Data-Worth Assessment for a Three-Dimensional Optimal Design in Nonlinear Groundwater Systems

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Abstract

Groundwater model predictions are often uncertain due to inherent uncertainties in model input data. Monitored field data are commonly used to assess the performance of a model and reduce its prediction uncertainty. Given the high cost of data collection, it is imperative to identify the minimum number of required observation wells and to define the optimal locations of sampling points in space and depth. This study proposes a design methodology to optimize the number and location of additional observation wells that will effectively measure multiple hydrogeological parameters at different depths. For this purpose, we incorporated Bayesian model averaging and genetic algorithms into a linear data-worth analysis in order to conduct a three-dimensional location search for new sampling locations. We evaluated the methodology by applying it along a heterogeneous coastal aquifer with limited hydrogeological data that is experiencing salt water intrusion (SWI). The aim of the model was to identify the best locations for sampling head and salinity data, while reducing uncertainty when predicting multiple variables of SWI. The resulting optimal locations for new observation wells varied with the defined design constraints. The optimal design (OD) depended on the ratio of the start-up cost of the monitoring program and the installation cost of the first observation well. The proposed methodology can contribute toward reducing the uncertainties associated with predicting multiple variables in a groundwater system.

Introduction

Groundwater models are commonly used in conjunction with field monitoring to assess the physical processes representing subsurface flow and solute transport. Such models simulate the groundwater dynamics in an aquifer by translating its physical, chemical, and biological characteristics into mathematical equations by simplifying assumptions (Holzbecher and Sorek 2006). These equations require data about aquifer characteristics—such as hydraulic properties, geological borders, boundary conditions, and sources and sinks—that will sufficiently aid in understanding groundwater dynamics (Bakalowicz 2005). However, the complexity of subsurface conditions may lead to a paucity in data describing the control parameters; this in turn will result in increased uncertainties with model simulations (El-Fiky 2010). The lack of data coupled with model prediction uncertainty makes it difficult for water resources managers and decision makers to plan a management strategy to secure the quantity and quality of groundwater (Tribbia and Moser 2008; Comte et al. 2016). Therefore, it is imperative to design a monitoring network that would reduce prediction uncertainties in order to improve the protection and management of aquifer systems (Storck et al. 1997). In this context, models can be used as test beds to identify new (optimal) monitoring locations that would increase the reliability of model simulations. This technique is generally referred to as an OD with several reported methods to guide the design of monitoring networks toward reducing uncertainties in model predictions (Rouhani and Hall 1988; Loaiciga 1989; Andricevic and Foufoula-Georgiou 1991; Cieniawski et al. 1995; Wagner 1995; Herrera et al. 2000; Reed et al. 2000; Tiedeman et al. 2003).

A recently developed method by Moore and Doherty (2005), and later extended within the Bayesian context by Christensen and Doherty (2008), evaluated the variance of prediction uncertainty using a linear propagation

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Article impact statement: A new method is developed to identify optimal groundwater sampling locations with minimal observation wells in space and depth.

Received May 2018, accepted October 2018. © 2018, National Ground Water Association.
doi: 10.1111/gwat.12835

NGWA.org Groundwater 1
of uncertainties associated with parameters that are formulated for distributed models. Using this method, an existing calibration data set is augmented by adding new observations. The worth of such an addition (subsequently referred to as data worth [DW]) on reducing model prediction uncertainty is then evaluated. Dausman et al. (2010) applied the DW-based OD on the Henry problem to define the optimal locations of salinity concentration and temperature that would reduce the uncertainty of predicting the displacement of a salt/fresh water interface caused by a change in the inflow rate to the system. Wallis et al. (2014) extended the DW-based OD for selecting multiple observations, and Wöhling et al. (2016) extended it further by using a genetic algorithm (GA) to incorporate multiple new observations of head and/or hydraulic conductivity to decrease the predictive uncertainty. Vilhelmsen and Ferre (2017) carried out yet another extension to simultaneously select multiple new measurements targeting multiple forecasts of interest. Note that the applications of the DW-based OD are largely restricted to a single or multiple observation location(s) in two dimensions. The simultaneous DW-based optimization of monitoring design with measurements in three dimensions has not been reported. Expanding the DW approach into three-dimensional (3D) space is particularly important for monitoring contaminant concentrations and to design a solute transport system.

In practice, groundwater contamination distribution varies spatially and vertically over a geologic domain. 3D OD aids in identifying optimal locations for monitoring contaminant concentration such as salt water intrusion (SWI). 3D models are especially useful when attempting to understand SWI, where the spread of intrusion typically occurs in three dimensions through lateral and vertical displacement (or upconing) of the interface. Neglecting three-dimensionality in geologic input data (i.e., hydraulic conductivity) can result in large uncertainties with regards to model predictions (Werner et al. 2013), influencing both the magnitude and the trend of the intrusion (Kerrou and Renard 2010). It can also lead to the overestimation of the toe penetration length (Lu et al. 2009) and subsequently to the miscalculating the DW of a proposed design when the OD target is to increase the reliability of a model in predicting the future position of the interface. To avoid such situations, ideally hydraulic conductivity values should be collected in all spatial directions in order to reduce prediction uncertainty. However, financial constraints and/or spatial limitations (e.g., in urbanized aquifers) reduce to the ability to directly measure the hydraulic conductivity from deeper parts of an aquifer (Hartmann et al. 2014). Using inverse modeling, hydraulic conductivity values can be estimated through the inverse solution of groundwater flow and/or solute transport equations for the value of an observed dependent variable (or an indirect observation) such as hydraulic head and/or contaminant concentration (Hoeksema and Kitamidis 1984). Dausman et al. (2010) recognized that salinity concentration defines the interface and that measuring salinity is crucial to understanding how the interface moves.

Total cost and utility of measuring head and/or salinity depends on the operation, implementation, and start-up costs of the monitoring plan, types of monitoring equipment, resolution of monitoring data, and data processing costs (Hericks et al. 2017). In practice, nearby existing supply wells are usually used as observation wells to monitor head and/or contaminant concentration data (Sen 2015). Using existing wells will result in avoiding extra costs for drilling new observation wells.

The DW methodology assumes that an initial baseline can be established using historical data and available information about the main hydrogeological characteristics of the aquifer under study (e.g., boundary conditions and source/sinks). The initial baseline is expected to provide insight into the overall water flow system and serve as a test bed to estimate hydraulic heads, conduct model simulations, and calculate the sensitivity of head and predictions to model parameters. However, uncertainties in model predictions are common when attempting to predict beyond the range of available input data, for example, in highly parameterized models with more unknown parameters than observations. In such nonlinear models, the entire range of possible values of observational data should be considered during the OD in order to calculate a wide range for the sensitivities of observations to the model parameters because the actual values of the observation data are unknown prior to collection (Leube et al. 2012). If the model nonlinearity is high, it may result in multiple plausible observation locations.

Several Monte Carlo (MC)-based techniques have been developed to account for model uncertainty. MC techniques can be used to generate a range of measurement values obtained by using different parameter sets (multiple realizations) that are conditioned by the calibration data set (Keating et al. 2010). Compared with other techniques such as the Markov Chain MC method (Harvey and Gorelick 1995), the generalized likelihood uncertainty estimation (Beven and Binley 1992), and calibration-constrained MC methods (Tavakoli et al. 2013), the subspace technique for calibration-constrained MC analysis referred to as the null-space Monte Carlo (NSMC) method (Tonkin and Doherty 2005) requires less computational time for generating a large number of calibration-constrained parameter fields. The NSMC method is best suited for groundwater modeling in highly parameterized systems due to the long model runtime and the large number of required model runs for such systems (Herckensrath et al. 2011). In this method, a set of random parameter realizations is first generated from a probability distribution defined from available prior information, for example, about the hydraulic conductivity. The generated random realizations are then projected onto the null-space and adjusted through model recalibration. This results in a set of calibration-constrained realizations (for details see Tonkin and Doherty 2009). For the OD, calibration-constrained realizations can be ranked and assigned weights according to their goodness of fit with observed data. Using Bayesian model averaging (BMA), model weights are determined via Bayes’ theorem from
the likelihood that the calibration data set is generated from realizations (Hoeting et al. 1999). When applying
the OD, the optimal location of a new observation can be
determined by averaging all possible locations obtained
using calibration-constrained realizations (Freeze et al.
1992). In this manner, parameter and prediction uncer-
tainties are both considered when attempting to find the
optimal locations for new observations.
An ideal DW-based OD should provide flexibility
concerning model dimensionality, allow for any desired
task-oriented formulation, target any measurement type
(direct and indirect), account for various sources of uncer-
tainty (e.g., geologic structure, heterogeneity, boundary
condition, and source/sink) while also ensuring that it is
cost effective. Existing DW-based OD methodologies fall
short of simultaneously providing these criteria for the
design of a monitoring network in a groundwater system.

In this study, we expand the DW-based OD method to
optimize simultaneous measurements of various data types
collected at different depths at a single and multiple spatial
locations, while considering model nonlinearity through a
BMA framework and minimizing costs. The cost-effective
solution involves obtaining adequate hydrogeological
information with a minimum number of observation wells.
In what follows, we describe the theory underpinning the
design methodology followed by the design method that is
applied to a case study that reduces the uncertainty in the
predictions of SWI by determining the optimal location(s)
that would allow for efficiently obtaining data within the
model domain.

Methods and Materials

The proposed methodology involves a BMA frame-
work with a 3D DW-based OD analysis that is imple-
mented to select an optimal observation data set that
would reduce model uncertainty (Figure 1).

BMA Framework

We denote a set of distributed groundwater models
Mk: k = 1, ..., K for predicting flow and/or solute
transport over a geologic domain. Each model is a
probability distribution model comprising the likelihood
function \( P(h_0 | M_k) \) of the observed data \( h_0 \)
and the model parameters \( P_k \) (e.g., hydraulic
conductivity). The posterior predictive distribution of the forecast of
interest \( \Delta \) is determined as a weighted averaged individual
prediction as expressed in Equation 1, where weights can be
determined using Bayes’ theorem on the basis of the
likelihood that the observed data \( h_0 \) are generated using
Equation 2 (Hoeting et al. 1999):

\[
P(\Delta | h_0) = \sum_{k=1}^{m} P(\Delta | h_0, M_k) P(M_k | h_0)
\]  

(1)

\[
P(M_k | h_0) \propto P(h_0 | M_k) P(M_k)
\]  

(2)

where \( P(M_k) \) is a probability mass function over the
model \( M_k \). The use of a defuse prior \( P(M_k) = 1/m \)
ensures that there is no subjective preference for any of
the model \( M_k \) (Wöhling et al. 2015). One can normalize
the weights of the models by applying Bayes theorem as expressed below:

\[
P(M_k | h_0) = \frac{P(h_0 | M_k) P(M_k)}{\sum_{k=1}^{m} P(h_0 | M_k) P(M_k)}
\]  

(3)

where \( P(h_0 | M_k) \) is the likelihood of observing the
calibration data set \( h_0 \) under model \( M_k \). It can be
determined based on its prior parameter distribution using
Equation 4:

\[
P(h_0 | M_k) = \int P(h_0 | M_k, P_k) P(P_k | M_k) dP_k
\]  

(4)

where \( P(P_k | M_k) \) is created by generating random param-
eter fields that meet calibration constraints. We use the
NSMC method to create random parameter fields \( P_k \).
Prior to incorporating the NSMC method, the hydraulic
conductivity field is parameterized by defining a large
number of pilot points \( P_k \) that cover the geologic
domain. The model is then calibrated to estimate the
values of the pilot points. The extent to which a pilot
point parameter can be informed (identified) by the exist-
ing observations can be measured by a singular value
decomposition of the Jacobian matrix that represents the
sensitivity of observations to the pilot point parameters
(for details see Doherty and Hunt 2010). The pilot points
corresponding to the singular values that are larger than
a given user-defined “truncated” value (5.0 \( \times \) 10^{-6}
in this work) span the calibration solution space. These pilot
points are deemed to be estimable on the basis of existing
observations. In contrast, pilot points that contain low or
zero singular values (that span the calibration null-space)
are considered inestimable. Using the NSMC method,
a set of random values is first generated from a prior
probability distribution of hydraulic conductivity fields.
The random values are placed on the pilot points (which
is called a random realization). The generated random
realization is then projected onto the null-space through
differencing the random values and the calibrated pilot
points’ values, and then readding the projected difference
onto the calibrated values. The projected parameter set is
then adjusted through a model recalibration in order to
respect calibration constraints made by existing observa-
tions. The result is a calibrated-constrained realization that
respects both the stochastic variability of the hydraulic
conductivity field as well as the calibration constraints
(for details see Tonkin and Doherty 2005). The posterior
probability distribution of a model prediction is then com-
puted on the basis of the generated calibrate-constrained
realizations.

Linear Model Calibration

We assume that each Bayesian model \( M_k \) is a linear
model that defines a relationship between its parameters

and its predictions using Equation 5 (Doherty 2015):

$$M_k : h = XP + \varepsilon$$

where $h$ denotes a $m \times 1$ vector of head observations comprising the calibration data set that are contaminated with noise $\varepsilon$ (i.e., error in field measurement), $P$ represents a $n \times 1$ vector of model (pilot points) parameters in the conceptual model, and $X$ is the action of model or model sensitivity (or Jacobian matrix). The unknown model parameters can be estimated by minimizing an objective function that is defined based on the sum of the squared-weighted residuals between the model results and (potential) observed data (or model-to-measurement misfit) as shown in Equation 6:

$$\varphi = (h - X\bar{P})^T Q (h - X\bar{P})$$

where $h$ is a $m \times 1$ vector of potential head observations, $\bar{P}$ is the vector comprised of unknown parameters (or parameter estimates), $T$ stands for the matrix transpose operation, and $Q$ is a diagonal matrix with squared observations weights ($\omega_i$) that is defined to be proportional to the inverse of the covariance matrix of the observations noise. The parameter vector minimizing the objective function (in Equation 6) can be determined using Equation 7:

$$\bar{P} = (X^T Q X)^{-1} X^T Q h$$

where $Q$ is the matrix with the squared (calculated) weights of observations. The potential wrongness (or error) of the estimated parameters $\bar{P}$ compared with the true parameter fields $P$ can be evaluated by Equation 8:

$$P - \bar{P} = I P - (X^T Q X)^{-1} X^T Q h$$
where \( I \) is the identity matrix. Let us further assume that \( s \) denotes a true model prediction. Then, the relationship between \( s \) and \( P \) is estimable using Equation 9:

\[
s = y^T P
\]

(9)

where \( y \) is a \( n \times 1 \) vector representing the sensitivity of the predictions to model parameters. If \( \bar{s} \) is a model prediction that is computed from \( \overline{P} \) using Equation 6, then the potential error in the computed prediction can be expressed by Equation 10:

\[
s - \bar{s} = y^T (P - \overline{P})
\]

(10)

However, the true parameter fields \( (P) \) and prediction \( (s) \) are unknown. Therefore, none of the potential wrongness of (or error in) the estimated parameter (in Equation 8) and the computed prediction (in Equation 10) can be calculable. If we assume that \( P \) and \( \varepsilon \) are independent and their covariance matrices are known, then the covariance of the parameter error can be expressed by Equation 11:

\[
C (P - \overline{P}) = (I - R) C (P)(I - R)^T + EC (\varepsilon) E^T
\]

(11)

where \( C(P) \) is a \( n \times n \) parameter covariance matrix representing innate parameter variability, which can be created using a Kriging variogram that is defined to represent the spatial distribution of the hydraulic conductivity fields, \( C(\varepsilon) \) is a \( m \times m \) matrix of measurement noise \( \varepsilon \), \( R = (X^T X)^{-1} X^T Q X \) is a so-called resolution matrix that describes the relationship between the estimated and true parameters, and \( E = (X^T Q X)^{-1} X^T Q \).

**Prediction Uncertainty Variance**

Combining Equations 10 and 11 leads to the expression of the variance of model prediction uncertainty as shown in Equation 12 (Christensen and Doherty 2008):

\[
\sigma_{\delta}^2 = y^T C (P) y - y^T C (P) X^T \left[ X C (P) X^T + C (\varepsilon) \right]^{-1} X C (P) y
\]

(12)

The first term on the right-hand side of the equation is the precalibration uncertainty for the predictions. The second term shows the amount that the prediction uncertainty is reduced by calibrating the model using measurements comprising the calibration data set.

**DW Analysis (Prediction of Single Variable)**

The variance of the model prediction uncertainty does not account for the values of parameters, measurements, or prediction. Instead, it comprises only the sensitivity of the model’s observations and predictions to the parameters, which are included in the \( X \) and \( y \) matrices, respectively. For the purpose of an OD, the change in the prediction uncertainty can be evaluated when a new observation (set) is added to the existing calibration data set. In general, when adding an observation, it reduces the model prediction uncertainty, while increasing the DW that this observation has on the calibration data set. The DW is measured using Equation 13 (More details can be found in Vilhelmsen and Ferre 2017):

\[
DW = \frac{\sigma_{dec}^2}{\sigma_{base}^2}
\]

(13)

where the \( \sigma_{dec}^2 \) is the decrease in the prediction uncertainty when adding a new observation point, and \( \sigma_{base}^2 \) is the predictive uncertainty pertaining to the existing calibration dataset. The DW is represented by a value ranging between 0 and 1; it reflects the impact of additional observations on the base predictive uncertainty (reduction). For example, the DW of 1 denotes that the prediction uncertainty is completely diminished by adding a new observation, whereas a DW of 0 indicates that a new observation does not reduce the prediction uncertainty.

**Value Index (VI) Analysis (Prediction of Multiple Variables)**

In practice, multiple variables are often of interest to simulate using a groundwater model, for example, predictions of stream flow, flow velocity, or contaminant migration. The DW for an observational well can be further evaluated if the target of an OD is to minimize the uncertainty associated with predicting multiple variables in a given groundwater system. For this purpose, the DW is first calculated for each prediction variable. A weight is then defined for each variable and subsequently applied to each DW. The weighted DWs are then combined into a single VI that indicates the value of monitoring each observational well according to the priority of the prediction variables (Equation 14) (see Vilhelmsen and Ferre (2017) for more details):

\[
VI_j = \sum_{i=1}^{n} w_i DW_{i,j}
\]

(14)

where \( j \) corresponds to an observation set, \( n \) stands for the number of prediction variables, \( w_i \) is the weight of the \( i \)th prediction variable, and \( DW_{i,j} \) is the DW of \( j \)th observation set to \( i \)th variable. Weighting prediction variables are a subjective choice of the modeler/manager and can be based on various factors such as economic worth of making predictions or prioritizing a prediction when making management plans.

**DW-Based 3D OD**

As outlined in Figure 1, the methodology consists of five interrelated steps outlined below.

(A) For each model \( M_k \), a series of matrices are formed to represent the following: (1) the sensitivity of the existing calibration data set to model parameters \((X_{old})\); (2) the sensitivity of predictions to model parameters \((Y)\); (3) the innate parameter variability \(C(p)\); (4) the measurement noise \(C(\varepsilon)\); and (5) the Jacobian matrix \(X\).
that consists of \( Y \) and \( X_{\text{old}} \). These matrices are used to calculate the base predictive uncertainty. In this work, the PREDUNC (PREdiction UNCertainty) program in the PEST suites of utilities was used to calculate the base predictive uncertainty (Doherty 2015).

(B) A set of arbitrary 3D locations for potential (yet to be collected) observations is specified and used as input to the DW analysis. In order to perform a 3D design, we create an \( X_{\text{new}} \) matrix containing the spatial locations of observation wells, where each row of the matrix represents a single observation location. Each observation location (i.e., an element of the \( X_{\text{new}} \) matrix) contains a corresponding sensitivity matrix \( L \) with \( m \) rows and \( l \) columns, where \( m \) is the maximum number of sampling (or measurement) depths, and \( l \) is the number of model parameters. The \( L \) matrix comprises the sensitivity of a single observation location to all model parameters with respect to the depth at which the measurements were taken. Each row of the \( L \) matrix contains the sensitivity of a measurement depth to all model parameters. Each column includes the sensitivity of a certain parameter to all measurement depths. All new measurements pertaining to the \( L \) matrices are added to the \( X_{\text{old}} \), and then the sensitivities to the parameters are estimated by calculating the Jacobian matrix \( X \) using PEST (Doherty 2015). The new measurements are then detached from the \( X_{\text{old}} \).

Note that for temporal monitoring design, the \( X_{\text{new}} \) matrix can be simply expanded by additional \( L \) matrices corresponding to different model stress periods. For this purpose, an \( L_t \) matrix is generated for each model stress period.

(C) The third step is to find an optimal set of \( n \) potential observation wells. Combinations of \( n \) rows are selected from the \( X_{\text{new}} \) matrix. The GA is then used to select multiple rows corresponding to the combinations in the \( X_{\text{new}} \) matrix (for details about GA see Wöhling et al. 2016). A \( p \times 1 \) vector of randomly sampled \( N \) rows is then generated, where \( p \) is a user-defined population size (\( p = 50 \) in this work). This vector forms the initial population of designs (\( i = 1 \)). Each element of this vector contains a random combination of \( n \) rows in the \( X_{\text{new}} \). The \( L \) matrices pertaining to these \( n \) rows are added to the \( X_{\text{old}} \), and subsequently the value of information (VI) of combinations are evaluated using PREDUNC5 (Doherty 2015). Note that each combination comprises \( n \) number of new observation locations, and each new observation contains measurements at multiple depths. Therefore, the \( X_{\text{old}} \) matrix is expanded by adding the maximum \( n \times m \) measurements. In the next step, a new population of design (size \( N \)) is generated by applying the standard GA selection schemes, that is, selection, mutation, and crossover (Wöhling et al. 2016). In the present study, we retained 40% of the population for the next generation (\( i = i + 1 \), muted 5% of population (which was allowed to increase if the population was too uniform), and allowed a 15% chance of selecting outside of the admissible location (similar to Vilhelmsen and Ferre 2017 and Wöhling et al. 2016). The new proposal design is then compared with the previously generated designs.

We ended the loop when the highest VI and the proposed designs were similar in the last 10 subsequent trials. We allowed a maximum of 1000 trials to repopulate the designs in order to achieve convergence. The converged design is recorded as the proposed design of size \( N \) (i.e., \( D_N \)) for model \( M_k \).

(D) The \( X_{\text{old}} \) matrix is amplified by the proposed design (\( D_{N_k} \)) of size \( N \) obtained using one of the models \( M_k \). The reduction in prediction uncertainty is then evaluated in all models \( M_k \) using the amplified \( X_{\text{old}} \) matrix, and then the VI is calculated for each model. The estimated VIs are multiplied by the models weights (estimated through the BMA), and then they are averaged. The averaged VI is the impact of the proposed design (\( D_{N_k} \)) on the prediction uncertainty reduction with respect to model nonlinearity.

The proposed design of size \( N \) is recreated for all models \( M_k \) \((k < K)\). Afterward, the averaged VI is calculated. Among all proposed designs for models \( M_k \) \((k = 1,2, \ldots K)\), the design that has the highest averaged VI is selected as the optimal (or best) design for placing \( N \) observations with respect to the uncertainty of the model input.

(E) After completing the previous steps, one can conduct a cost-effective analysis to determine the optimal number of new observation wells (i.e., optimal size of design). A design is considered cost effective when the reduction of prediction uncertainty (i.e., increase in DW) outweighs its cost. The cost is herein defined by the number of required observation wells. The cost-effective analysis is accomplished by calculating the cost (Po) of each design of size \( N \), and the design that provides the most information at the smallest operation cost is considered the OD. This is estimated by the ratio: DW/Po. The design that has the highest DW/Po ratio is deemed the most cost-effective design.

Application

Description of Study Area

The performance of the proposed method was evaluated by simulating flow and solute transport in an actual aquifer system. Located along the Eastern Mediterranean (Figure 2), the pilot aquifer (covering an area of approximately 42 km²) underlies Beirut city (Lebanon) and its suburbs. The study area has a 16.5 km of shorelines encompassing rocky beaches, sandy shores, and cliffs. It is bounded by several faults to the east and south and partly by an intermittent river to the south.

Hydrogeology

The hydrogeology of the pilot aquifer consists of Cretaceous karst limestone overlaid by Upper Tertiary and Quaternary unconsolidated deposits (Peltekian 1980). The approximately 700-m-thick, fractured Cenomanian-Quaternary system is dominated by hard and compact limestone and dolomite interbedded with chert, and intercalations of marl (Khair 1992). According to the available geologic cross sections (Figure 3), the rock
sequence of Cenomanian-Turonian age (Walley 1997) can be divided into three subunits, as follows: (1) the Afqa-Dolomite member that consists of crystalline, dolomitic, marly dolomitic, and reefal limestone; (2) the Aaqoura member comprises a sequence of thinly bedded limestone, marly limestone, dolomite, and marly dolomite strata; and (3) the Mnaitra member that is composed of thick and compact limestone and fossiliferous strata with several chert bands and nodules across different horizons. These are also known as the C4a, C4b, and C4c formations, respectively (Saint Marc 1974). The aquifer can be divided into seven zones to describe its geologic surface and subsurface (Figure 4 and Table 1). The upper geologic layer consists of a mix of C4c and Quaternary formations in the north (zone 1), and a mix of the C4a and Quaternary formations in the middle and to the south (zone 2). The middle layer contains the C4a formation (aquifer) to the east (zone 3), the C4c formation (aquifer) in the north (zone 4), and the C4b formation (aquitard) along the western coastline (zone 5). The lowest geologic layer with a thickness of approximately 250 m comprises of the C4c formation (aquifer) in the north (zone 4), the deep C4c formation (aquifer) along the western coastline (zone 6), and the deep C4b formation (aquitard) to the east (zone 7) (Table 1).

The Cenomanian formation and Cenomanian-Quaternary systems are permeable with a specific yield of 0.03 and 0.15, respectively (United Nations Development Programme 1970). The infiltration rates are high in the quaternary deposits (Khair 1992). The freshwater influx to the aquifer in the year 1969 was primarily through a reportedly high recharge, equivalent to 20% to 30% of the precipitation (21% reported by United Nations Development Programme 1970; 27% reported by Khair et al. 1994; and 30% reported by Uhayl 1971). The increase in urbanization since 1969 has nevertheless decreased the recharge potential to near nil by 2018 (Safi et al. 2018).

Statement of Problem and Needs for a Monitoring Plan

The upper part of the pilot aquifer is highly vulnerable to SWI with many locations already experiencing high salinity because of groundwater overexploitation (Rachid et al. 2017; Safi et al. 2018). This limits the fresh water resources available in the aquifer and is pushing authorities to consider alternatives such as tapping the deeper parts of the aquifer. In this context, groundwater modeling can guide decision makers toward sustainable abstraction without accelerating SWI. Moreover, it can help protect the deeper parts, where the lack of subsurface characterization will inevitably increase uncertainties associated model predictions. Therefore, it is imperative to design a monitoring network with optimal locations to constrain/reduce model uncertainties.

To design a monitoring network for the pilot aquifer requires simulating and understanding the dynamics of SWI in response to future groundwater abstractions from the deep parts of aquifer. Hence, emulating realistic future conditions of the pilot aquifer is an important step in the OD analysis. In the current application, a scenario was defined whereby it was assumes the authorities will start extracting groundwater from the freshwater resources in the deep aquifer (zone 6) starting in March 2018. Extraction was assumed to occur through pumping from 50 wells at 200 m$^3$/d (Figure 5). In this scenario, the bottom elevation of the pumping wells reaches to depths
of 360 m below mean sea level (BSL) (in the middle of layer 3 in zone 6). A groundwater model was then used to simulate the future SWI in response to the groundwater exploitation from the upper and deeper parts of aquifer. The objective of the model was to find the best locations that will provide the needed information for model prediction of: (1) the displacement of the salt/fresh water interface in zone 6 caused by groundwater abstraction in the entire aquifer (prediction variable 1) and (2) the increase in the salinity concentration in the newly installed pumping wells in zone 6, which will be caused by the landward displacement of the interface. With regards to the latter, salinity concentrations were predicted at two points (A and B) specified in front of the pumping wells. Points A and B were located at depths of 360 m (BSL) (similar to the bottom elevation of the pumping wells in the scenario) (Figure 4). Salinity prediction at point A was denoted as prediction variable 2, and salinity prediction at point B was referred to as prediction variable 3.

Model Setup

SEAWAT code (Guo and Langevin 2002) was used to simulate the salinity migration in the pilot aquifer and to perform the OD in zone 6. SEAWAT is a variable density groundwater flow modeling code, representing flow and solute transport processes that are solved jointly by Modular Three-Dimensional Finite-Difference Groundwater Flow Model and Model Transport in 3 Dimensional. The criteria considered in the code selection process centered on its ability to: (1) simulate the 3D nature of the vertical and lateral encroachment of salinity in confined and unconfined aquifers; (2) characterize various types of time-dependent boundary conditions; (3) simulate steady-state and long-term transient flow and solute transport with the least numerical instability; (4) link to an inversion code (such as PEST) to quantify uncertainties; and (5) contain reasonable computational resources.

Our model comprised a transient stress period of 50 years subdivided into 50 subperiods of 1-year duration, extending from March 1969 up to March 2019 (the future state). The first stress period (March 1969) was used as the calibration-time period because that period had the most information with regards to head observations within the aquifer, with a total of 35 head observations tapping into the upper geologic layer (Figure 3).

In the setup of the SEAWAT model, the sea boundary to the north and west was specified as a constant head and concentration boundary condition with an average salinity level of 35 g/L. The eastern boundary was assumed to be a no-flow boundary due to aquitards and Faults 1 and 3 in the vicinity. The horizontal discretization contained 4251 active cells designed in 115 rows and 75 columns, where each grid cell represents a square of 100 by 100 m (Figure 6). Groundwater abstraction for the upper aquifer

Figure 3. Geologic cross sections CC’ and EE’.
Table 1  
Geologic Formations in the Pilot Aquifer with Corresponding Hydraulic Conductivity Ranges (Ukayli 1971; Peltekian 1980)

<table>
<thead>
<tr>
<th>Model Domain</th>
<th>Thickness (m)</th>
<th>Geologic Formation</th>
<th>Type</th>
<th>Log Hydraulic Conductivity (m/day)</th>
<th>Zone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Layer 1</td>
<td>~100 to 150</td>
<td>C4c-Quaternary</td>
<td>Aquifer</td>
<td>−1.69 to 2.69</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>C4a-Quaternary</td>
<td>Aquifer</td>
<td>−1.26 to 2.69</td>
<td>2</td>
</tr>
<tr>
<td>Layer 2</td>
<td>~150</td>
<td>C4a</td>
<td>Aquifer</td>
<td>−1.3 to 2.69</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>C4c</td>
<td>Aquifer</td>
<td>−1.69 to 2.69</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>C4b</td>
<td>Aquitard</td>
<td>−5 to −3</td>
<td>5</td>
</tr>
<tr>
<td>Layer 3</td>
<td>~250</td>
<td>C4c</td>
<td>Aquifer</td>
<td>−1.69 to 2.69</td>
<td>4¹</td>
</tr>
<tr>
<td></td>
<td></td>
<td>C4c</td>
<td>Aquifer</td>
<td>−1.69 to 2.69</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>C4b</td>
<td>Aquitard</td>
<td>−5 to −3</td>
<td>7</td>
</tr>
</tbody>
</table>

¹The same zone used to characterize the C4c in the second and third layers.

The freshwater influx to the aquifer in the year 1969 was assumed to be primarily through the recharge equivalent of 30% of precipitation (Ukayli 1971). This rate linearly decreased to zero for 2019 due to increase in impervious pavements over time (Safi et al. 2018). The lateral flow was assumed to be zero due to faults and aquitards in the vicinity of the aquifer.

**Pilot Points Parameterization**

The geology of the aquifer is poorly characterized due to limited information about its characteristics, but its hydraulic conductivity can vary considerably over small distances (Safi et al. 2018). Therefore, the use of only a few zones to represent the hydraulic conductivity field may not accurately represent the flow conditions. To remedy this problem, a set of pilot points was assigned to each geologic zone to represent the spatial variability in the hydraulic conductivity. Altogether, a total of 564 pilot points was defined for the entire model domain (i.e., the vector \( P_k \) in the BMA). We used an exponential Kriging variogram to spatially interpolate the pilot points values over the geologic domain. The value of 500 m (i.e., the separation distance between two pilot points) was defined for the range of the variogram (Doherty and Hunt 2010). Using PEST, the pilot points parameterization approach was used in conjunction with SVD and Tikhonov regularization to estimate the values of the pilot points using the existing head observations (Doherty 2015).

The calibrated pilot point model was then used to calculate the sensitivities of the observations and model prediction variables (1, 2, and 3) to the pilot point values. According to the sensitivity results, the pilot points...
parameters were categorized into three subsets: subset (1) contained the pilot points that spanned the solution space or were correlated with the pilot points that lied in the solution space (zones 1 to 4); subset (2) comprised the null-space located pilot points that inform the predictions (zone 6); and subset (3) included the pilot points that lied in the null-space and did not inform the predictions (zones 5 and 7).

**Bayesian Model Averaging**

A random realization of the hydraulic conductivity field was generated for each subset on the basis of the prior probability distribution of the subset’s hydraulic conductivity. The generated random realizations of all subsets were then combined into one realization that contained the full set of the pilot points parameters. The NSMC method was then used to calculate a calibrated-constrained realization on the basis of the combined generated random realization. This process was repeated several times to create multiple calibrated-constrained realizations.

The prior probability distributions were defined according to the available geologic knowledge for each subset, as follows: subset (1) used a probability distribution that was estimated based on the results of the calibrated pilot points model. Subset (2) used a fuzzy theory set (Zadeh 1965) to create a fuzzy probability distribution for the mean log hydraulic conductivity value. For this subset, a trapezoidal distribution was used to define the membership functions of the log mean hydraulic conductivity values for the pilot points (for details about fuzzy set theory see Bardossy et al. 1990). Accordingly, the membership function was set to 1 for the range of log hydraulic conductivities between 1.5 and 2 m/day, (lower and upper support limits). The membership value was set to zero for the lower and upper limits of the log mean hydraulic conductivity values, which were defined as 0.31 and 2.69 m/day, respectively. Subset (3) did not use any probability distribution and the hydraulic conductivity value was defined as constant for all pilot points.

Our BMA application encompassed the generation of only 10 calibration-constrained realizations due to the large run-time of the model. The Bayesian models ($M_k$) constructed on the basis of the realizations were then used to simulate SWI for the near future. For the purpose of model averaging, similar weights were assigned to the models ($M_k$) because all had almost the same sensitivity to the existing observed data after the model calibration.

**OD of New Observation Wells**

The OD analysis involved defining the best locations for a set of new observation wells in zone 6. The design assumed that the observation wells were available every 150 m in zone 6 (i.e., spatial location), and a total of 192 potential locations were specified in that zone (Figure 4). Each observation well was assumed to contain two measurements of field data, namely: (1) the head and salinity concentration at a depth of 300 m BSL and (2) the head and salinity concentration at a depth of 360 m BSL. Hence, the measurements vary spatially and vertically over zone 6 (i.e., three dimensions). The values of these measurements were extracted from the last stress period (i.e., the year 2019) of the Bayesian models and assigned to the observation wells.

The OD framework was then applied to define: (1) the OD to make all predictions under different prediction weights when having one observation well where head and salinity are measured and (2) the OD for multiple locations (i.e., $N = 1, 2, \ldots, 5$ observation wells) whereby measurements of head and salinity are taken at both single and multiple depths to reduce the uncertainty with the prediction of the salt/fresh water interface. Several scenarios were considered to define a variation in the predictions’ weights, listed as follows: scenario a: $w_1 = w_2 = w_3 = 0.33$; scenario b: $w_1 = 0.6$, $w_2 = 0.3$, $w_3 = 0.1$; scenario c: $w_1 = 0.6$, $w_2 = 0.1$, $w_3 = 0.3$; scenario d: $w_1 = 0.8$, $w_2 = w_3 = 0.1$; and scenario e: $w_1 = 1$, $w_2 = w_3 = 0$; where $wi$ is the weight assigned to the previously defined prediction variables ($i = 1, 2, \ldots, 3$).

Finally, a cost-effective analysis was made to find the optimal size of the design, involving new observations, $N = 1, 2, \ldots, 5$. It was assumed that the scaled start-up cost ($Ps$) of a monitoring project varies between 0 and 1, regardless of the number of planned observation wells. The cost of implementing the first observation well ($P1$) was assumed to vary between 0% and 100% of the start-up cost. Variations between start-up costs and implementation costs were considered using a triangular distribution. The cost of implementing additional observation wells ($P_i$) was assumed to be constant for all observation wells.

**Figure 5. Potential new observations locations for monitoring head and salinity in zone 6, along with the locations of new pumping wells and model prediction points A and B at the depth of 360 m BSL.—colored contours are the average of log hydraulic conductivity between all stochastic models—x and y axes are in units of meters.**
costs are captured by the P1/Ps ratio. The implementation cost decreases by 0, 10, 20, and 30% with any additional observation well (P1+). This variation is captured by P1+/P1. A set of random values was then generated based on the aforementioned cost criteria for implementing a given monitoring project. A uniform distribution was used to randomize the costs within their defined ranges. The reduction in predictive uncertainty in relation to the cost of sampling the head and salinity was evaluated by dividing the DW by the cost of the project implementation for simultaneous observation wells (i.e., N = 1, 2, ..., 5).

Results and Discussion

Bayesian Models

Calibration-constrained realizations were generated for the hydraulic conductivity field. The model-to-measurement misfit ranged from 251 to 258 m² with an average residual absolute error of approximately 2 m across all of the stochastic models Mk. Figure 7 shows the calibration-constrained log hydraulic conductivity fields along with the position of the salt/fresh water interface in zone 6 after a 50-year simulation based on the 10 generated stochastic models. The stochastic results are only shown for zone 6, where the OD was performed. The uncertainty with the estimated log hydraulic conductivity had a noticeable impact on the computation of the position of the interface. In four of the models (models M1, M5, M6, and M10), the log hydraulic conductivity values were low along the coastline, and therefore the landward displacement of the interface caused by groundwater abstraction was small as compared with that in other models that contained large hydraulic conductivity values. In the other models, it was apparent that the high hydraulic conductivity values tended to exacerbate SWI, subsequently decreasing the depth to the interface. Since the large uncertainties in the estimated hydraulic conductivities of zones 6 increased the uncertainties in the model prediction, it was necessary to quantify the prediction uncertainty.

We used the range of the simulated values obtained using the generated calibration-constrained realizations of the hydraulic conductivity to quantify the uncertainty in predicting the displacement of the salt/fresh water interface after the 50-year simulation. The histogram of the log hydraulic conductivity values approached an almost or a near normal distribution that ranged from 0.85 to 2.33 m/day (Figure 8a). The distribution was negatively skewed due to the limitations imposed by the low hydraulic conductivity values through a trapezoidal membership function during the randomization process. Figure 8b shows the histogram of the predicted values for the displacement of the interface where the transitional mixing zone was limited to 1 g/L (threshold for drinking water). The predicted values are shown as percent displacement of the interface from its initial position with respect to the coastline, which involved calculating the percent increase in the volume of salinity concentration of greater than 1 g/L in freshwater due to groundwater abstraction. The histogram for the model prediction exhibited a large level of uncertainty in the predicted percent change of the position of the interface, ranging from 40% to 70% (Figure 8b). The prediction histogram had a shape similar to that of the log hydraulic conductivity value (Figure 8a vs. Figure 8b). This underlines the
Figure 7. Log hydraulic conductivity distribution in zone 6 using 10 stochastic models $M_k$: $k = 1, 2 \ldots 10$; along with the position of the interface with 75% (black), 50% (gray), and 25% (white) of sea water concentration (35 g/L) after 50 years simulation.

importance of estimating or knowing the hydraulic conductivity in the deep parts of a coastal aquifer to compute the displacement of the interface.

**OD for Measurements with Single Depth at a Single Observational Well**

The OD analysis determined if the uncertainty in the model predictions would be affected by adding measurements of head and salinity obtained from a single depth at a single observational well in zone 6. The DW of these measurements was found to be sensitive to both the spatial location of the observational well and to the depth at which the measurements were taken (Figure 9). Head and salinity measurements made near points A and B were found to be more effective at informing model predictions of salinity levels at these points as compared with measurements made at other locations (Figure 9b and Figure 9c). The DW increased slightly (~20% more) when the head and salinity measurement depth increased by 20% (Figure 9b vs. Figure 9e and Figure 9c vs. Figure 9f). Measurements acquired at depths shallower than points A and B (e.g.,
300 m BSL) informed the predictions more effectively (i.e., variables 2 and 3) when they were sampled from an observational well located on the seaward side of these points (Figure 9b and Figure 9c). Conversely, measurements taken at the same depths as points A and B (e.g., 360 m BSL) had more impact on reducing the predictions’ uncertainties when they were sampled from an observational well located on the landward side of these points (Figure 9e and Figure 9f).

The observation wells that were located parallel to the coastline and within the transition zone predicted better the displacement of the salt/fresh water interface (i.e., prediction variable 1) (Figure 9a and Figure 9d). A single observational well that had the highest impact on the uncertainty reduction of prediction variable 1 was located approximately 500 m from the coastline if the measurements were taken at a depth of 300 m BSL (Figure 9a). When the measurement depth increased by 60 m, an observational well that was located 200 m further landward provided the most information for predicting variable 1 (Figure 9a vs. Figure 9d). However, adding a single observational well had less impact on reducing the uncertainty associated with predicting variable 1 as compared with variables 2 and 3. Our findings showed the importance of considering the three dimensionalities in an OD when predicting SWI.

**OD for Measurements with Multiple Depths at a Single Observational Well**

Figure 10 shows the contoured VI averaged over the 10 models for the three prediction variables (the 3D displacement of the interface and the salinity levels at points A and B), given a set of specific weight distributions assigned to these three variables. Different locations for an observational well were specified according to the different weights of the prediction variables. The optimal location of a single observational well lied close to the points A and B when all prediction variables were assigned a similar weight (Figure 10a). The DW of a single observational well was twice as high when predicting salinity levels at points A and B than when predicting the interface displacement (Figure 10a vs. Figure 10a and c). Although increasing the weight assigned to predicting variable 1 (w1) twofold increased the VI in the observational wells located within points A and B, the best location for a single observational well was found to be still close to these points (Figure 10b and c). An eightfold increase in w1 as compared to w2 and w3 shifted the best location approximately 500 m upward (Figure 10d). It was found that it was better to collect head and salinity data in the middle of zone 6 within points A and B; that would provide a more accurate prediction of the interface displacement (variable 1) (Figure 10e) because most models showed the highest intrusion occurring along the middle of the coastline (Figure 7). Compared with observational wells located elsewhere, the observational wells located in the middle of zone 6 sensed more concentration changes as the interface approached landward. Our results suggest that designing the location of only one observational well did not noticeably reduce the uncertainty in the prediction of the interface displacement (i.e., variable 1). Moreover, the uncertainty with predicting salinity levels at points A and B appears to approach nil as the number of new observational wells reaches the number of prediction variables (i.e., two observation wells for two point-source prediction variables). In contrast, more observational wells provided substantial information on estimating the interface displacement.

**OD for Measurements with Multiple Depths at Multiple Observational Wells**

The performance of the methodology was examined for a larger number of observations in the case where the weight was given solely to the model prediction of the 3D displacement of the interface (i.e., prediction variable 1). The optimal locations for $N = 2, \ldots$ and 5 simultaneous new observations (i.e., proposed design) were specified separately for each of the 10 models $M_k$. Altogether, 10 designs were proposed for every N. Figure 11 shows the effectiveness of each of the proposed designs in reducing the prediction uncertainty when considering the impact of model nonlinearity. Boxplots were used as a means of comparing the proposed designs.
across the 10 models. In these plots, the averaged DW using all models is shown for each design along with the estimated DW for that model. Each boxplot contains 10 estimated DWs and the highest DW in each boxplot corresponds to the original model \( M_k \) for which the design \( D_k \) was proposed.

The results suggest that the best proposed design having \( N=2 \) observations corresponded to model \( M_5 \) (Figure 11a). The corresponding design (\( D_5 \)) had the highest mean DW and the smallest variance compared with the designs obtained from other models, suggesting that the uncertainty with regards to the hydraulic conductivity values did not significantly affect identifying the optimal locations when \( N \) was equal to 2 observations. For the designs with 3 observations, the highest mean estimate of DW (of 0.72) corresponded to \( D_1, D_6, D_7, \) and \( D_{10} \).
Figure 10. Black solid triangle represents the OD location for a single observation with measurement of head and salinity at multiple depths for multiple prediction variables—(a) \( w_1 = w_2 = w_3 = 0.33 \); (b) \( w_1 = 0.6, w_2 = 0.3, w_3 = 0.1 \); (c) \( w_1 = 0.6, w_2 = 0.1, w_3 = 0.3 \); (d) \( w_1 = 0.8, w_2 = w_3 = 0.1 \); and (e) \( w_1 = 1, w_2 = w_3 = 0 \). \( w_i \) is prediction weight—colored contours are VI that was averaged over the models \( M_k \) locations of points A and B are shown in black rectangles.

(NGWA.org A. Safi et al. Groundwater)
Figure 11. Estimated DW of proposed designs $D_k$ from models $M_k$ ($k = 1, 2 \ldots 10$) for $N = 2, 3, 4, \text{ and } 5$ observation wells in plots a to d respectively: x axis corresponds to a proposed design specified using a model $M_k$, and y axis is the estimated DW when applying a proposed design on all models—Black dot is the mean estimate of the DW, and red line is the median of the estimated DWs for each design using all models.

decreases by increasing the number of observations (Figure 11), the OD size varies according to the cost criteria. The most effective criterion was found to be $P_1/P_s$ (the cost of implementing the first observation according to the cost at start-up). An inspection of the cost-effective results shows that the optimal size of a design should include a maximum of two observations when the cost of implementing the first observation is more than 50% of the start-up cost of the monitoring project (Figure 13c and d). Under this condition, the optimal size can be increased by decreasing the cost of implementing an additional observation to <80% of the cost of operating the first observation (i.e., $P_1^+/P_1$) (Figure 13a and b). With $P_1/P_s < 30\%$, the optimal size can be increased up to 4 observations if $P_1^+/P_1$ is <0.8 (Figure 13a). The implementation of 5 observations seems to be the most cost effective only if the cost of implementing the first observation is much lower than the start-up cost, which may not be plausible.

Conclusion

In this study, we extended an existing linear DW method that optimizes the process of locating multiple new observational locations (yet to be collected) in order to reduce the uncertainty in predicting multiple variables in a groundwater system. Compared with previous studies that used two-dimensional locations for the observations, our method also optimizes simultaneous for measurements occurring at different depths at a single or multiple locations (i.e., three dimensions) at a minimum cost. We also suggested the use of BMA, which was used to define weights for each Bayesian model that contains a set of stochastic parameters. The capability to produce the calibration dataset (also considered as a model prediction) by the stochastic parameters was used to calculate the weight of each Bayesian model. The final outcome of the OD was a set of proposed locations for an observational set that accounts for the nonlinearity of the model.

We applied the proposed methodology on a pilot heterogeneous coastal aquifer that lacks hydrogeological information for its deep geologic layers. The target of the design was to find the best locations for placing 1, 2, 3, 4, or 5 new observations that could contribute to the reduction of the prediction uncertainties. Two types of prediction were used as the optimization targets: capturing the increase in salinity at two points located in the deep part of the aquifer and the displacement of the interface caused by groundwater abstraction. The types of observations that were accounted for included head and salinity at different locations in three dimensions. The following findings were deduced from the OD results of our case study:

• It is important to sample data at different depths and locations if the target of a monitoring design is to predict a solute transport over a 3D geologic domain.
Figure 12. OD for \( N = 1, \ldots, 5 \) new observations with measurement of salinity at multiple depths for predicting the displacement of the salt/water interface. \( \blacktriangle \) is a proposed location for a new observation. The colored contours represent the DW for a single observation with measurement at multiple depths.

- Model nonlinearity has a slight impact on proposing a single location for an observational well with head and salinity measurements. This impact increases by increasing the design size.
- The DW of an observational well with head and salinity measurements depends primarily on its spatial proximity to the coastline, while the depth of measurement is secondary but also important. The proposed spatial location for an observational well moves toward the coastline when the depth of measurement decreases.
- When the depth of measurement is less than the depth of a point at which the predicted salinity concentration is required, a proposed observation should be located adjacent to that point and seaward (i.e., in the direction of the sea). Conversely, when the depth of the measurement is equal to or greater than the depth of the point, the observation should be located adjacent to the point and landward (i.e., further inland).
- To reduce uncertainty with the future interface, observational wells should be located close to the coastline if the measurements to be taken are located at shallow depths. Conversely, observational wells can be located farther from the coast as measurement depth increases.
- The proposed spatial locations for (head and salinity) observations change with the design size. The locations become more similar when the number of planned additional observations increases.
- The optimal size for the monitoring plan depends mostly on the ratio between the start-up cost of the monitoring project and the cost of drilling the first observation well, while the implementation cost of additional observational wells is secondary but also important.
- For the current application, a maximum of two observation wells is needed to obtain a cost-effective monitoring plan if the cost of implementing the first observation, and the monitoring data are more than 50% of the start-up cost of the monitoring project. If the cost of implementing the first observation is equal or greater than the start-up cost, only one observation is recommended for a cost-effective design. Monitoring data from multiple observation wells are the most cost-effective if the start-up cost is much more than the cost of implementing the first observation (e.g., using nearby pumping wells as observation wells).

This study stresses that the effectiveness of the proposed methodology to secure the optimal results hinges on properly weighing the stochastic models when nonlinearity is high. This underlines the importance of a priori knowledge of the system, before designing a monitoring network to produce an effective and successful model calibration. While an increase in the number of planned observations can reduce the prediction uncertainties during the model calibration stage, the cost-effectiveness of a monitoring design was found to be mostly contingent on the cost of operating the first observational location. The results of this study can be used for future field-studies to guide adaptations and implementing sampling strategies in aquifers. Owing to the long model
run-times to simulate SWI, we limited the OD to steady-state measurements of head and salinity. With transient measurements, it is expected that the proposed locations for observation wells will shift landward with the movement of the interface.

Acknowledgments
This study is part of a program on climate change and sea water intrusion along the Eastern Mediterranean funded by the International Development Research Center (IDRC) of Canada at the American University of Beirut Grant No. 106706-001. Special thanks are extended to Dr. Charlotte Macalister at IDRC for her support and feedback in implementing this program and to Dr. Steen Christensen for his invaluable help, which greatly improved this manuscript.

Authors' Note
The author(s) does not have any conflicts of interest or financial disclosures to report.

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