Simultaneous Identification of Number, Location, and Release History of Groundwater Contamination Sources

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Abstract: In previous studies, a 0-1 mixed integer nonlinear programming optimization model (0-1MINLPOM) could only identify the location and release intensity for groundwater contamination sources (GCSs), and the location of each GCS was regarded as a 0-1 integer variable, selected from several locations determined in advance. However, in actual situations, the locations usually cannot be accurately isolated to a few GCSs and the number of GCSs is often unknown, so 0-1MINLPOM was improved in this study. Based on the estimation that there is a maximum of three GCSs in the study area, an improved 0-1 MINLPOM was established to simultaneously identify the number of GCSs (treated as 0-1 integer variable), the location (treated as integer variable) and release history of GCS (treated as continuous variables).
The simulation model was constructed as an equality constraint embedded improved 0-1 MINLPOM. In the improved 0-1 MINLPOM solution process, repeatedly calling the simulation model would have incurred a massive computational load and taken a long time. Thus, a surrogate model based on kriging and extreme learning machine (ELM) was established respectively for the simulation model to avoid this shortcoming. The results show that the accuracy of the kriging surrogate model (Krig-SM) was higher compared with the ELM surrogate model (ELM-SM). The improved 0-1 MINLPOM could identify the number, location, and release history of GCSs simultaneously. The accuracy of identifying the number of GCSs was 100%, and the accuracies of identifying the locations and release history were above 91.67% and 90.14%, respectively.

**Keywords:** Groundwater contamination, Mixed integer nonlinear programming, Source number, Surrogate model.

**Introduction**

Groundwater contamination beneath the surface of the earth is characterized by concealment and hysteresis of discovery, which leads to a lack of understanding of GCSs, including their number, location, and release history in aquifers (Mahinthakumar and Sayeed, 2005; Jha and Datta, 2013). The release history refers to the contaminant release intensity in each release period (Atmadja and Bagtzoglou, 2001; Sun et al., 2006). As a consequence, the design of groundwater contamination remediation schemes is difficult, and the responsibility for contamination liabilities and the assessment of contamination risks are also problematic (Lapworth et al., 2012; Om and Bithin, 2013). Developing an effective method for identifying GCSs would address these
problems, and have important significance (Bagtzoglou and Atmadja, 2005).

The identification of GCSs roughly started in the 1980s and many methods have been proposed for identifying GCSs, including optimization approaches (Gorelick et al., 1983; Mahar and Datta, 2000; Zhao et al., 2015), direct approaches (Skaggs and Kabala, 1994), probabilistic and geostatistical simulation approaches (Neupauer and Wilson, 2000), and analytical solution and regression approaches (Sidauruk et al., 2010).

Among those methods, simulation optimization methods have been used widely to characterize GCSs. Gorelick et al. (1983) applied a method based on least squares regression and linear programming to identify the location and release intensity of GCSs. Woodbury and Ulrych (1996) employed the minimum relative entropy method to recover the release and evolution history of contamination plumes in one-dimensional systems. Mahar and Datta (2000) used a nonlinear optimization model to identify the location and release intensity for GCSs in an unsteady flow system. Mahinthakumar and Sayeed (2006) employed hybrid optimization method to reconstruct release histories of GCSs. Jha and Datta (2011) applied a simulation-optimization based a variant of simulated annealing algorithm to identify the unknown source flux magnitude, duration and timing. Zhao et al. (2015) employed optimization approaches to identify the location and release intensity for GCSs. Xu and Gómez-Hernández (2016) used an ensemble Kalman filter to identify the source location, release time, and release concentration of GCSs.

Various preconditions are considered when applying simulation optimization methods to identify GCSs including: (1) the locations and number of GCSs are known to identify the release history (Zhao et al., 2016); (2) the suspected locations (3-4 specific locations) of the GCSs are known to identify the true location and release history (Ayvaz, 2010; Guo et al. 2018); (3) the
number (usually set to one) and all locations where the contaminant may be released are known to identify the true location and release history (Yeh et al., 2014; Xu and Gómez-Hernández, 2016).

Although good research results have been achieved in the application of simulation optimization methods for pollution source identification, there is no study have reported the application of simulation optimization methods for simultaneously identifying the number, location, and release history of GCSs. Guo et al. (2018) used the 0-1 MINLPOM to identify the location (treated as a 0-1 integer variable) and release intensity (treated as a continuous variable) of GCSs, but the number of GCSs was not identified. Thus, in the present study, an improved 0-1 MINLPOM was developed to simultaneously identify the number, location, and release history of GCSs. In the improved 0-1 MINLPOM, the number of GCSs is regarded as a 0-1 integer variable, and the locations and release history of GCSs are regarded as an integer variable and continuous variable, respectively.

When solving the improved 0-1 MINLPOM, the simulation model must be called repeatedly for calculations, thereby incurring a large calculation load and requiring a long time. This problem can be solved by establishing a surrogate model of the simulation model. The previously studies to identify GCSs based on surrogate models include kriging (Simpson et al., 2001; Luo and Lu, 2014a), radial basis function, support vector regression and extreme learning machines (Jiang et al., 2015; Hou and Lu, 2018).

The kriging and ELM methods were used to establish surrogate models for the simulation model, and the surrogate model with higher accuracy was embedded in the improved 0-1 MINLPOM. The improved 0-1 MINLPOM based on the surrogate model was then used to simultaneously identify the number, location, and release history of GCSs.
Methodology

Kriging

The kriging method is also known as the spatial local interpolation method and it is based on variation function theory and structural analysis for obtaining unbiased optimal estimates of regionalized variables in a finite region (Bargaoui and Chebbi., 2009; Kleijnen., 2009). In recent years, the kriging method has been extended as a surrogate modeling method with applications in many fields of engineering (Ryu et al., 2002; Kleijnen and van Beers, 2005; Coetzee et al., 2012).

A kriging surrogate model is established according to the following principle:

\[ Y(x) = \sum_{i=1}^{n} f_i(x) \cdot \beta_i + Z(x) = f^T \cdot \beta + Z, \]  

where: \( x \) is the input value of the training samples, \( f(x) = [f_1(x), f_2(x), \ldots, f_p(x)]^T \) are known determinate regression functions, \( \beta = [\beta_1, \beta_2, \ldots, \beta_p]^T \) is a regression coefficient matrix estimated from the training samples, \( p \) is the number of determinate regression functions, and \( Z(x) \) is a random part deviation of the regression model, which must satisfy the following conditions:

\[
\begin{align*}
E(Z(x)) &= 0 \\
D(Z(x)) &= \sigma^2 \\
\text{cov}(Z(x_i), Z(x_j)) &= \sigma^2 R(x_i, x_j) 
\end{align*}
\]

where \( R(x_i, x_j) \) is the spatial correlation function between any two sampling points \( x_i \) and \( x_j \), \( \sigma^2 \) is variance of \( Z(x) \):

\[ R(x_i, x_j) = \exp\left(-\sum_{k=1}^{n} \theta_k |x_i^k - x_j^k|\right) \quad (i = 1, 2, \ldots, n, j = 1, 2, \ldots, n), \]

where \( \theta_k \) is the undetermined coefficient and \( x_i^k \) is the k-dimensional coordinate of the \( i \)th sample.

Using the input and output data for \( n \) known sample points, the output value corresponding to
any point \( x^* \) in the predicted feasible domain is:

\[
Y(x) = f^T \beta^* + r^T(x)R^{-1}(y - f^T \beta^*),
\]

(4)

where \( r(x) \) is the correlation vector of point \( x^* \) and \( n \) sampling points \( \{x_1, x_2, \ldots, x_n\} \), \( y \) is the matrix \( n \times m \), \( n \) is the number of sampling points, \( m \) is the dimension of the output value, and \( \beta^* \) is the undetermined coefficient of the regression part, which can be obtained by the optimal linear unbiased estimation:

\[
r^*(x) = \begin{bmatrix} R(x_1, x_1) & L & R(x_1, x_2) \\ & M & O \\ R(x_n, x_1) & L & R(x_n, x_2) \end{bmatrix}
\]

(5)

\[
\beta^* = (f^T R^{-1} f)^{-1} f^T R^{-1} y,
\]

(6)

where \( R \) is the correlation matrix comprising the correlation coefficients of \( n \) sampling points:

\[
R = \begin{bmatrix} R(x_1, x_1) & L & R(x_1, x_2) \\ & M & O \\ R(x_n, x_1) & L & R(x_n, x_2) \end{bmatrix}.
\]

(7)

The variance estimate value \( \sigma^2 \) is determined by:

\[
\sigma^2 = \frac{1}{n} (y - f^T \beta^*)^T R^{-1} (y - f^T \beta^*)
\]

(8)

\[
\max_{\sigma^2} \left[ -n \ln \sigma^2 + \ln |R| \right].
\]

(9)

The surrogate model can be established by solving the nonlinear unconstrained optimization problem defined above.

Extreme learning machine

Huang et al. (2004) proposed the ELM method to improve backward propagation neural networks in order to improve the low efficiency of learning, while also simplifying the settings for the learning parameters. Compared with the backward propagation neural network algorithm, the ELM method has advantages in terms of its rapid learning speed and good generalization performance (Huang et al., 2006). ELM is a type of machine learning method based on a
feedforward neural network (Huang et al., 2015), as illustrated in Fig. 1.

Fig. 1. Schematic diagram showing the single hidden layer feedforward neural network of ELM.

Thus, ELM was used to establish the surrogate model for the simulation model because of these advantages. A single hidden layer feedforward neural network comprises an input layer, hidden layer, and output layer. The input layer is fully connected to the hidden layer, and the hidden layer is fully connected to the output layer. The input layer contains $m$ neurons corresponding to each input variable, the hidden layer has $n$ neurons, and the output layer has $k$ neurons corresponding to each output variable. The connection weight matrix between the input layer and the hidden layer is $\omega$. The connection weight matrix between the hidden layer and output layer is $\beta$. The threshold value matrix for the hidden layer neurons is $b$. These matrices are expressed as follows:

$$\omega = \begin{bmatrix} \omega_{11} & \ldots & \omega_{1m} \\ \vdots & \ddots & \vdots \\ \omega_{m1} & \ldots & \omega_{mm} \end{bmatrix}_{m \times n}$$ (10)

$$\beta = \begin{bmatrix} \beta_{11} & \ldots & \beta_{1k} \\ \vdots & \ddots & \vdots \\ \beta_{k1} & \ldots & \beta_{kk} \end{bmatrix}_{k \times n}$$ (11)

$$b = \begin{bmatrix} b_1 \\ \vdots \\ b_n \end{bmatrix}_{n \times 1}$$ (12)

where $\omega_{ij}$ is the connection weight between the $i$-th neuron in the input layer and the $j$-th neuron in the hidden layer, and $\beta_{jk}$ is the connection weight between the $j$-th neuron in the hidden layer and the $k$-th output variable.
hidden layer and the $l$-th neuron in the input layer.

The input matrix $X$ and output matrix $Y$ comprise training data sets containing $M$ samples, as follows.

$$
X = \begin{bmatrix}
x_{11} & \cdots & x_{1M} \\
\vdots & & \vdots \\
x_{n1} & \cdots & x_{nM}
\end{bmatrix}_{n \times M} 
$$ (13)

$$
Y = \begin{bmatrix}
y_{11} & \cdots & y_{1M} \\
\vdots & & \vdots \\
y_{n1} & \cdots & y_{nM}
\end{bmatrix}_{n \times M} 
$$ (14)

The activation function for the hidden layer neurons is $g(x)$ and the output $T$ from the network is:

$$
T = \begin{bmatrix} t_1 & \cdots & t_M \end{bmatrix} 
$$ (15)

$$
t_j = \begin{bmatrix} t_{ij} & \cdots & t_{Mj} \\
\vdots & & \vdots \\
t_{ij} & \cdots & t_{Mj}
\end{bmatrix}_{M \times 1} = \begin{bmatrix} \sum_{i=1}^{M} \beta_{ij}g(\omega_{ij} + b_i) \end{bmatrix}_{M \times 1} 
$$ (16)

The objective when training a single hidden layer neural network is to minimize the error between the output data and training data, which can be expressed as follows.

$$
\sum_{j=1}^{M} \left\| t_j - y_j \right\| = 0 
$$ (17)

$$
\omega_i = [\omega_{ij} \ L \ \omega_{in}] ; x_j = [x_{ij} \ L \ x_{jn}]^T ; y_j = [y_{ij} \ L \ y_{jn}]^T 
$$ (18)

Thus, we calculate $\omega$, $\beta$, and $b$ while ensuring that Eq. (19) holds:

$$
\begin{bmatrix}
\sum_{i=1}^{M} \beta_{ij}g(\omega_{ij} + b_i) & \cdots & \sum_{i=1}^{M} \beta_{ij}g(\omega_{ij} + b_i) \\
\vdots & & \vdots \\
\sum_{i=1}^{M} \beta_{ij}g(\omega_{ij} + b_i) & \cdots & \sum_{i=1}^{M} \beta_{ij}g(\omega_{ij} + b_i)
\end{bmatrix}_{M \times M} = Y^T 
$$ (19)

The input weight and the hidden layer threshold are determined randomly before training and they remain unchanged during the training process, so the hidden layer's output matrix $H$ is uniquely determined. Training a single hidden layer neural network can be transformed into solving the linear equations in Eq. (21) to obtain the output weight matrix. The final output layer
connection weight matrix can be obtained by solving the least squares solution of Eq. (20):

\[ H\beta = Y^T \] (20)

\[ \min_{\beta} \| H\beta - Y^T \| \] (21)

\[ H = \begin{bmatrix} g(\omega_1 x_i + b_i) & L & g(\omega_1 x_i + b_i) \\ M & O & M \\ g(\omega_1 x_M + b_i) & L & g(\omega_1 x_M + b_i) \end{bmatrix}_{M \times n} \] (22)

\[ \hat{\beta} = H^*Y^T, \] (23)

where \( H^* \) is the Moore-Penrose generalized inverse matrix of the hidden layer output matrix H.

The output corresponding to any input can be predicted after calculating the output layer connection weight matrix.

**Numerical applications**

**Site overview**

A hypothetical contaminated site was considered as a case study in order to analyze the application of the research methods for the identification of groundwater contamination sources. The hypothetical contaminated site was designed to represent the conditions that may occur in most actual problems, including the aquifer geometry, boundary conditions, initial conditions, hydrogeological parameters, and groundwater flow characteristics (Prakash and Datta, 2014; Datta et al., 2017). Contaminant is considered to be divalent manganese ion which do not undergo chemical and biological conversion. The study area comprised a two-dimensional, heterogeneous, isotropic, submerged aquifer with irregular boundaries, where the groundwater flow was transient. There three parameter zones in the aquifer and their medium types were as follows: I, coarse sand, II, medium sand and III, fine sand. The boundary conditions, locations of observation wells and potential contamination source area (all locations where the contaminant might have been released) are shown in Fig. 2, and the other parameters related to the aquifer are presented in Table 1. The hydraulic head of the northwest specific head boundary was 27 m and the hydraulic head of the...
southeast specific head boundary was 25 m. The vertical direction of the study area received uniform recharge from atmospheric rainfall with a recharge amount of 730 mm/a. The initial concentration of the contaminant in the study area was 0 g/L.

In order to test the effectiveness of the identification method developed in this study, two hypothetical cases are designed in the paper. The aquifer parameters of hypothetical contaminated site were the same for the two cases, as shown in Table 1.

Table 1. Parameters for the aquifers in the hypothetical contaminated site

| Parameters                        | Values | Value | I   | II  | III |
|-----------------------------------|--------|-------|-----|-----|-----|
| Hydraulic conductivity in x-direction, Kxx (m/d) | 42     | 30    | 18  |     |     |
| Hydraulic conductivity in y-direction, Kyy (m/d) | 42     | 30    | 18  |     |     |
| Effective porosity, θ             | 0.25   | 0.2   | 0.18|     |     |
| Longitudinal dispersivity, αL (m) | 40     | 40    | 40  |     |     |
| Transverse dispersivity, αT (m)   | 8      | 8     | 8   |     |     |
| Saturated thickness, b(m)         | 35     | 35    | 35  |     |     |
| Grid spacing in x-direction, Δx (m)| 3      | 3     | 3   |     |     |
| Grid spacing in y-direction, Δy (m)| 3      | 3     | 3   |     |     |
| Length of the simulation period, Δt (month) | 6      | 6     | 6   |     |     |
| Volume flux per unit area, W (m/d) | 0.0004 | 0.00035 | 0.0003 |     |
| Initial concentration (g/L)       | 0      | 0     | 0   |     |     |

The study area was discretely divided into 5262 grids. Since the divided grid is treated as an integer variable, the location information of the potential contamination source area in the study area is represented by grid numbers. The horizontal and longitudinal locations of the potential contamination source areas and their corresponding grid numbers are shown in Table 2.

Table 2. The horizontal and longitudinal locations and their corresponding grid numbers

(a) horizontal

| Number | Location | Number | Location | Number | Location | Number | Location |
|--------|----------|--------|----------|--------|----------|--------|----------|
| 19     | 55.5     | 32     | 94.5     | 45     | 133.5    | 58     | 172.5    |
| 20     | 58.5     | 33     | 97.5     | 46     | 136.5    | 59     | 175.5    |
| 21     | 61.5     | 34     | 100.5    | 47     | 139.5    | 60     | 178.5    |
| 22     | 64.5     | 35     | 103.5    | 48     | 142.5    | 61     | 181.5    |
| 23     | 67.5     | 36     | 106.5    | 49     | 145.5    | 62     | 184.5    |
| 24     | 70.5     | 37     | 109.5    | 50     | 148.5    | 63     | 187.5    |
| 25     | 73.5     | 38     | 112.5    | 51     | 151.5    | 64     | 190.5    |
| 26     | 76.5     | 39     | 115.5    | 52     | 154.5    | 65     | 193.5    |
| 27     | 79.5     | 40     | 118.5    | 53     | 157.5    | 66     | 196.5    |
| 28     | 82.5     | 41     | 121.5    | 54     | 160.5    | 67     | 199.5    |
| 29     | 85.5     | 42     | 124.5    | 55     | 163.5    | 68     | 202.5    |
The real number, location, and release history of GCSs for the two cases are designed as follows:

Case one: the number of GCSs was one, the location of the GCSs is as shown in Fig. 2a and release intensities of the GCSs during two release periods are as shown in Table 3a.

Case two: the number of GCSs was two, the locations of the two GCSs are shown in Fig. 2b and release intensities of the two GCSs during two release periods are as shown in Table 3b.

Table 3. Details of the real GCSs for case two: (a) case one; (b) case two.

| Number of sources | Grid number | Release intensity (g/d) |
|-------------------|-------------|------------------------|
|                   | Horizontal | longitudinal | SP1 | SP2 |
| 1                 | 35         | 52          | 260  | 140 |

(b) case one

| Number of sources | Sources | Grid number | Release intensity (g/d) |
|-------------------|---------|-------------|------------------------|
|                   |         | Horizontal | SP1 | SP2 |
| 2                 | S1      | 35         | 190 | 120 |
|                   | S2      | 41         | 170 | 160 |
During the inverse identification of GCSs process, the number, location, and release history of the GCSs for the two cases were regarded as unknown, what known is only the potential contamination source area (shown in Fig. 2a and Fig. 2b). For hypothetical case one and case two it is estimated that the maximum number of GCSs is 3. The number, location, and release intensity of the GCSs were then identified based on optimization model.

All the GCSs are assumed to have same activity initiation times (the first day of simulation) and the time lag between the initial release contaminant of the GCSs and first contaminant concentration measurement of the two cases are shown in Table 4. The activity duration of two cases is 300 days. The entire activity duration of the sources is divided into two equal stress periods of 150 days. Concentration is measured every 90 days, starting from the first concentration measurement time. The contaminant flux from the sources is assumed to be constant over a stress period. The total simulation time for contaminant transport is 900 days. A total of eight periods concentration measurements from each of the fourteen observation wells are utilized for the two cases.

| Case   | Activity duration (d) | Time lag between the initial release contaminant of the source and first concentration measurement (d) | Concentration measurement intervals (d) |
|--------|-----------------------|-------------------------------------------------------------------------------------------------|----------------------------------------|
| one    | 2×90                  | 270                                                                                             | 90                                      |

Table 4. Test case scenarios
Numerical simulation model

In order to identify the GCSs, a numerical simulation model of the groundwater flow and contaminant transport was established based on the specific conditions of the study area. The governing partial differential equations for the groundwater flow and the contaminant transport of the transient flow in the two-dimensional aquifer system are defined as follows (Pinder and Bredehoeft, 1968; Singh and Datta, 2006):

The governing partial differential equation for the groundwater flow is as follows:

$$\frac{\partial}{\partial x} \left( K_y \frac{\partial H}{\partial x} \right) + W = \mu \frac{\partial H}{\partial t} \quad (x, y) \in \Omega \quad i, j = 1, 2 \quad t \geq 0,$$

where $\mu$ is the specific yield, $H$ is the hydraulic head, $K_y$ is the hydraulic conductivity, $\Omega$ is the simulated area range, and $W$ is the volumetric flux per unit volume.

The governing partial differential equations for the contaminant transport are as follows:

$$\frac{\partial C}{\partial t} = \frac{\partial}{\partial x} \left( D_c \frac{\partial C}{\partial x} \right) - \frac{\partial}{\partial y} \left( u_i C \right) + \frac{R}{\theta} \quad (x, y) \in \Omega \quad i, j = 1, 2 \quad t \geq 0,$$

$$u_i = \frac{K_y}{\theta} \frac{\partial H}{\partial y} \quad i, j = 1, 2,$$

where $\Omega$ is the simulated area range, $\theta$ is the effective porosity of the aquifer medium, $C$ is the contamination concentration, $D_c$ is the dispersion coefficient, $u_i$ is the average linear velocity of the groundwater flow determined by Darcy’s law, and $R$ is the source or sink term.

After establishing the groundwater flow and contaminant transport numerical simulation model, the GMS software was used to solve the simulation model.

In contrast to the actual problem, the hypothetical example had no actual data measurements. Therefore, it was necessary to forward run the contaminant transport simulation model and obtain the contaminant concentration data for the observation wells during each simulation period for use as the data measurements during the identification process. Fig. 3 shows the contaminant plume distributions on day 450 and day 900 of each case. Fig. 4 shows the values measured for each
observation well in each simulation period for the two contaminated cases.

Fig. 3. Contaminant plume distributions: (a), (b) 450 d, 900 d for case one, respectively; (c), (d), 450 d, 900 d for case two, respectively.
Fig. 4. Values measured of the observation wells for two cases: (a) case one; (b) case two.

Surrogate models of the numerical simulation model

A surrogate model with satisfactory accuracy has almost the same input-output relationship as the simulation model. The advantage of applying a surrogate model instead of a simulation model is that the surrogate model is not only simpler to call, but it can reduce a lot of computational load and time required when solving the optimization model (Simpson et al., 2001; Luo and Lu, 2014a Luo et al., 2014b; Jiang et al., 2015).

The horizontal and longitudinal grid numbers and release history corresponding to a maximum number of the GCSs were used as the input variables for the surrogate model (six grid numbers variables and six release intensity variables comprising a total of 12 input variables). The contaminant concentrations in each observation well during each simulation period were treated as the output variables in the surrogate model. Using the Latin hypercube method, the grid numbers and release history of each zone were sampled in their feasible domains. The feasible domain ranges are shown in equations (30). In total, 420 groups were sampled where the grid numbers and release intensities for the 420 groups were used sequentially as inputs for the simulation model. The corresponding outputs comprising the concentrations in all of the observation wells in each period were obtained by running the contaminant transport numerical simulation model. 360 groups of input–output data were selected as the training samples for the surrogate model and 60 groups of input-output data were selected as the testing samples for the surrogate model. The kriging and ELM methods were coded with Matlab, and the Krig-SM and ELM-SM were then established based on the training samples.
The accuracy of the two surrogate models was tested based on three evaluative coefficients comprising the certainty coefficient ($R^2$), root mean square error (RMSR), and mean relative error (MRE).

$R^2$ was calculated as follows.

$$R^2 = 1 - \frac{\sum_{i=1}^{n} (y_i - \hat{y}_i)^2}{\sum_{i=1}^{n} (y_i - \bar{y})^2}$$  \hspace{1cm} (28)

RMSR was calculated as follows.

$$RMSR = \sqrt{\frac{\sum_{i=1}^{n} (y_i - \hat{y}_i)^2}{n}}$$  \hspace{1cm} (29)

MRE was calculated as follows.

$$MRE = \frac{1}{n} \sum_{i=1}^{n} \left| \frac{y_i - \hat{y}_i}{y_i} \right| \times 100\%$$  \hspace{1cm} (30)

In Eqs 28–30, $y_i$ is the output value of the $i$th sample obtained from the contaminant transport numerical simulation model, $\hat{y}_i$ is the output value of the $i$th sample obtained from the surrogate model, and $\bar{y}$ is the average of the output values for $n$ samples obtained from the contaminant transport simulation model. Smaller values for the MRE and RMSR coefficients, as well as values of $R^2$ closer to 1 indicate that the surrogate model is more accurate at simulating the output of the simulation model, thereby demonstrating that the surrogate model can be used to replace the simulation model.

**0-1 MINLPOM**

The optimization model is the basis for the identification of GCS. Therefore, establishing the optimization model is an essential step in the research process (Cristo et al., 2008; Guneshwor et al., 2018). The optimization model comprises three important components: an objective function, decision variables, and constraints.
Based on measured values of groundwater contaminant the optimization model for identifying
the characteristics of GCSs was established. The number, location, and release history of the GCSs
were decision variables in the optimization model. The number of GCSs was treated as integer
variable, which could only be a value of 0 or 1. The location (grid numbers) and release history
were treated as integer variables and continuous variables, respectively. The fitting error between
the measured and simulated contaminant concentrations in the observation wells during each
simulation period was minimized as the objective function. The contaminant transport numerical
simulation model (replace with the surrogate model) was embedded in the optimization model as
an equality constraint to ensure that the optimization model satisfied the contaminant transport law
in a groundwater system during the optimization process. The feasible domains for the number,
location, and release history of GCSs comprised the inequality constraints. The objective function
and constraints constituted the 0-1 MINLPOM used to identify the GCSs. The 0-1 MINLPOM of
two cases are expressed as follow:

\[
\begin{align*}
\min z(\phi, x, y, q) &= \sum_{i=1}^{3} \sum_{k=1}^{t_i} (C_i^k(t) - C_i^k(0))^2 \\
\phi_i &= \begin{cases} 
0 & i = 1, 2, 3 \\
1 & \text{else}
\end{cases} \\
\sum_{i=1}^{3} \phi_i &\leq 3 \\
\phi_1 &\leq \phi_2 \leq \phi_3 \\
19 \leq x_i \leq 70, 18 \leq y_i \leq 55 & i = 1, 2, 3 \quad x_i, y_i \in \mathbb{N}^* \\
\gamma_1 &\leq y_2 \leq y_3 \\
0 \leq q_m &\leq 300 & m = 1, 2, \ldots, 6 \\
C_i^k(t) &= f(q_m)
\end{align*}
\]

(31)

where \( \phi_i \) is the 0-1 integer variable representing whether the current location is GCS, 1
indicates that a real GCS is present in the current location, 0 indicates that no GCS is present in
the current location, \( x_i \) is the horizontal grid numbers, \( y_i \) is the longitudinal grid numbers, \( i \)
represents the \( i \)-th potential GCS, \( q_m \) is the release intensity of the GCSs during each release
period, \( m \) represents the \( m \)-th release intensity variable (each GCS contains two release intensity variables), \( C'_t(t) \) is the simulated concentration of the contaminant at the observation point, and \( C'_t(0) \) is the measured value of the contaminant concentration at the observation point.

The 0-1MINLPOM for identifying the GCSs in each case was exactly the same, except the contaminant concentrations values measured in observation wells corresponding to the two cases were different.

**Results and discussion**

Comparative analysis of the accuracy of the surrogate models

The accuracies of ELM-SM and Krig-SM were compared, and the surrogate model with higher accuracy was selected and embedded in the 0-1 MINLPOM to be called when solving the 0-1 MINLPOM, thereby allowing the number, location (grid numbers), and release history of the GCSs to be identified simultaneously.

In order to test the accuracy of the two surrogate models, 60 groups of input test samples were entered in the simulation model and 60 groups of output samples were obtained. The same 60 groups of input test samples were then entered in the two surrogate models and 60 groups of output samples were obtained from the two surrogate models. The accuracies of the two surrogate models were evaluated by comparing the output samples from the simulation model and those from the two surrogate models. In order to clearly compare the accuracies of the two surrogate models, the contrast histograms were plotted for MRE, RMSR, and \( R^2 \) (Fig. 5).
Fig. 5. Evaluative coefficients of surrogate for the two cases: (a) mean relative error (MRE); (b) Root mean square error (RMSR); (c) Certainty coefficient ($R^2$).

The minimum certainty coefficients with Krig-SM of case one and case two were all above
The maximum and minimum certainty coefficients with ELM-SM were 0.9934 and 0.9255, respectively (show in Fig. 5), which demonstrate that Krig-SM obtained a higher $R^2$ than ELM-SM. Fig. 5 show that the RMSR and MRE values were smaller with Krig-SM than ELM-SM. Thus, Krig-SM obtained a better approximation of the simulation model and it had higher accuracy. Therefore, Krig-SM was embedded in the 0-1 MINLPOM for simultaneously identifying the number, location, and release history of GCSs.

Analysis of GCSs identification results

Krig-SM is embedded in the 0-1 MINLPOM and the GA was used to solve the 0-1 MINLPOM to identify the number, location, and release history of GCSs for two cases. A detailed introduction to GAs was provided by Guo et al. (2018) and Zwickl (2006). The parameter settings for the GA are shown in Table 5.

When solving the 0-1 MINLPOM, it takes about 2200 hours to call the simulation model for 240000 times, while it only takes about 0.21 hours, to call the surrogate model for 240000 times. Thus, using the surrogate model instead of the simulation model for calculation could save 99% of the computational load and the computational time required.

The identification results of GCSs for the two cases are shown in Table 6 and Fig. 6. Case one: The identification results for GCSs are shown in Fig. 6a and Table 6a. A value “1” (shown in Fig. 6a) in the identification results indicates that the number of GCSs in the study area was one. The location and the release intensities of GCSs are shown in Table 6a. The identification accuracy of the number and location was 100%, and the identification accuracy of release history was above 93.59%.
Case two: The identification results for GCSs are shown in Fig. 6b and Table 6b. Those two values of “1” (shown in Fig. 6b) in the identification results indicate that the number of GCSs was two. The location and the release intensities of the two GCSs are shown in Table 6b. The identification accuracy of the number was 100%, and the identification accuracy of location and release history was above 91.67% and 90.14%, respectively.

Table 5. Parameter settings for the genetic algorithm

| Parameter               | Setting         |
|-------------------------|-----------------|
| Population size         | 200             |
| Scaling function        | Rank            |
| Selection function      | Stochastic uniform |
| Mutation function       | Constraint dependent |
| Crossover function      | Scattered       |
| Direction               | Forward         |
| Stall generations       | 1200            |
| Other parameter settings| Default         |

Table 6. Source number, location, and release history results for case two

(a) case one

| Example 1 | Number of sources | S-location (m) | S-Release intensity (g/d) |
|-----------|-------------------|----------------|--------------------------|
|           |                   | X  | Y  | SP1 | SP2 |
| Real value| 1                 | 35 | 52 | 260 | 140 |
| Identified results | 1 | 35 | 52 | 271.73 | 131.03 |
| Relative error | 0.00% | 0.00% | 0.00% | 4.51% | 6.41% |

(b) case two

| Example 2 | Number of sources | S1-location (m) | S2-location (m) | S1-Release intensity (g/d) | S2-Release intensity (g/d) |
|-----------|-------------------|-----------------|-----------------|--------------------------|--------------------------|
|           |                   | X  | Y  | X  | Y  | SP1 | SP2 | SP1 | SP2 |
| Real value| 2                 | 35 | 52 | 41 | 36 | 190 | 120 | 170 | 160 |
| Identified results | 2 | 35 | 55 | 41 | 33 | 207.89 | 110.42 | 154.73 | 175.78 |
| Relative error | 0.00% | 0.00% | 5.77% | 0.00% | 8.33% | 9.41% | 7.98% | 8.98% | 9.86% |
The identification results of GCSs for the two cases showed that the proposed method performed well at simultaneously identifying the number, location, and release history of the GCSs.
Comparing with the 0-1 MINLPOM proposed by previous researchers, the improved 0-1 MINLPOM proposed in this study is more applicable to the identification of GCSs, and can simultaneously identify the number, location, and release history of the GCSs.

In this study, the actual number of GCSs was less than the estimated maximum number of three. If the actual number of GCSs is greater than or equal to the maximum number of GCSs, the solution is increasing the original estimated maximum number of GCSs. Then establishing a new surrogate model and optimization model to identify the information for the GCSs until the number of GCSs identified is less than the estimated maximum number of GCSs (in a feedback correction process).

However, the number of input variables of the numerical simulation model will increase as the estimated maximum number of GCSs increases and the nonlinear relationships will become more complicated. When faced with simulation model with more complex nonlinear relationship, the accuracy of the surrogate model established by the traditional shallow learning method (Kriging and extreme learning machine, etc) may be unsatisfactory. Thus, methods for establishing surrogate model with better accuracy should be studied. Deep learning neural networks and deep reinforcement learning neural networks have strong fitting capabilities for simulation models with complex nonlinear relationships and have great potential to establish surrogate model in future research.

Conclusions

In this study, an improved 0-1 MINLPOM was developed to simultaneously identify the
number, location, and release history of GCSs. Based on the study two conclusions were obtained:

First, the results showed that the surrogate model established using the kriging method was more accurate than the ELM. Thus, Krig-SM can be used instead of the simulation model to embed in the improved 0-1 MINLPOM during solution process. The surrogate model provided a highly accurate approximation of the simulation model, and applying Krig-SM instead of the simulation model in calculation can also save 99% of the computational load and the computational time required.

Second, in previous methods employed for the identification of GCSs, the improved 0-1 MINLPOM was used to identify only the location and release history of the GCSs. In the present study, the number of GCSs was treated as integer variable with a value of 0 or 1, the location was treated as integer variable and release history of GCSs were treated as continuous variables. The number, location, and release history of GCSs were simultaneously identified using the 0-1 improved MINLPOM. The results showed that the identification results were very similar to the true values of the GCSs characteristics. The accuracy of identifying the number of GCSs was 100%, and the accuracies of identifying the locations and release intensities were over 91.67% and 90.14%, respectively. This method can effectively solve the problem of identifying the number of GCSs, as well as simultaneously identifying their locations and release history.

**Data Availability Statement**

All of the data, models, or code generated or used in this study are available from the corresponding author by request.
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Declarations

The authors have no relevant financial or non-financial interests to disclose.

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