{k=1}^K q_k(u_\alpha) m = m. \quad (7)$$

254

255 Finally methods based on a deterministic model of geology ( $\mathbf{M}_2$ ) and a probabilistic model  
256 ( $\mathbf{M}_3$ ) are repeated with the corrected deterministic ( $\mathbf{M}_{2c}$ ) and probabilistic ( $\mathbf{M}_{3c}$ ) models. The  
257 correction is based on the method described in Sect 2.3. Table 1 provides a summary of the  
258 proposed methods.

259

[\[Table 1 about here.\]](#)

260

261

262

### 263 **3 Case Study: Murrin Murrin Nickel-Cobalt Laterite Deposit**

264 Murrin Murrin East (MME) is a nickel-cobalt laterite deposit located in Western Australia, at  
265 about 60 km southeast of Laverton and at an average elevation of approximately 420 metres  
266 above mean sea level. The orebody is approximately 1,500 meters long, 600 meters wide, and  
267 30 meters thick. Figure 2 shows a satellite image of the orebody together with the location of  
268 the boreholes.

269

270

[\[Fig. 2 about here.\]](#)

271

272

### 273 **3.1 Geological Description**

274 Laterite deposits are formed during chemical weathering of ultramafic rocks near the surface  
275 of the earth. At MME, nickel laterite deposits occur as laterally extensive, undulating blankets  
276 of mineralisation with strong vertical trends covering basement ultramafic rocks (Murphy  
277 2003). Proximity of the deposit to a major salt lake probably influenced the development of the  
278 regolith (Markwell 2001). Based on the geochemical interpretation and the logging information  
279 from geologists, lateritic weathering of the ultramafic rocks at MME has produced a profile  
280 that may be broadly divided into four geological units (Camuti and Riel 1996; Markwell 2001;  
281 Monti and Fazakerley 1996). The sequence of these units from the bottom to the top is as  
282 follows: (i) ultramafic rocks (UM) at the base of weathering overlain by (ii) a Saprolite zone  
283 (SA) overlain by (iii) a Smectite zone (SM) and finally capped by (iv) a Ferruginous zone (FZ,  
284 Fig. 3(b)). **UM** occurs as a thin layer of unweathered ultramafic rock at the base of the deposit.  
285 Due to the undulating nature of different layers in this deposit, some outcrops of UM can be  
286 recognized at the surface of the deposit. **SA** consists mainly of Lizardite and Smectite. SA is  
287 enriched in Mg, but low amounts of Fe are present in this zone. A strong contrast between Mg  
288 in SM and SA allows easier domaining of the two units (Markwell 2001). **SM** consists mainly  
289 of Smectite and is confined to the shoulders of Saprolite domes. SM is enriched in Ni and Co  
290 (Fig. 3) and depleted in Mg. **FZ** is composed predominantly of Goethite, Kaolin, and  
291 Maghemite. FZ is less enriched in Ni, but it can host significant amounts of Co, especially at  
292 the transition to the Smectite zone (Fig. 3).

293 Table 2 shows the mineral assemblage in each geological unit at MME, obtained from several  
294 representative samples (Markwell 2001).

295 [\[Table 2 about here.\]](#)

296

297

### 298 **3.2 Dataset**

299 In total, 17,512 samples (of 1 meter length) from 926 RC holes (Fig. 2) make up the database  
300 for this study. Four rock types (FZ, SA, SM, and UM) are considered to define compositionally  
301 homogeneous domains. Three major (Fe, Al, and Mg) and two target (Ni and Co) elements are  
302 the variables of interest in this study. Since the data are compositional, a filler variable is  
303 introduced to achieve closure and to retain the intuitive relationship between each component  
304 and the mass of its associated element. The data set was subdivided into two subsets: 453 holes  
305 (8,694 samples) are considered for validation and 473 holes (8,818 samples) for geostatistical  
306 modelling, called input data from hereon. The input data are used for constructing the various  
307 geostatistical models. Locations of the validation and input boreholes can be seen in a cross  
308 section of the deposit for northing 300m (Fig. 3(a)). Figure 4 depicts the histograms of different  
309 geochemical components in the validation and input sets. Figure 5 shows centred ternary  
310 diagrams (Buccianti et al. 1999; Pawlowsky-Glahn et al. 2015) of the sub-compositions. In this  
311 case study geochemical compositions are composed of different populations. The multi-  
312 population character is consistent with the presence of four geological units (Fig. 4 and Fig. 5).  
313 These geological units control the statistical and spatial distribution of the geochemical  
314 compositions. Figure 6 shows the vertical curves of the rock type proportions and clr-  
315 transformed of the geochemical components.

316 There is a zoned weathering profile (Fig. 6), in which Mg as a mobile element effectively has  
317 been removed while Fe and Al, as less mobile elements, have been enriched residually in the  
318 Ferruginous zone. On the other hand, Co (one of the value elements), has been accumulated at  
319 the transition between Ferruginous and Saprolite zones (Talebi et al. 2017) where Smectite  
320 units are mainly located. Finally, the Smectite and Saprolite zones between fresh (ultramafic  
321 parent rock) and weathered zones (Ferruginous zones), have been enriched in Ni (the other  
322 value element). Figure 7 shows the scatterplots of the clr-transformed components. Complex  
323 relationships (nonlinearity, multi-population, and presence of outliers) can be recognised  
324 between different components. Based on the vertical curves (Fig. 6) and the scatterplots of the  
325 clr-transformed components (Fig. 7), Al-Fe, Ni-Co, and Mg-Ni are positively correlated. On  
326 the other hand, Al-Mg, Al-Ni, Co-Filler, Fe-Mg, Ni-Filler, are in terms of their clr's negatively  
327 correlated. From Fig. 4 to Fig. 7, it is clear that the validation and input sets are statistically  
328 and spatially similar.

329

330 [\[Fig. 3 about here.\]](#)

331 [\[Fig. 4 about here.\]](#)

332 [\[Fig. 5 about here.\]](#)

333 [\[Fig. 6 about here.\]](#)

334 [\[Fig. 7 about here.\]](#)

335

336

### 337 3.3 Compositional Contact Analysis

338 In order to evaluate the variation of the variables of interest at the transition zone between two  
339 geological domains, a contact analysis was conducted. To do this, statistics of the variables of  
340 interest such as means and standard deviations are plotted as functions of distance from the  
341 contact zone (Ortiz and Emery 2006). However, compositional data have spurious correlations  
342 and real variation in one component might cause apparent variations in others, so contact  
343 analysis of raw components should be avoided (Tolosana-Delgado et al. 2016). Although log-  
344 ratio transformed data can be treated as real data and compositional contact analysis can be  
345 implemented on the clr-transformed data, contact analysis of the pwlr-transformed data  
346 provides the geologists with an enriched view of the variations at the contact zone. Figure 8  
347 shows the compositional contact analysis between the two dominant rock types, FZ and SA.  
348 Diagrams in the last row and last column show means and standard deviations as functions of  
349 the distance from the contact zone for the clr-transformed data and raw components  
350 respectively while the remaining diagrams show those of the pwlr-transformed data  
351 (row/column).

352

353

[\[Fig. 8 about here.\]](#)

354

355

356 A comparison of the contact diagrams of the raw components with the associated clr-  
357 transformed ones shows that the behaviour of the means at the transition zone has been  
358 exaggerated unrealistically in the raw contact analysis. Except for Filler which shows different  
359 behaviour in the clr plot, other components show similar behaviour in raw and clr analysis. Al,  
360 Fe, and Mg show abrupt transitions (variation of means as a function of distance) while the

361 local variations are stable (standard deviations as a function of distance). All the diagrams of  
362 the Co (raw, clr, and pwlr) show an increase of the mean and the standard deviation  
363 (proportionality effect) at the transition zone. The reason for this behaviour would be the  
364 presence of Smectite units at the transition from Ferruginous to Saprolite zone, not recognised  
365 during logging (Smectite units are enriched in Co). Pairwise log-ratio of the Co/Ni and Al/Fe  
366 means and standard deviations are stable across geological units. The results suggest that these  
367 two geological domains (FZ and SA) can be considered as stationary for some subcompositions  
368 such as Al and Fe, but domaining would be necessary for other subcompositions or ratios, for  
369 example Mg and Ni. Similar assessments were conducted on the other contacts between the  
370 four geological domains and results supported the necessity of domaining. Given the results of  
371 compositional contact analysis and statistical analysis of compositional data in Sect 3.2, it is  
372 reasonable to partition the study area to four homogeneous geological units (four rock types)  
373 prior to geostatistical modelling.

374

### 375 **3.4 Deterministic and Probabilistic Geological Models**

376 The geological domains (rock types) for the validation data were simulated via a plurigaussian  
377 model (Armstrong et al. 2011). Overall 100 realisations were generated and used to determine  
378 the probability of occurrence for each rock type (Fig. 9(b) to Fig. 9(e)) and the most probable  
379 rock type (Fig. 9(f)). Proportions of the rock types in the most probable map (Table 3) show  
380 that, under-representation of the SM has occurred. Since SM units are highly mineralised in  
381 Co, prediction of these units with high accuracy is of great importance. The method described  
382 in Sect 2.3 was used to adjust the proportions of geological domains in the simulated models.

383 The last column in Table 3 shows the adjusted proportions in the most probable rock type map,  
384 demonstrating an improvement in the proportion of the SM units. Figure 9(g) to Fig. 9(j) show

385 cross-sections of the validation boreholes, coloured by the adjusted probabilities, for northing  
386 300m. A visual comparison of Fig. 9(c) and Fig. 9(h) indicates a clear improvement in the  
387 proportion of the SM units, more representative of reality (Fig. 9(a)).

388

389 [\[Table 3 about here.\]](#)

390

391

392 [\[Fig. 9 about here.\]](#)

393

394

395

## 396 **4 Results and Discussion**

397 Figure 10 shows the histograms and scatterplots (coloured by kernel density estimate) of the  
398 ilr-transformed input data. As the ilr-transformed data (in this study) are not multivariate  
399 normal (Fig. 10), a transformation to normal space is needed prior to geostatistical simulation.

400 Transformation of ilr transformed data into normal space in  $\mathbf{M}_0$  was based on GA. Figure 11  
401 shows the scatterplots of the normal scores obtained by GA. Although the marginal  
402 distributions are normal, the scores are not multivariate normal. An analysis of the  
403 omnidirectional experimental semivariograms and cross-variograms further revealed that these  
404 normal scores are spatially correlated, with Tercan's  $\bar{\tau}$  and  $\bar{\kappa}$  (Tercan 1999) equal to 0.1973  
405 and 0.8147, respectively. To make the simulation step more straightforward, normal scores  
406 data were transformed to spatially orthogonal factors via MAF. MAF transformation improved  
407 spatial orthogonality with  $\bar{\tau}$  and  $\bar{\kappa}$  equal to 0.0782 and 0.9689, respectively. Orthogonal factors

408 were simulated at the entire validation holes via the TB algorithm. The simulated results were  
409 back-transformed to the simplex afterwards to recover outputs in the original scale as  
410 percentages.

411 On the other hand,  $\mathbf{M}_1$  transforms the ilr-transformed data to multivariate normal space via FA.  
412 Due to the complexity of the data and the number of the observations and variables,  
413 multivariate normality was not achieved by a single FA. Two successive FA with the same  
414 parameters ( $\sigma_0 = 0.1$  and  $\sigma_1 = 1.1$ ) were required to achieve multivariate normality. Figure 12  
415 shows the normal scores are obviously close to multivariate normal and uncorrelated, so  
416 statistically independent. Spatial structural analysis (variography) showed further that the  
417 normal scores are spatially orthogonal, with Tercan's  $\bar{\tau}$  and  $\bar{\kappa}$  equal to 0.0656 and 0.9873,  
418 respectively, so they could be simulated independently. The scores were simulated at the  
419 validation holes independently via TB algorithm and back-transformed to composition  
420 afterward.

421 [\[Fig. 10 about here.\]](#)

422 [\[Fig. 11 about here.\]](#)

423 [\[Fig. 12 about here.\]](#)

424

425

426 For simulation based on a deterministic geological model input compositions were divided into  
427 four subsets based on their associated rock types (FZ, SM, SA, and UM). Compositions in each  
428 subset were transformed to ilr space and subsequently transformed to multivariate normal space  
429 via FA independently. The validation holes were also divided into four mutually exclusive and  
430 exhaustive parts based on the most probable rock types (Fig. 9(f)) and the corrected most  
431 probable rock types (Fig. 9(k)), provided the simulation grids for  $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$  respectively.

432 Normal scores from each subset (for example, flow anamorphosed FZ data) were simulated at  
433 the associated part of the simulation grid (for example, locations in the validation holes with  
434 the most probable rock known as FZ). Finally simulated results for each domain were back-  
435 transformed to the original space independently.

436 Unlike the deterministic approaches ( $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$ ), in the probabilistic approach normal scores  
437 from each subset of input compositions were simulated at the entire simulation grid. This  
438 process provided four sets of simulated geochemical compositions associated with the four  
439 geological domains (FZ, SM, SA, UM). Final simulated compositions obtained by weighting  
440 the simulated compositions associated with the different domains by the probabilities of  
441 occurrence of each domain (Eq. 6). Two sets of probabilities are available, the raw probabilities  
442 (Fig. 9(b) to Fig. 9(e)) obtained by the PGS model ( $\mathbf{M}_3$ ) and adjusted probabilities ( $\mathbf{M}_{3c}$ )  
443 obtained by the proposed correction method in Sect 2.3 (Fig. 9(g) to Fig. 9(j)).

444 The simulated compositional models based on the six proposed approaches were validated  
445 against reality available at the validation boreholes (Fig. 3(a)). Figure 13 shows the global  
446 histogram reproduction of the six methods. Best realisations were achieved by  $\mathbf{M}_1$  followed by  
447  $\mathbf{M}_{2c}$ . Visual comparison of histogram reproduction of  $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$  reveals that adjusting  
448 proportions (Sect 2.3) has improved the histogram reproduction especially for Co component.  
449 Worst reproductions were achieved by  $\mathbf{M}_0$ .

450 [\[Fig. 13 about here.\]](#)

451

452

453 To check the ability of the methods to reproduce the complex patterns in the simplex space,  
454 ternary diagrams were plotted and compared to each other (Fig. 14). Due to the averaging  
455 nature of  $\mathbf{M}_3$  and  $\mathbf{M}_{3c}$ , different populations have been mixed up, which is not acceptable. On

456 the other hand,  $\mathbf{M}_0$  was not able to reproduce different populations and complex patterns inside  
457 the simplex space. Here,  $\mathbf{M}_1$  generated the best results followed by  $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$ .

458 The smoothing effect of the  $\mathbf{M}_3$  and  $\mathbf{M}_{3c}$  is more obvious from the experimental variograms of  
459 the simulated models (Fig. 15). In Fig. 15 continuous black lines are input data, dashed black  
460 lines are validation data and grey lines are realisations. The sills of the variograms for one of  
461 the simulated component (Ni) are systematically less than the sills for input and validation data,  
462 representing a systematic reduction of the spatial variability. Adjusting probabilities of  
463 geological domains has improved the variogram reproduction for the deterministic approaches  
464 (compare  $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$ ).

465 Risk quantification in grade-tonnage curves (GTC) and predicting true curves are critical for  
466 feasibility studies and capital investment in mining projects. The accuracy of the six methods  
467 was investigated based on their capability for reproducing input GTCs and predicting true  
468 (validation data in this study) GTCs. Figure 16 and Fig. 17 show the GTCs for Ni and Co  
469 components respectively. The smoothing effect of the  $\mathbf{M}_3$  and  $\mathbf{M}_{3c}$  is again clear from the GTCs  
470 of both Ni and Co. Comparing GTCs of  $\mathbf{M}_2$  and  $\mathbf{M}_{2c}$ , an improvement can be recognised due  
471 to the implementation of the proposed technique for adjusting underrepresented domains.  
472 GTCs for Co component generated by  $\mathbf{M}_0$  is not acceptable.  $\mathbf{M}_1$  generated the most satisfactory  
473 results followed by  $\mathbf{M}_{2c}$ .

474 Finally, the six proposed methods were compared to check the presence of any systematic bias  
475 in the predicted proportions of samples above cut-offs (Fig. 18). In multi-element deposits,  
476 there are more than one target elements (in this study two target elements: Ni and Co). In Fig.  
477 18, each cell of the maps is associated with two cut-offs (one for Ni and one for Co). For each  
478 cell the expected proportion of samples above the two cut-offs were calculated from all  
479 realisations and subsequently subtracted from the real associated proportion calculated from

480 validation data. These maps can be compared to show the capability of the proposed methods  
481 for avoiding any serious bias (over or underestimations of ore tonnages).  $M_0$ ,  $M_3$ , and  $M_{3c}$   
482 show clear systematic over and underestimations in some range of cut-offs. In this regard,  $M_1$   
483 generated the best (least bias) results, followed by  $M_{2c}$ .

484 According to the aforementioned criteria,  $M_1$  outperformed other proposed techniques for  
485 geostatistical simulation of geochemical compositions. One reason is that  $M_1$  considers spatial  
486 correlations of geochemical components across geological boundaries. More important reason  
487 is the ability of geostatistical simulation via FA to reproduce multi-population characteristic of  
488 the input data which is consistent with the presence of several geological units.

489

490 [\[Fig. 14 about here.\]](#)

491 [\[Fig. 15 about here.\]](#)

492 [\[Fig. 16 about here.\]](#)

493 [\[Fig. 17 about here.\]](#)

494 [\[Fig. 18 about here.\]](#)

495

496

## 497 **5 Conclusion**

498 This paper compares various geostatistical approaches for simulation of geochemical  
499 compositions and their application to mineral resource evaluation. Results and several  
500 validation tests showed that the classical transformation to normal space (gaussian  
501 anamorphosis) is not capable of reproducing complex statistical patterns inside data and should  
502 be replaced with more advanced transformations. The method for modelling geochemical

503 compositions based on a probabilistic geological model exhibits smoothing effects due the  
504 averaging nature of the algorithm. Although this method generates satisfactory results for  
505 kriged maps (Emery and González 2007a; Emery and González 2007b; Talebi et al. 2015), it  
506 should be avoided for simulation purposes. The proposed technique for adjusting  
507 underrepresented domains improved the result of simulation and should be used in the cases  
508 where there are important geological domains with small proportions such as SM in this case  
509 study. Flow anamorphosis is a vital element for geostatistical modelling of geochemical  
510 composition due to its invariance properties and capability for reproducing complex patterns  
511 in data such as: outliers, multi-population, nonlinearity, and heteroscedasticity. In the case  
512 study presented, a simulation involving a global flow anamorphosis without domaining was  
513 best capable of reproducing all performance targets (histograms, variograms, grade and  
514 tonnage curves). In the authors' opinion, this remarkable property might occur again in other  
515 settings in which domains emerge as the effect of chemical processes mostly involving the  
516 composition modelled.

517

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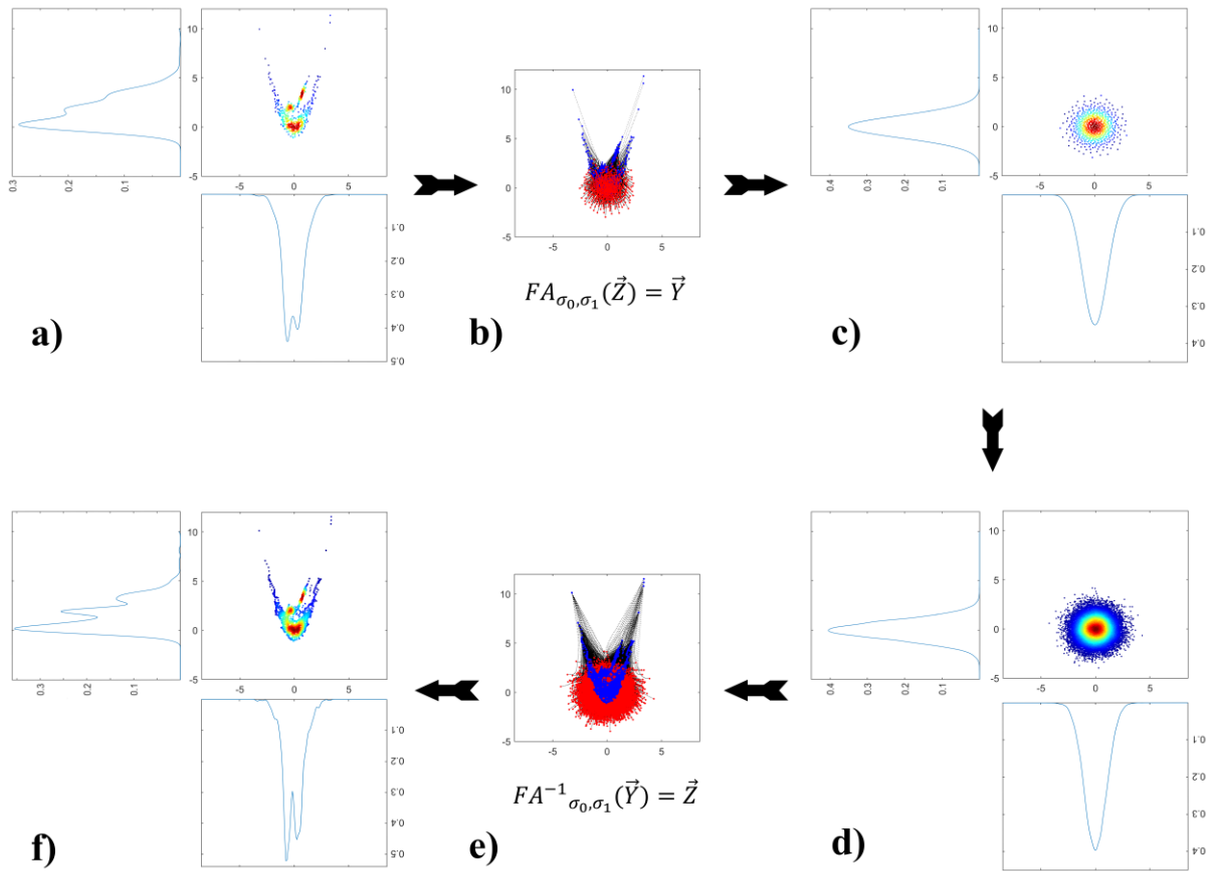
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**Fig. 1** Geostatistical simulation via Flow Anamorphosis

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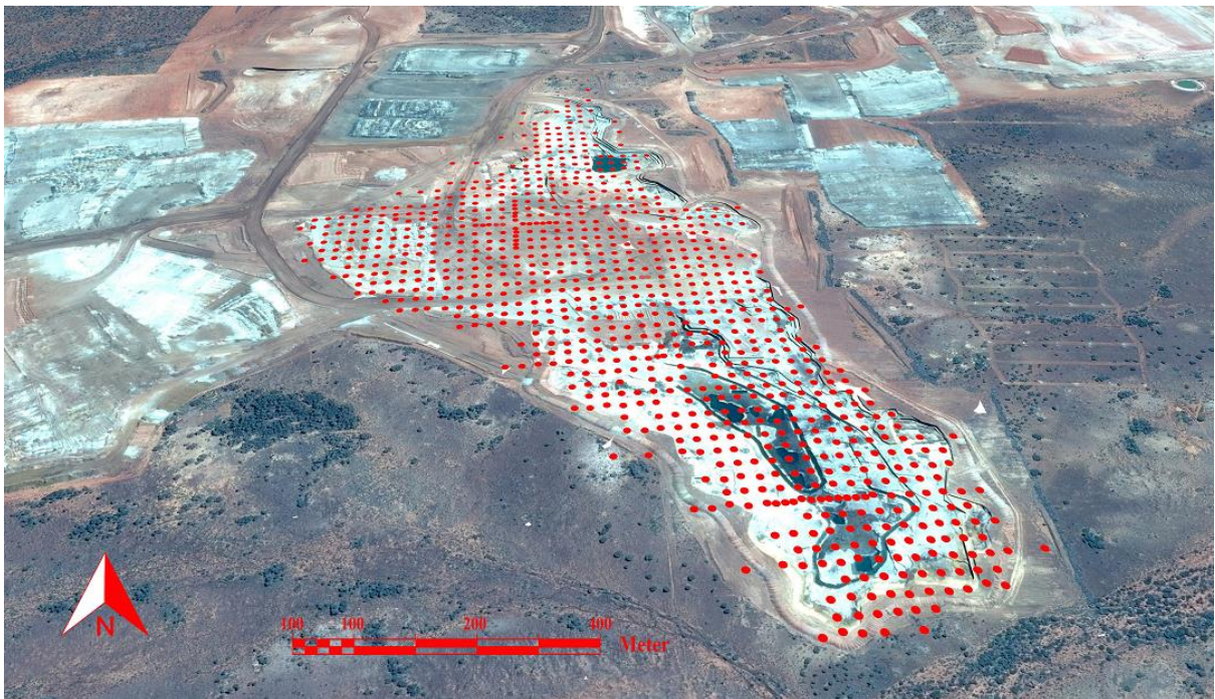
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**Fig. 2** Borehole location map of the Murrin Murrin East (MME)

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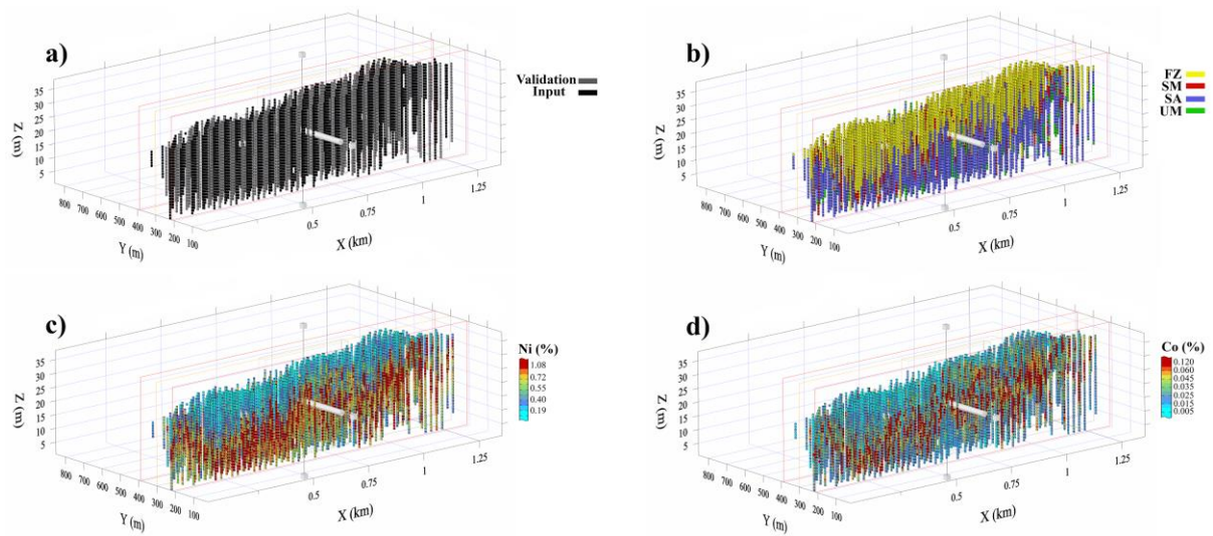
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761 **Fig. 3** Cross sections of boreholes for northing 300m and 100m thickness: locations of input and validation  
762 boreholes (a), spatial distributions of different rock types (b) Ni grade (c) and Co grade (d)

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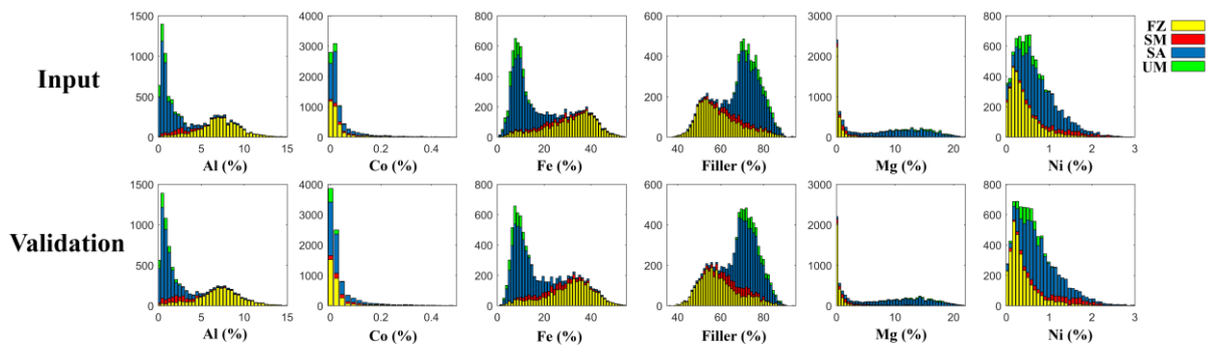
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772 **Fig. 4** Histogram of the geochemical components of input and validation sets, coloured by the proportion of rock  
773 types in each bin

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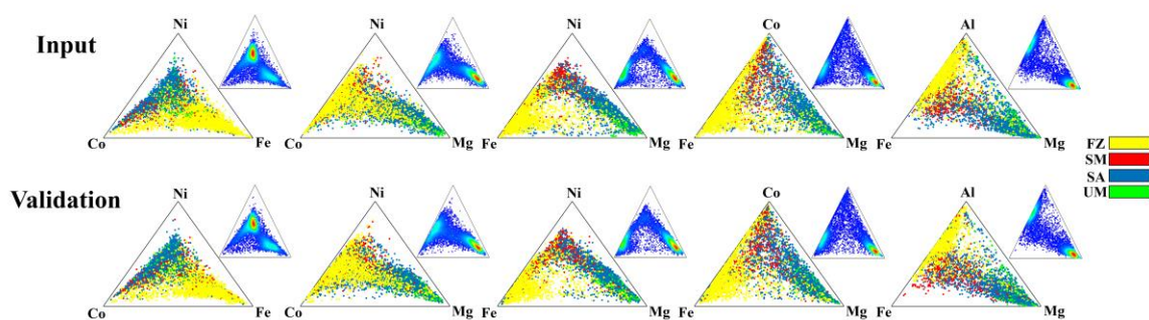
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781 **Fig. 5** Ternary diagrams of the geochemical compositions of input and validation sets, coloured by the rock types

782 (large triangles) and kernel density (small triangles)

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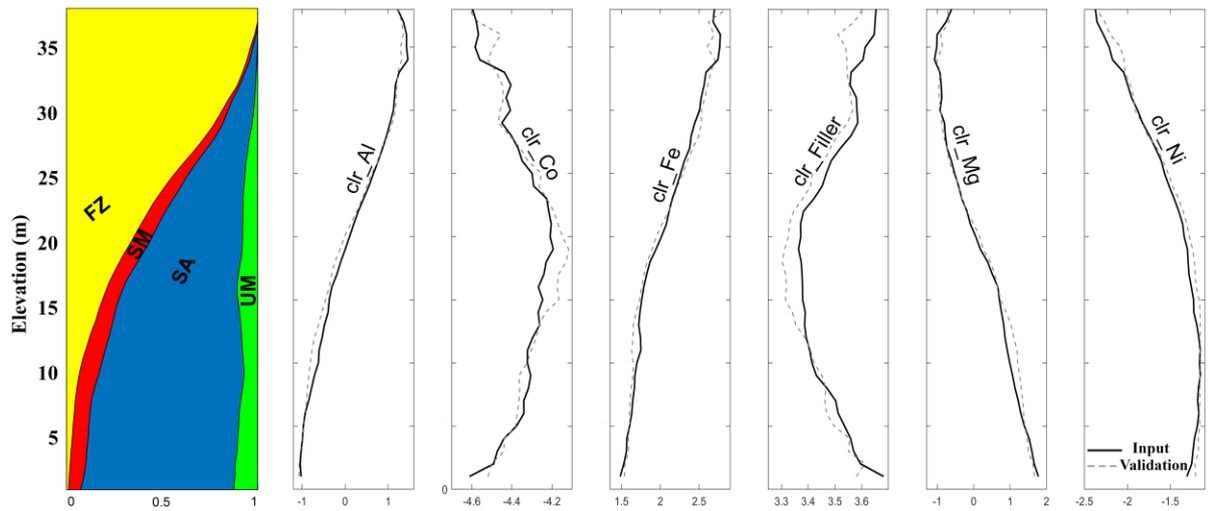
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792 **Fig. 6** Vertical proportion curves of different rock types and clr-transformed geochemical components

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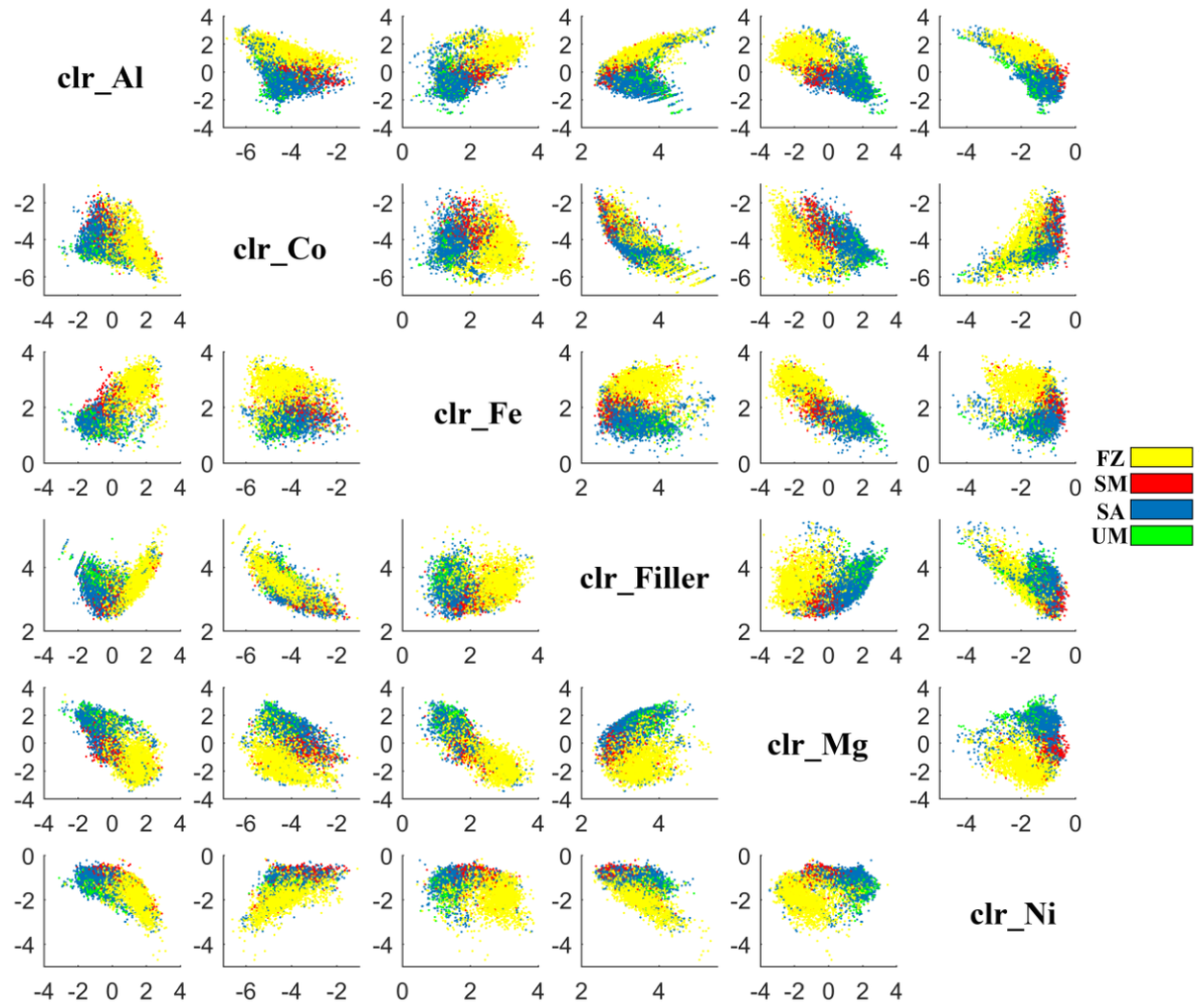
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801 **Fig. 7** Scatterplots of clr-transformed geochemical components (upper triangle is input and lower triangle is  
 802 validation set)

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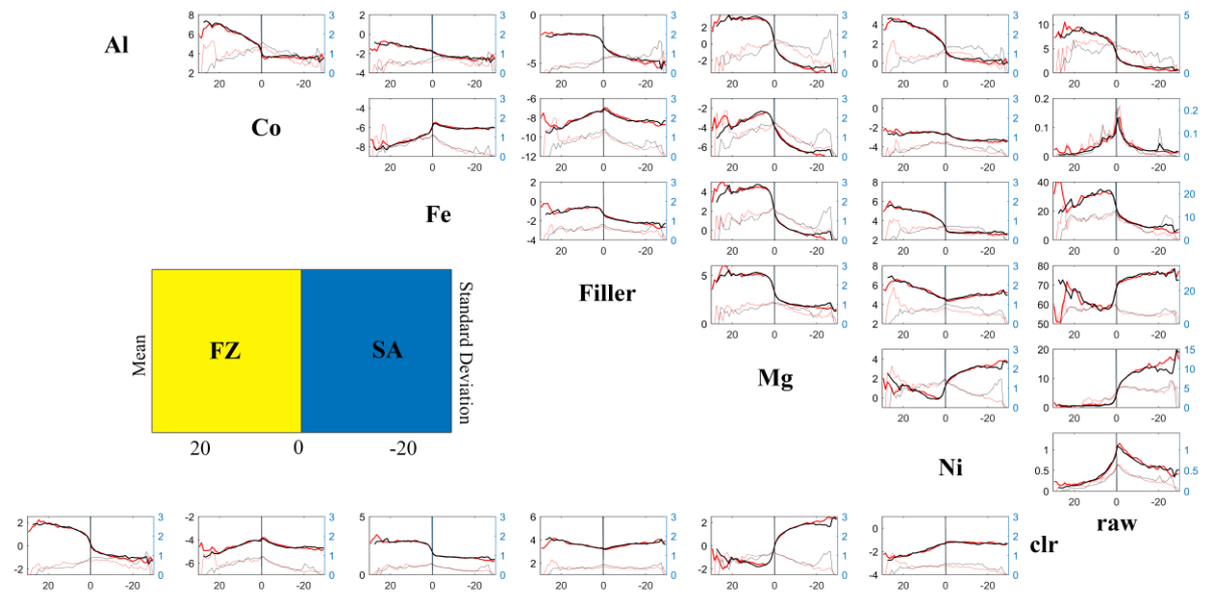
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811 **Fig. 8** Compositional contact analysis for two dominant geological domains (FZ and SA). Mean values and  
 812 standard deviations are represented by continuous and dashed lines respectively (black for input set and red for  
 813 validation set)

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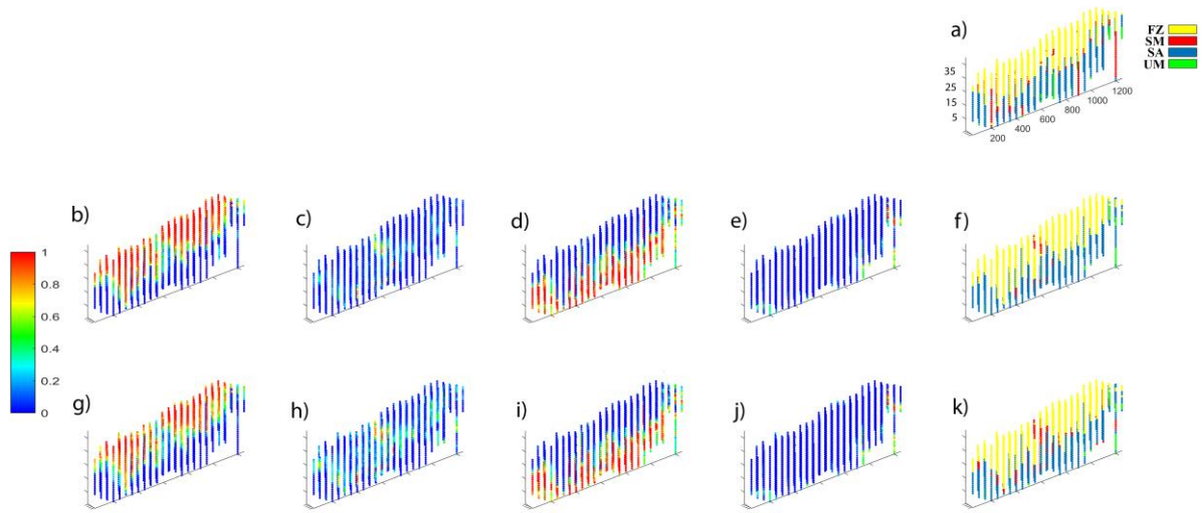
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821 **Fig. 9** Cross sections of validation boreholes for northing 300m and 50m thickness. a) true rock types. b) to e)  
 822 probability of FZ, SM, SA, and UM respectively. f) most probable rock types. g) to j) adjusted probability of FZ,  
 823 SM, SA, and UM respectively. k) adjusted most probable rock types

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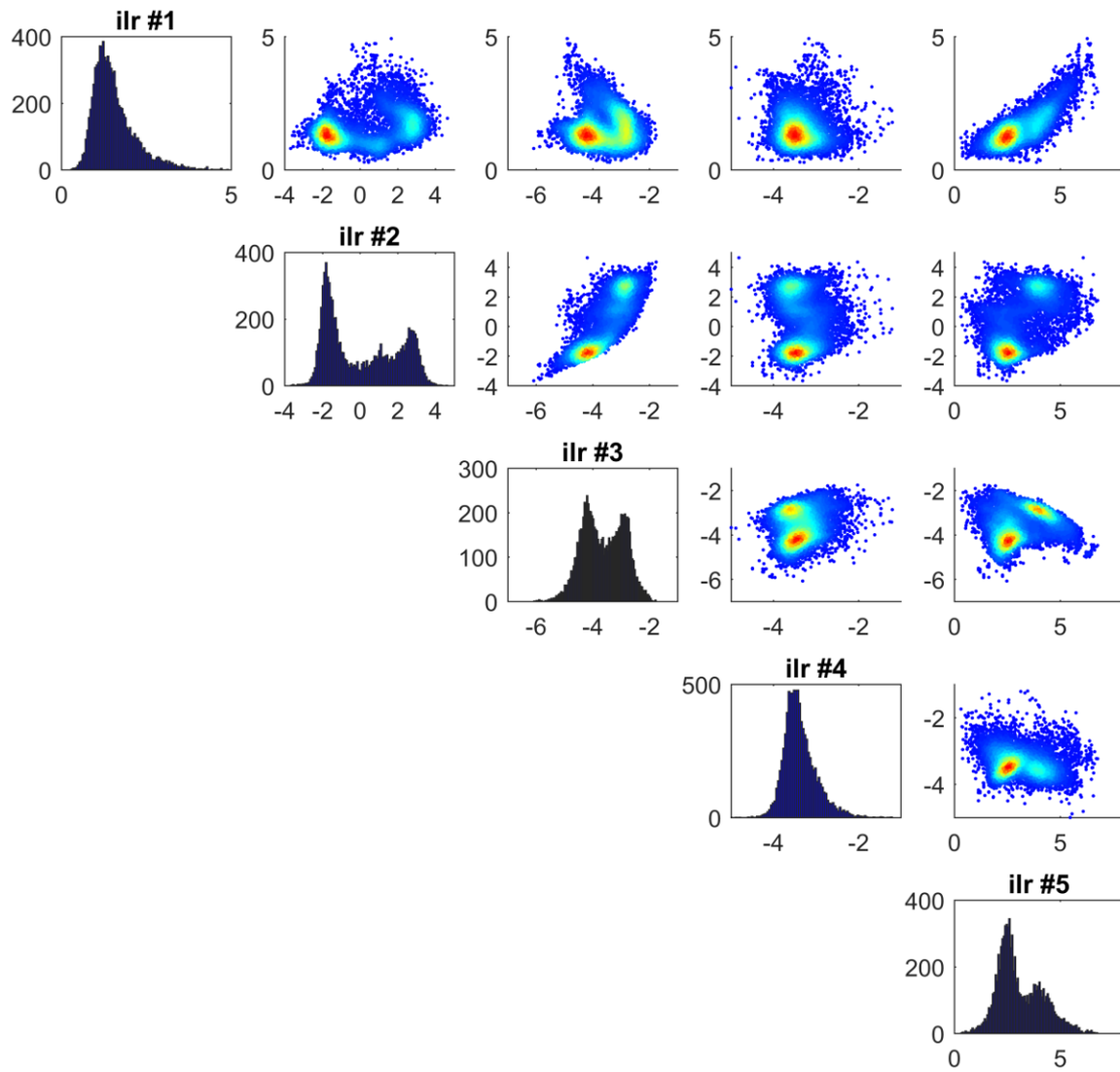
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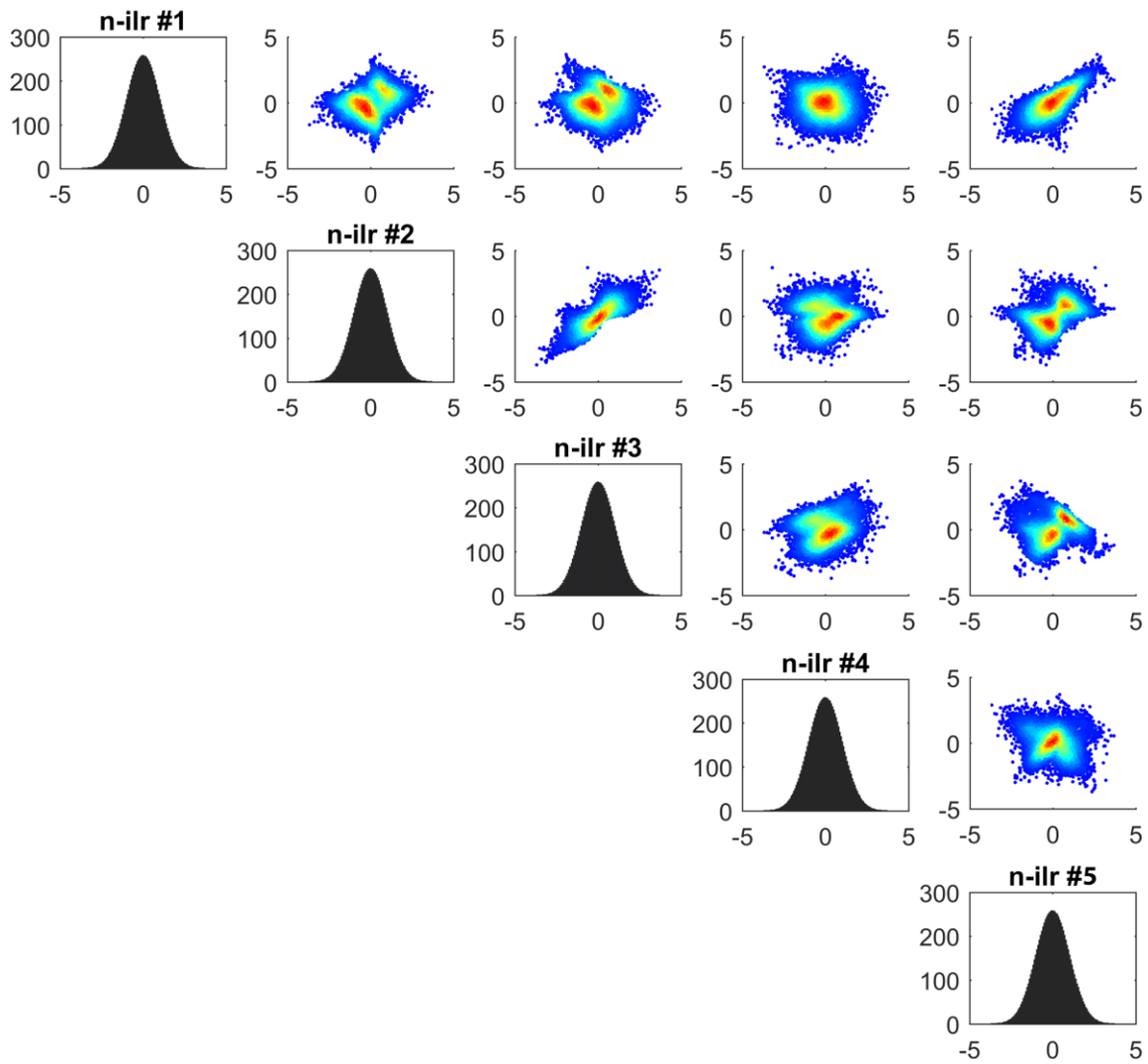
**Fig. 10** Histograms and scatterplots of ilr-transformed input data (coloured by kernel density estimate)

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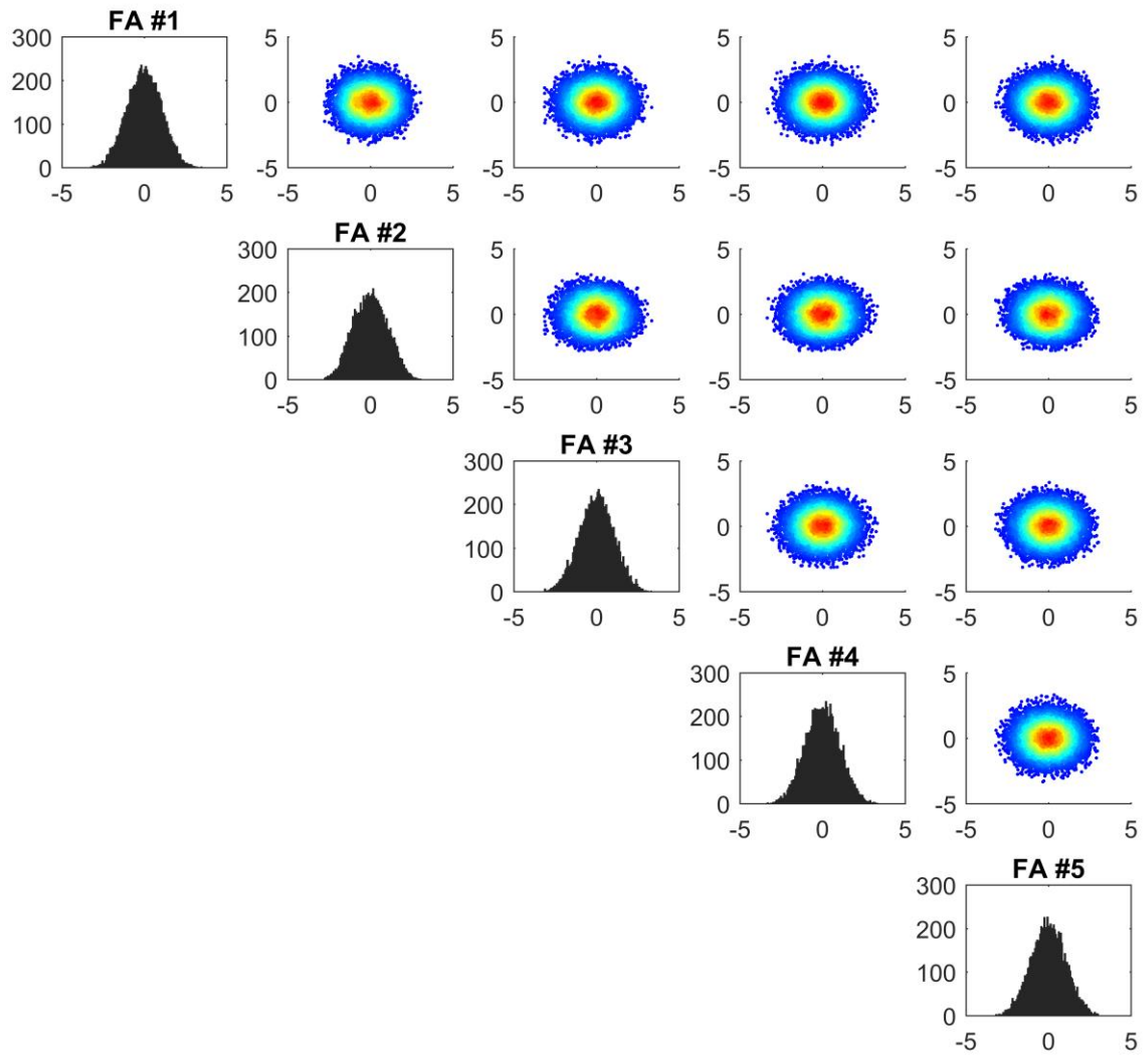


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842 **Fig. 11** Histograms and scatterplots of the transformed data to normal space via a GA (coloured by kernel density  
843 estimate)

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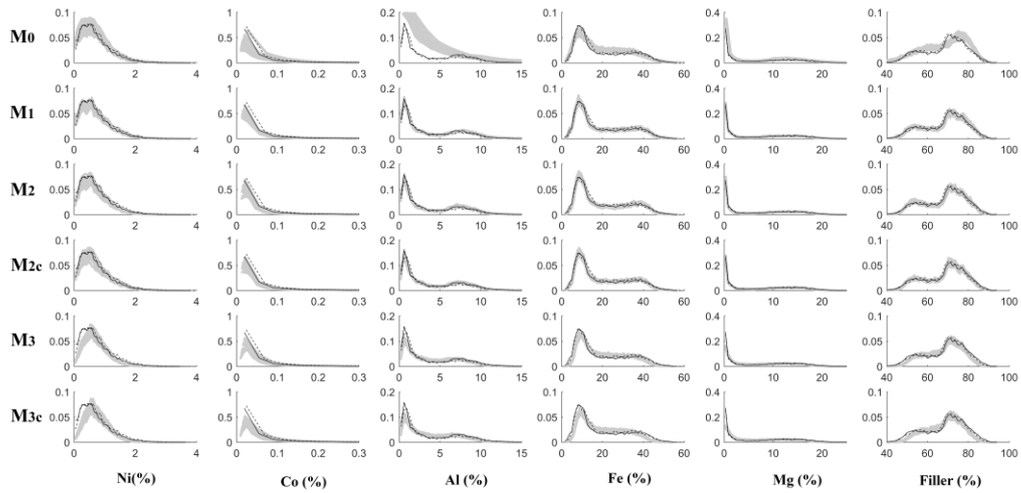
847 **Fig. 12** Histograms and scatterplots of the transformed data to normal space via FA (coloured by kernel density

848 estimate)

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853 **Fig. 13** Histogram reproduction of the six proposed methods for simulation of geochemical compositions.

854 Continuous black lines are input data, dashed black lines are validation data, and grey lines are realisations

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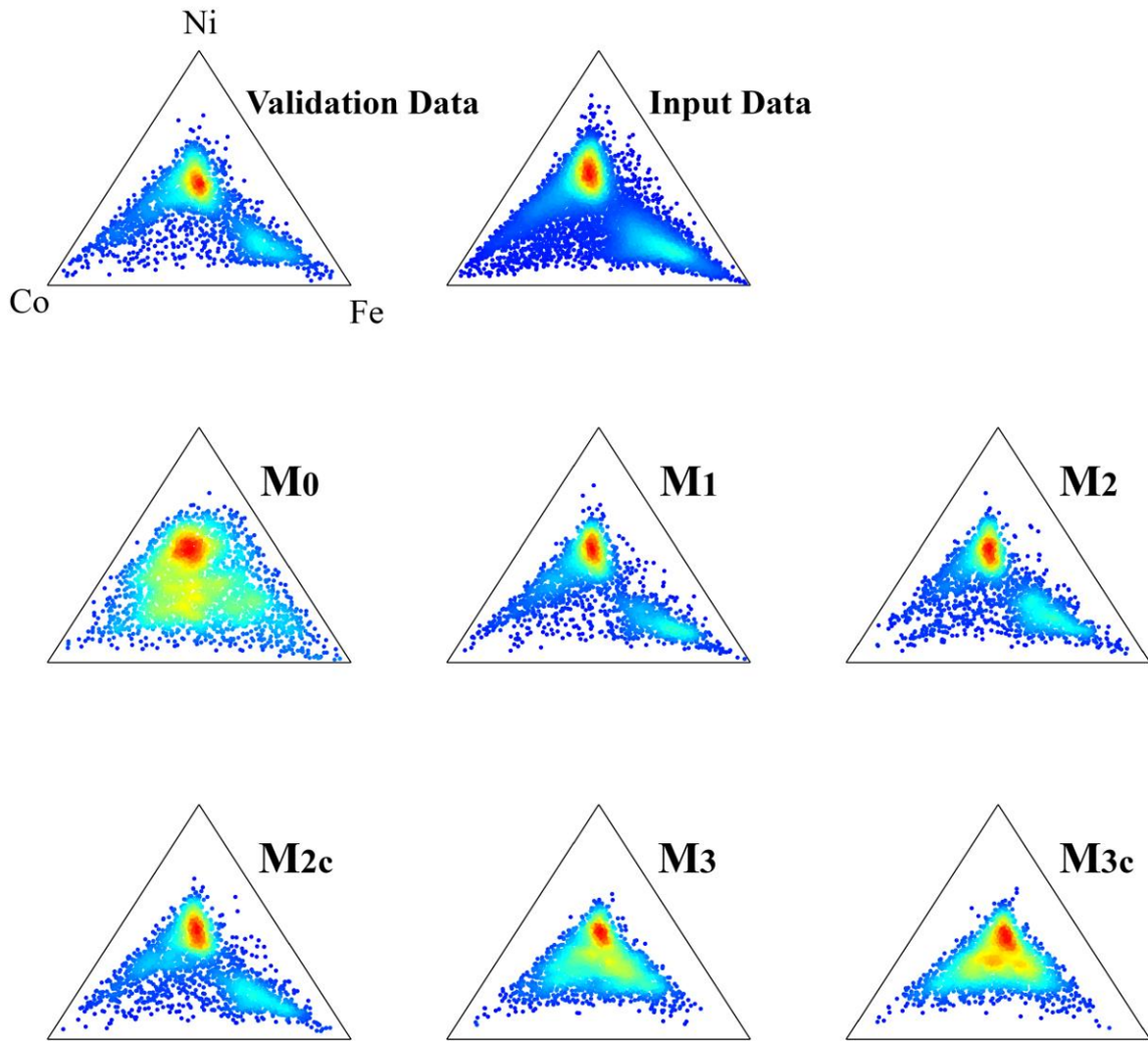
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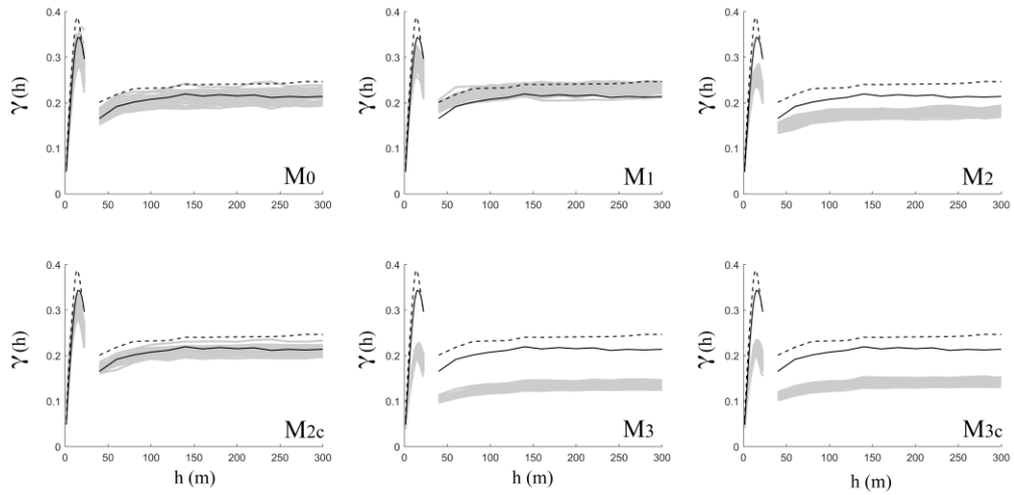
863 **Fig. 14** Ternary diagrams of input and validation data (three components: Ni, Co, Fe), and one realisation  
 864 (randomly selected) from each method

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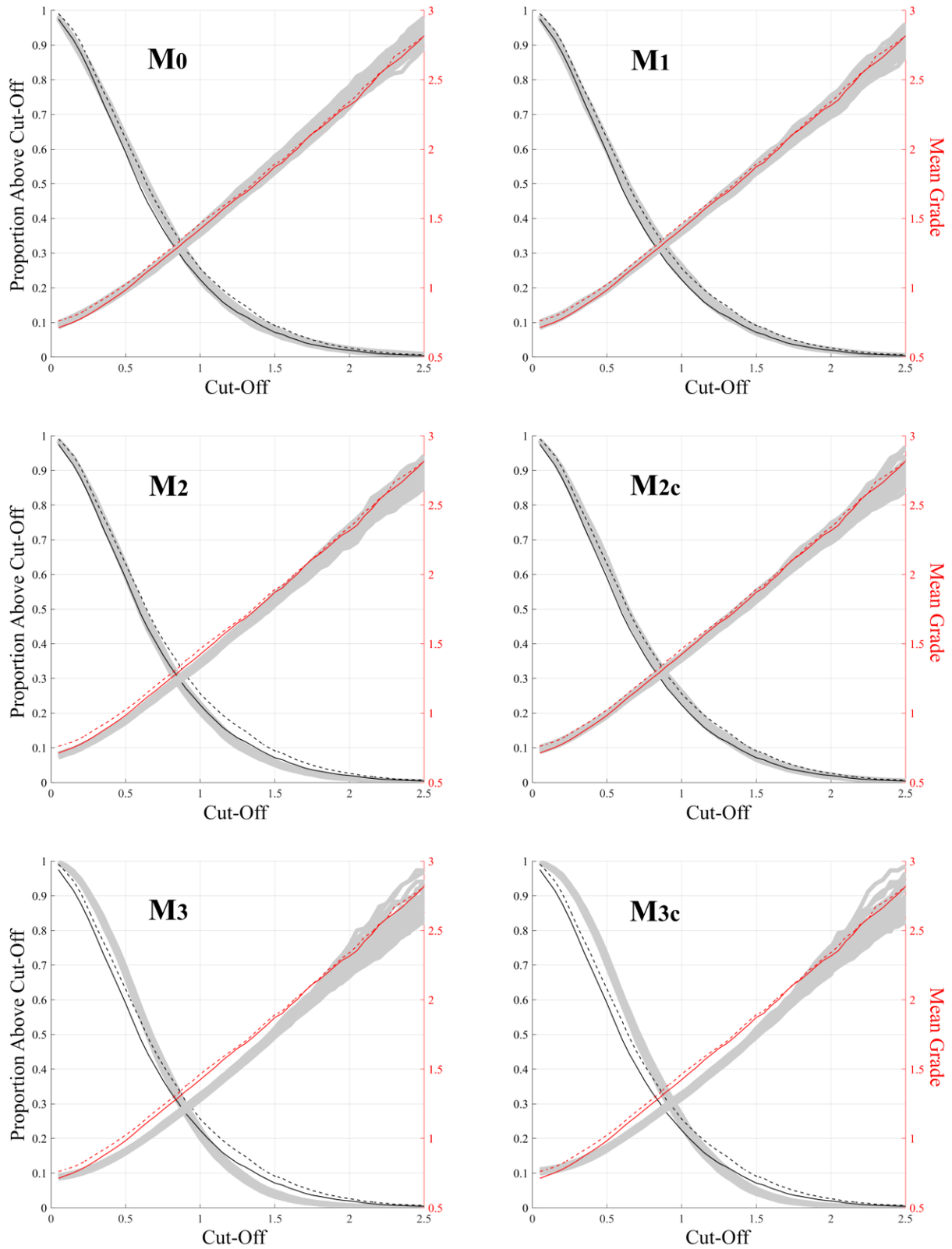
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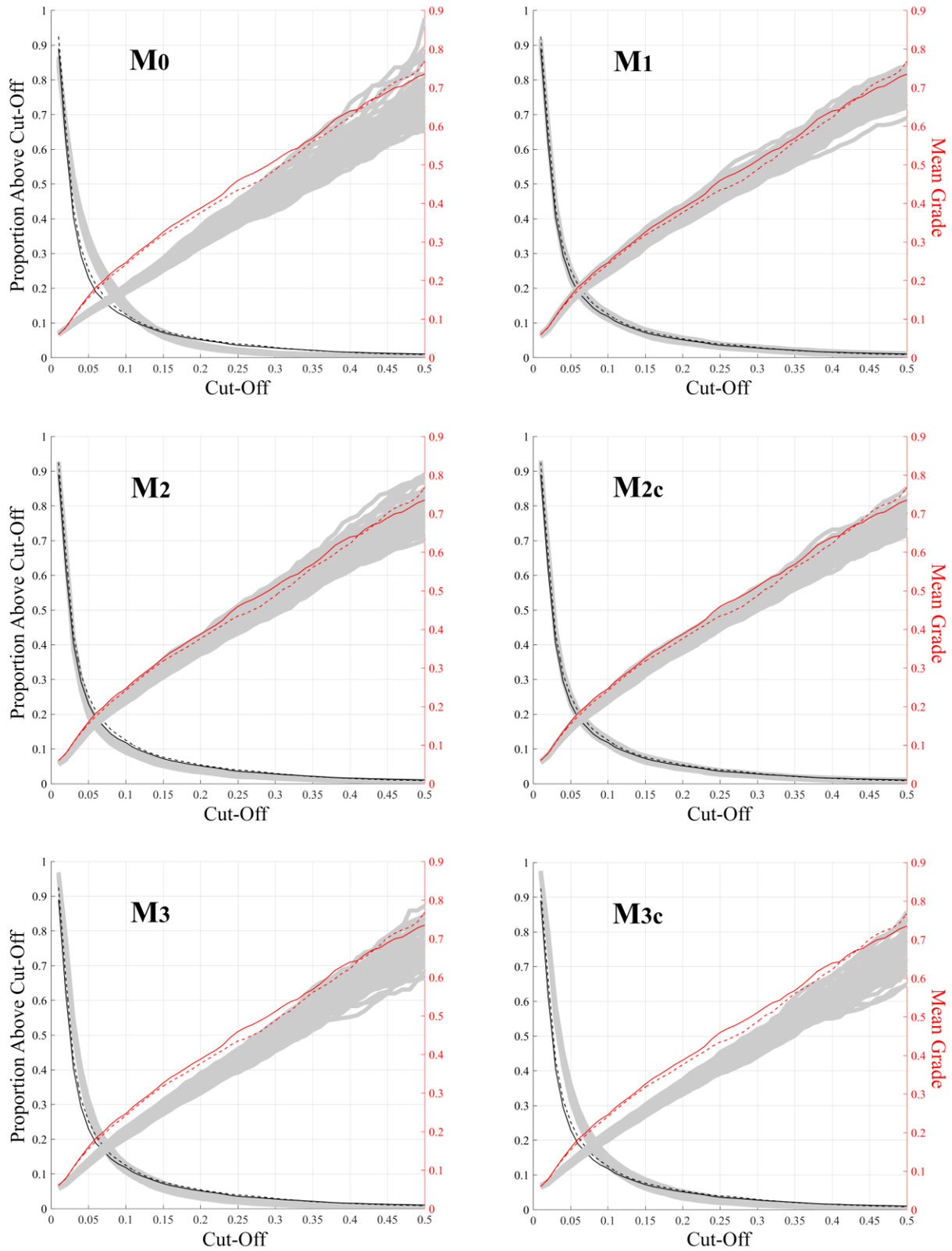
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870 **Fig. 15** Experimental variogram reproduction (Ni component) of the six proposed methods in vertical (short range)  
871 and horizontal (long range) directions.



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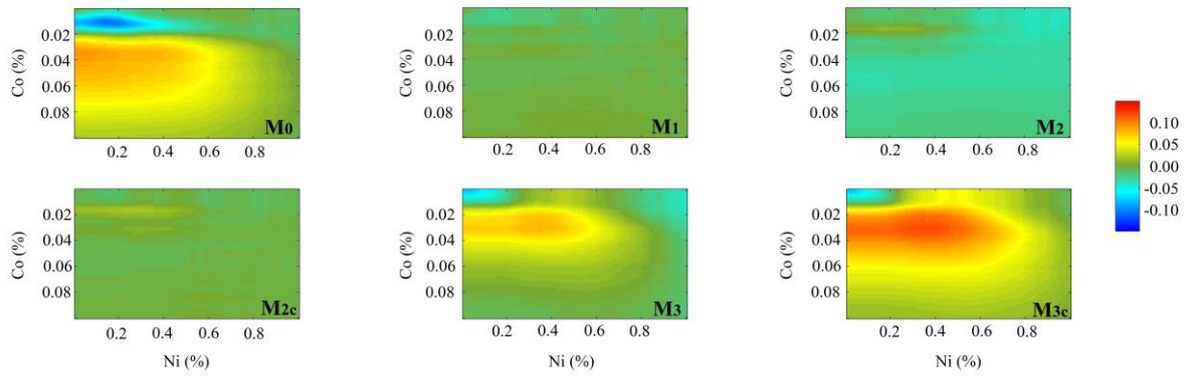
873 **Fig. 16** Grade-tonnage curves (for Ni component) of the six proposed methods. Continuous black lines are the  
 874 proportion of samples above Ni cut-offs while continuous red lines are the average grades for input data. Dashed  
 875 lines are for validation data while grey lines are different realisations.



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877 **Fig. 17** Grade-tonnage curves (for Co component) of the six proposed methods Continuous black lines are the  
 878 proportion of samples above Co cut-offs while continuous red lines are the average grades for input data. Dashed  
 879 lines are for validation data while grey lines are different realisations.

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882 **Fig. 18** The difference between the expected proportions above cut-offs (Ni and Co), calculated from realisations,  
883 and real proportions above cut-offs, calculated from the validation data

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Table 1 Proposed methods and the related features

<b>Code</b>	<b>Geological control</b>	<b>Adjust proportions</b>	<b>Transform to real space</b>	<b>Transform to normal</b>	<b>Factorization</b>	<b>Spatial simulation</b>
<b>M<sub>0</sub></b>	No geological control	-	ilr	GA	MAF	TB
<b>M<sub>1</sub></b>	No geological control	-	ilr	FA	-	TB
<b>M<sub>2</sub></b>	Deterministic geological model	-	ilr	FA	-	TB
<b>M<sub>2c</sub></b>	Deterministic geological model	✓	ilr	FA	-	TB
<b>M<sub>3</sub></b>	Probabilistic geological model	-	ilr	FA	-	TB
<b>M<sub>3c</sub></b>	Probabilistic geological model	✓	ilr	FA	-	TB

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Table 2 Ore mineralogy of different geological units at MME

<b>Geological unit</b>	<b>Kaolin</b>	<b>Goethite</b>	<b>Hematite</b>	<b>Maghemite</b>	<b>Chlorite</b>	<b>Smectite</b>	<b>Lizardite</b>
<b>FZ</b>	<50%	<50%	< 10%	<50%	-	-	-
<b>SM</b>	<2%	< 10%	<2%	<2%	< 10%	>75%	<2%
<b>SA</b>	-	< 10%	<2%	<2%	<2%	<50%	<50%
<b>UM</b>	-	-	-	-	-	-	-

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Table 3 Proportions of rock types

	<b>Validation</b>	<b>Input</b>	<b>Most probable map</b>	<b>Most probable map (adjusted)</b>
<b>FZ</b>	0.352	0.367	0.417	0.376
<b>SM</b>	0.092	0.068	0.027	0.084
<b>SA</b>	0.485	0.486	0.479	0.462
<b>UM</b>	0.071	0.079	0.077	0.078

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