

1 **Numerical modeling and data-worth analysis for characterizing the
2 architecture and dissolution rates of a multicomponent DNAPL source**

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12 **Key Points**

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14 • Aqueous-phase concentrations monitored in a field experiment were simulated to quantify
15 NAPL distribution and dissolution rates
16 • Depletion profiles of the most soluble DNAPL component accurately constrained the source
17 zone architecture
18 • Multiscale heterogeneity of source zone architecture controlled the uncertainty of estimated
19 mass transfer coefficients

20
21 **Abstract**

22
23 A numerical solute transport model was history matched to a high-resolution monitoring dataset
24 to characterize a multicomponent source of nonaqueous phase liquids (NAPLs) and evaluate the
25 uncertainty of estimated parameters. The dissolution of NAPL mass was simulated using the
26 SEAM3D solute transport model with spatially-varying NAPL saturations and mass transfer rate
27 coefficients, representing the heterogeneous architecture of the source zone. Source zone
28 parameters were simultaneously estimated using PEST from aqueous-phase concentrations
29 measured in a multilevel monitoring transect and from mass recovery rates measured at extraction
30 wells during a controlled field experiment. Data-worth analyses, facilitated by PEST ancillary
31 software, linked maximum aqueous-phase concentrations of all compounds to reductions in prior
32 uncertainty of mass transfer coefficients. In turn, transient concentrations of the most soluble
33 NAPL fraction constrained the source mass estimation. Accurately estimating the source mass and
34 reducing prior uncertainties was possible by removing concentrations measured during early
35 NAPL dissolution stages, identified as prior-data conflicts using the iterative ensemble smoother
36 PESTPP-iES. Prior-based Monte Carlo analyses highlighted model limitations for representing
37 sub-grid-scale heterogeneity of source zone architecture and NAPL dissolution, yet history-
38 matching of final dissolution stages measured at multilevel ports eliminated parameter bias and
39 produced long-term projections of source depletion with multistage behavior. Including mass
40 discharge constraints further improved the accuracy of source mass estimation, complementing
41 multilevel monitoring constraints on the source architecture and mass transfer coefficients.

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47 **Plain Language Summary**

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49 Quantifying dissipation timeframes of DNAPL source zones and contaminant discharge rates is
50 fundamental for environmental-management support. Both variables depend on the distribution of
51 DNAPL mass (architecture) within the source zone, and cannot be quantified by direct observation
52 methods. We elucidated upon the worth of multilevel monitoring for characterizing the source
53 zone architecture of a field experiment with inverse numerical modeling of contaminant transport
54 and DNAPL dissolution. Uncertainties on estimated DNAPL distribution and dissolution rates
55 were primarily associated to variability in dissolved concentration trends at multiple scales.
56 Dissolved concentration peaks measured during early DNAPL dissolution stages were found
57 responsible for inducing model parameter and predictive errors. Yet the depleting signature of the
58 least soluble component accurately constrained the source zone architecture, combining mass
59 recovery rates with multilevel monitoring to reduce model uncertainties. Hence, our approach and
60 results have beneficial implications for management support of aged source zones undergoing final
61 depletion stages.

62 **Index Terms and Keywords**

63 1831 Groundwater Quality, 1846 Model calibration (3333), 1873 Uncertainty quantification
64 (3275), 4314 Mathematical and computer modeling, 1815 Monitoring, forecasting, prediction
65 (4315)

66 DNAPL source zone, numerical modeling, uncertainty quantification, DNAPL mass transfer,
67 source zone architecture, contaminant mass discharge

68
69 **1. Introduction**

70 Remediation and long-term dissipation of contaminant source zones comprised of dense
71 nonaqueous phase liquids (DNAPLs) in the subsurface encompasses technical challenges related
72 to uncertainty of DNAPL spatial distribution and dissolution rates (Kueper et al., 2014; Mayer &
73 Hassanzadeh, 2005; NRC, 2005). Entrapped DNAPL mass and saturation distributions in the
74 porous medium, referred to as the source zone “architecture”, are key parameters controlling
75 source-zone longevity and depletion behavior (Dekker and Abriola, 2000; DiFilippo & Brusseau,
76 2008). Typical multistage and nonmonotonic depletion profiles observed in monitoring data reflect
77 the gradual dissolution of NAPL accumulations with characteristic saturations (Brusseau et al.,
78 2013; Kokkinaki et al., 2014; Stewart et al., 2021). Accumulations of low-saturation ganglia
79 allowing for groundwater flow-through account for peaks of discharge concentrations at early
80 NAPL dissolution stages, whereas high-saturation pools with negligible hydraulic accessibility
81 account for dissolution tailing at the final stages of a source lifespan (Christ et al., 2010; Kueper
82 et al., 2014; Yang et al. 2018). Thus, estimating the depletion behavior and remedial timeframes
83 of a source zone requires knowledge on NAPL architecture, which is difficult to characterize with
84 direct observation methods (Engelmann et al., 2019), but can be estimated from field tests and
85 monitoring data using mathematical models of NAPL dissolution (Falta et al., 2005a, 2005b;
86 Stewart et al., 2021).

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92 Numerical modeling methods coupling groundwater flow and contaminant transport with NAPL
93 dissolution have been used to estimate saturation distributions and mass transfer rate coefficients
94 from monitoring data (Frind et al., 1999; Mobile et al., 2012; Saenton & illangasekare, 2004).
95 Researchers have focused on estimating NAPL architecture or depletion timeframes from
96 synthetically-generated source zones and aqueous-phase concentrations using several
97 mathematical approaches to simulating mass transfer. Several studies considered either a local
98 equilibrium assumption (LEA) or Gilland-Sherwood models of interphase mass transfer (Kang et
99 al., 2021a, 2021b; Marble et al., 2008; Saenton & Illangasekare, 2004). The applicability of LEA
100 in decision-support models is questionable because heterogeneity of aquifer hydraulic properties
101 and DNAPL architecture can induce flow bypassing and mass transfer rate limitations, resulting
102 in nonequilibrium concentrations typically observed at field sites (Falta, 2003; Kokkinaki et al.,
103 2013; Powers et al., 1992, 1994). Similarly, Gilland-Sherwood models rely on correlations
104 between empirical coefficients and soil particle sizes that were determined under specific bench-
105 scale conditions, which may not be applicable to field-scale problems with different hydraulic
106 conditions (Powers et al., 1994; Saenton & Illangasekare, 2007). Additional uncertainties on LEA
107 and Gilland-Sherwood models include grid discretization requirements, as both approaches have
108 been validated with pore-scale experimental data (Agaoglu et al., 2015; Falta, 2003).

109
110 Upscaled models have been developed to simulate NAPL dissolution kinetics over a representative
111 elementary volume (REV) incorporating source zone metrics (Christ et al., 2010; Marble et al.,
112 2008; Parker & Park, 2004; Saenton & Illangasekare, 2007; Stewart et al., 2021; Zhu & Skyes,
113 2004). These metrics include NAPL mass and descriptions of source zone architecture in the form
114 of areal dimensions of NAPL accumulations or the ganglia-to-pool (GTP) mass ratio metric
115 (Abriola et al., 2013; DiFilippo & Brusseau, 2011). Because these metrics are difficult to measure
116 at contaminated sites, upscaled models incorporating a spatially-varying lumped-process mass
117 transfer coefficient have also been used to interpret monitoring data and predict source depletion
118 timeframes (Guo et al., 2020; Marble et al., 2008; Mobile et al., 2012; Park & Parker, 2005). These
119 models simplify the heterogeneity of porous media, aqueous-phase velocities, NAPL architecture,
120 and dispersivity, into a single lumped-process parameter at the REV scale (Falda, 2003; Imhoff et
121 al., 1993; Luciano et al., 2018). Although scale-dependent mass transfer rate coefficients may
122 simplify grid discretization requirements, the parameterization of NAPL source zones for inverse
123 numerical modeling and uncertainty quantification with spatially-correlated random parameter
124 fields is not straightforward (Arshadi et al., 2020; Kang et al., 2021a, 2021b; Kock & Nowak,
125 2015, 2016)

126
127 Given that NAPL source zones have complex spatial morphologies with sharp saturation
128 transitions at fine scales, traditional interpolation and geostatistical methods used in groundwater
129 flow modeling may be not suitable for parameterizing NAPL source zones (Arshadi et al., 2020;
130 Kang et al., 2021a). Alternative methods proposed for parameterizing NAPL source zones include
131 deep learning algorithms trained with images of saturation distributions generated with multiphase
132 flow simulations on highly-resolved permeability fields (Arshadi et al., 2020; Kang et al., 2021a,
133 2021b), posing additional data requirements and uncertainties on porous media characteristics and
134 model parameters (Abriola, 1989; Agaoglu et al., 2015; Miller et al., 1998). Moreover, these
135 parameterization methods have been tested with synthetically-generated source zones to estimate
136 categories of NAPL saturations through inverse modeling conditioned by borehole data (Arshadi
137 et al., 2020), or by aqueous-phase concentrations under LEA (Kang et al., 2021a, 2021b). Although

138 these methods can generate physically-based, spatially-correlated categorical parameters, they are
139 computationally expensive and require further validation and verification with field data.

140
141 Numerical models with parsimoniously parameterized source zones have proved useful for
142 characterizing NAPL architecture and/or lumped-process mass transfer coefficients (Marble et al.,
143 2008; Mobile et al., 2012; Saneton & Illangasekare, 2004). Moreover, combining multilevel
144 monitoring with recovery rates of contaminant mass (or with conventional monitoring wells) can
145 be valuable for characterizing heterogeneous NAPL architectures, as spatially-varying
146 contaminant fluxes may be mapped to soil horizons harboring NAPL mass within a source zone
147 (McMillan et al., 2018). Several studies have incorporated Gilliland-Sherwood or upscaled mass
148 transfer functions in discretized NAPL zones or in dual-domain models to estimate grid-scale
149 parameters from multilevel monitoring data and/or mass discharge/flux measurements (Christ et
150 al., 2010; Falta, 2003; Frind et al., 1999; Guo et al., 2020; Mobile et al., 2012; Park & Parker 2005;
151 Saenton & Illangasekare, 2004). Although previous investigations have demonstrated the utility
152 of parameterization parsimony for characterizing NAPL source architecture and dissolution rates
153 with inverse modeling, the uncertainty of grid-scale parameters arising from the assimilation of
154 high-resolution monitoring data has not been investigated.
155

156 A primary objective in this numerical modeling study was to evaluate the worth of aqueous-phase
157 concentrations monitored at a multilevel sampling (MLS) transect in combination with total mass
158 discharge (MD) rates to quantify and reduce the uncertainty of the mass, architecture, and
159 dissolution rates of a multicomponent DNAPL source. The field experiment considered in this
160 study involved the creation a small-scale source zone with a heterogeneous architecture at the
161 Borden experimental site (Broholm et al., 1999). Broholm et al. (2005) quantified the initial mass
162 of the DNAPL mixture using multicomponent NAPL dissolution theory and mass balance
163 analyses. Mobile et al. (2012) estimated the source architecture and dissolution rates from MD
164 profiles and from a single, incomplete MLS nest, using an inverse modeling technique. They
165 constrained the initial NAPL mass by the known amount and the source zone dimensions by the
166 measured post-experimental footprint (Mobile et al., 2012). Our study expanded on both previous
167 analyses by incorporating the entire MLS transect to infer the source footprint and to quantify
168 initial NAPL mass, saturation distribution, and mass-transfer rate coefficients, while examining
169 causality of parameter uncertainty. A secondary objective was to investigate model limitations for
170 reproducing the observed system behavior, further elucidating upon monitoring data assimilation
171 for source zone characterization with inverse numerical modeling, while minimizing the
172 propensity for biasing timeframes of source zone persistence.
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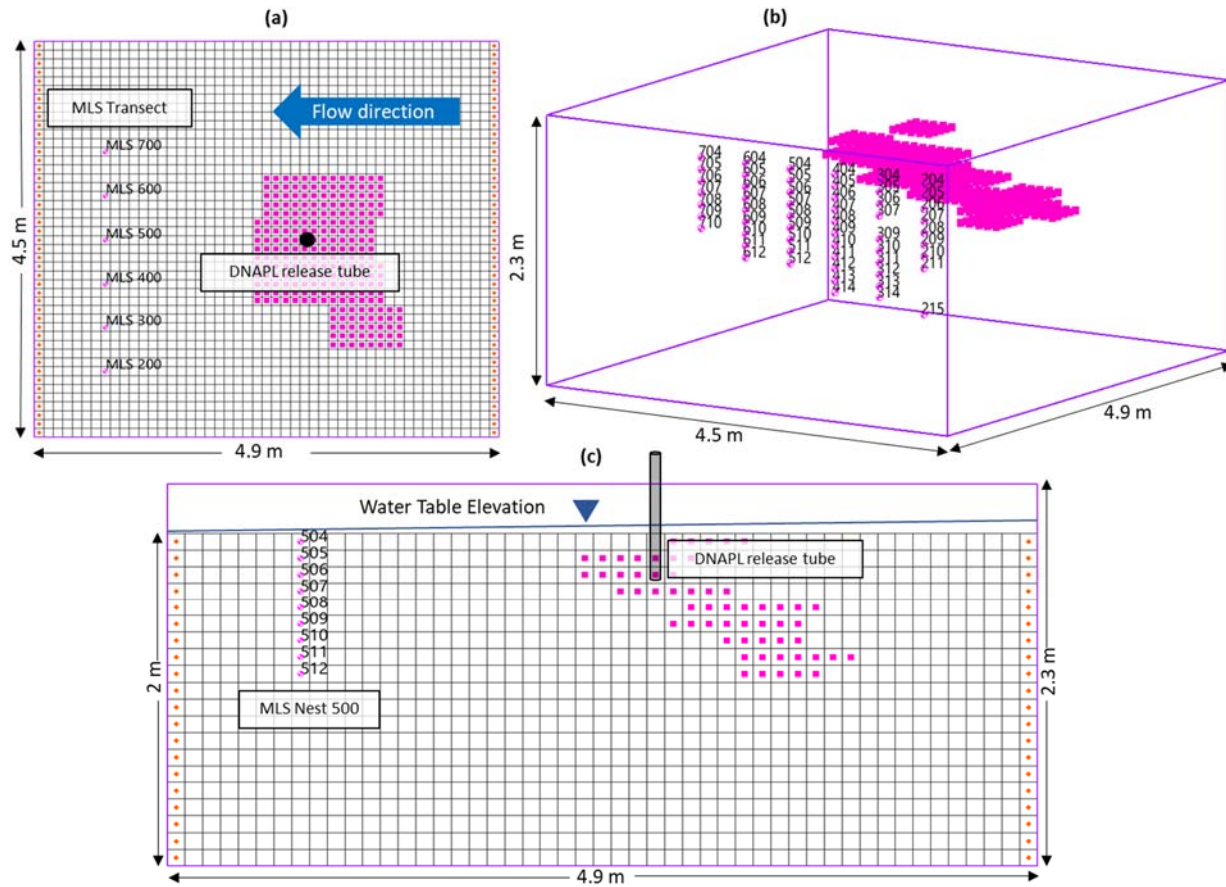
174 **2. Materials and Methods**

175 **2.1. Overview of Field Experiment of Multicomponent DNAPL Dissolution**

176 This study incorporated a dataset documenting the dissolution of a DNAPL mixture in a field
177 experiment (Broholm et al., 1999). At the Borden experimental site in Canada, a 5-liter (7.7 kg)
178 mixture of dyed solvents (10% TCM, 40% TCE, and 50% PCE by volume) was injected at
179 approximately 0.05 m below the water table into a 55 m³ (5.5 m long x 4.5 m wide x 2.3 m deep)
180 unconfined aquifer test cell (Figure 1) comprised of medium- to fine-grained lacustrine sand with
181 occasional beds of coarse sand/gravel and silt. Groundwater flow through the test cell, bounded
182

184 laterally by sheet piling on four sides, was maintained by a network of five upgradient injection
 185 and five downgradient extraction wells. The network was operated at approximately 360 L/d,
 186 generating a mean groundwater velocity of 0.13 m/d and horizontal hydraulic gradient of $0.023 \pm$
 187 0.0024. Aqueous-phase contaminant concentrations were monitored for 220 days in a fence of
 188 MLS ports (Figure 1) located approximately 2.1 m downgradient from the DNAPL release
 189 location. After 220 days of natural dissolution monitoring, a 5.5-day pulse of methanol was
 190 injected in the test cell to evaluate dissolution enhancement processes (Broholm et al., 1999;
 191 Broholm, 2006). The horizontal spacing of MLS ports was 0.5 m with a vertical spacing of 0.1 m,
 192 with screen lengths less than 1 cm.

193



194
 195 **Figure 1.** Configuration of aquifer test cell and its representation in the numerical model. (a) Plan view of all MLS
 196 nests and NAPL grid blocks encompassing the entire source zone footprint in the numerical model. (b) Test cell

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198 Flow-weighted concentrations were monitored at the extraction wells with screen lengths spanning
 199 the average height (1.82 m) of the saturated zone for 291 days. The cell was excavated 291 days
 200 after the mixture release to map the DNAPL distribution over 5-cm vertical intervals (Figures S1
 201 and S2) (Broholm et al., 1999). Broholm et al. (2005) compared estimates of initial NAPL mass
 202 calculated from the post-excavation source footprint and by multicomponent NAPL dissolution
 203 theory with mass balance analysis of effluent data and area- and depth-integrated MLS
 204 concentrations. Their best mass estimates ranged from 6.7 to 7.5 kg by averaging of MLS data,
 205 and differences between theoretical NAPL dissolution rates and effluent data were attributed to
 206 pre-flushing TCM dissolution and volatilization losses. Also, dissolved concentrations below

equilibrium levels observed throughout the experiment were attributed to dilution effects (Broholm et al., 2005). These results support kinetic mass transfer modeling coupled with uncertainty analyses for indirect characterization of NAPL source zones, as detailed, pore-scale process modeling and site characterization in this context is unfeasible for remedial-decision support at hazardous waste sites.

2.2. Numerical Modeling of Groundwater Flow and Contaminant Transport

A steady-state groundwater flow model was developed with MODFLOW2000 (Harbaugh et al., 2000). The aquifer test cell was discretized in 23 layers with uniform grid blocks measuring 10 cm along all dimensions. As shown in Figure 1, the extent of the model domain matched the size of the test cell along the vertical dimension (Z-axis) and the horizontal dimension perpendicular to the flow direction (Y-axis). The horizontal dimension parallel to flow (X-axis) was reduced from 5.5 m (test cell length) to 4.9 m for computational efficiency. Constant values of hydraulic conductivity, flow boundary conditions, and transport parameters were assigned to match field conditions, which were characterized with tracer tests and soil cores by Broholm et al. (1999) and analyzed with numerical modeling in Mobile et al. (2012). Model layers 1 through 3 were inactivated because the water table fell below the elevation of layer 3 during the monitoring period. Model layer 4 encompassed MLS port 504, which showed evidence of NAPL presence at 0.4 m below the top of the aquifer test cell despite the reported average depth of the water table during the experiment at ~0.48 m (Broholm et al., 1999).

Dissolution of the multicomponent DNAPL source and aqueous-phase contaminant transport were simulated with SEAM3D (Waddill & Widdowson, 2000). Interphase mass transfer [$M T^{-1} L^{-3}$] was simulated using a linear driving force model:

$$J_i = k_i^N (C_i^{eq} - C_i) \quad (1)$$

where $k_i^N [T^{-1}]$ is a lumped mass transfer rate coefficient specific to each NAPL phase constituent i , $C_i^{eq} [M L^{-3}]$ is the equilibrium solubility calculated according to Raoult's Law, and $C_i [M L^{-3}]$ is the aqueous phase concentration. Equation 1 is coupled in SEAM3D to the following relationship representing NAPL dissolution from the soil medium into the aqueous phase:

$$J_i = -\rho_b \frac{dC_i^N}{dt} \quad (2)$$

where $\rho_b [M L^{-3}]$ is the bulk density of the soil and $C_i^N [M M^{-1}]$ is the NAPL mass of compound i per unit mass of dry soil. A modified version of SEAM3D incorporates the upscaled NAPL dissolution model developed by Parker and Park (2004) to simulate transient mass transfer rates:

$$k_i^N = k_{i,0}^N \left(\frac{\bar{q}}{\bar{K}} \right)^\alpha \left(\frac{M_i(t)}{M_{i,0}} \right)^\beta \quad (3)$$

where $k_{i,0}^N$ = initial mass transfer rate coefficient [T^{-1}], \bar{q} = average Darcy velocity [$L T^{-1}$], \bar{K} = average hydraulic conductivity [$L T^{-1}$], $M(t)/M_0$ = transient ratio of NAPL mass [$M M^{-1}$], α and β are dimensionless empirical parameters. Previous investigations have reported a linear relationship

248 between k_i^N and \bar{q} , with $\alpha = 1$ (Parker & Park, 2004; Park & Parker, 2005). The transient mass
 249 ratio raised to the empirical depletion exponent β represents a reduction of NAPL/water interfacial
 250 areas over time, regulating tailing of discharge concentrations with reducing mass transfer rates as
 251 the source mass is depleted (Parker & Park, 2004; Stewart et al., 2020). A previous modeling study
 252 of the same Borden experiment indicated model insensitivity to the β exponent, attributed to a lack
 253 of extensive monitoring of decreasing discharge concentrations (Mobile et al., 2012). In this work,
 254 both the α and β parameters were set to zero to focus the uncertainty and data-worth analyses on
 255 the spatially-variable parameters $k_{i,0}^N$ and C_{0}^N . Adjusting C_{0}^N parameters allowed to estimate the
 256 initial mass (M_0^N [M]) of the entire NAPL mixture and quantify its uncertainty, as SEAM3D
 257 generates an output of remaining NAPL mass in the source zone every time step using Equation
 258 4, by adding the mass of all grid blocks where a C_{0}^N parameter value was assigned:
 259

$$M_0^N = \sum_{\text{Zone 1}}^{\text{Zone 23}} V^N \times C_0^N \times \rho_b \quad (4)$$

260 where V^N [L^3] = NAPL zone volume. The source zone was represented with 23 NAPL zones
 261 (Figures S1 and S2) positioned upgradient of MLS ports which showed contaminant breakthrough,
 262 suggesting the upgradient presence of NAPL mass (Figures 2, 3, and 4). Each NAPL zone was
 263 comprised of 25 grid blocks (Figures S1 and S2) and was assigned one pair of adjustable
 264 parameters, k_0^N and C_0^N , representing uniform mass distribution and dissolution within each NAPL
 265 zone. The areal dimensions of all NAPL zones were designed as 0.5 m x 0.5 m on the horizontal
 266 plane representing the horizontal spacing of MLS ports, whereas vertical layers of 0.1 m
 267 represented the vertical spacing between ports. The location of NAPL zones was determined by
 268 contaminant travel times analyzed from MLS breakthrough data. Overall, the source distribution
 269 in the model encompassed the observed post-excavation footprint (Figures S1 and S2), which
 270 likely developed through vertical and downward NAPL migration throughout the experiment
 271 (Broholm et al., 1999, 2005).

273 2.3. Parameter Estimation and Uncertainty Quantification

274 For each NAPL zone, C_0^N and k_0^N were simultaneously estimated from monitoring data. In
 275 addition, two global k_0^N multipliers to identify compound-specific mass transfer coefficients for
 276 TCE and PCE ($k_{TCE,0}^N$ and $k_{PCE,0}^N$) were estimated as multipliers of $k_{TCM,0}^N$ for a total of 48
 277 adjustable source zone parameters. History-matching targets included 1,556 measurements of
 278 dissolved TCM, TCE, and PCE concentrations monitored at the MLS transect (Figure 1), out of
 279 4,770 measurements comprising the entire MLS dataset. The 1,566 MLS targets corresponded to
 280 23 ports including concentration measurements through 130 days (Figures 2, 3, and 4), when
 281 extraction well redevelopment abruptly increased the water table by 1 m causing significant data
 282 noise through 220 days (Broholm et al., 1999; Mobile et al., 2012). These MLS targets were
 283 grouped by sampling port and a weight of 1 was assigned to each aqueous-phase concentration
 284 measurement within each port. Additional constraints included 78 measurements of mass
 285 discharge rates monitored at extraction wells for 220 days (before methanol remediation was
 286 implemented). Mass discharge measurements were grouped by contaminant (i.e., three MD
 287 groups) and assigned uniform weights, balancing the initial error contribution of each group to the
 288 objective function (Φ). Specifically, individual measurements of MD were assigned a weight of
 289 250 within the TCM and TCE MD groups, whereas individual PCE MD measurements were
 290

292 assigned a weight value of 500. This weighting strategy was designed to balance the visibility of
293 MD groups with MLS ports for Φ minimization.

294
295 An initial parameter estimation with PEST_ HP (Doherty, 2020), which uses a parallelizable
296 gradient-based optimization process, only included the MLS targets. All k_0^N and C_0^N parameters
297 were log-transformed to facilitate the nonlinear optimization process. Bounds for C_0^N parameters
298 were designed as a function of initial NAPL saturation ($S_0^N [\%]$) of the pore space calculated as:
299

$$S_0^N = \frac{\rho_b C_0^N}{\rho_N \theta} \quad (5)$$

300
301 where ρ_N [M L⁻³] is the NAPL density and θ is the soil porosity. Bounds for C_0^N parameters were
302 set as $0.05 < S_0^N (\%) < 25$ for most NAPL zones, and as $0.05 < S_0^N (\%) < 5$ for NAPL zones below
303 layer 10 and in layer 4. The prior (pre-history matching) NAPL mass value was set as ~120 %
304 greater than the known initial mass (7.7 kg). This prior value was established to evaluate whether
305 initial history-matching of MLS data with PEST_ HP could result in a total NAPL mass estimate
306 close to the known value. Bounds for all k_0^N ($k_{0,TCM}^N$) parameters were set as $0.01 < k_0^N (\text{d}^{-1}) < 7.5$
307 following an order-of-magnitude range obtained through a simplified mass transfer correlation
308 defined in Frind et al. (1999), where the dissolution of a large-scale DNAPL mixture in the Borden
309 aquifer was simulated using a similar grid scale. Bounds for k_0^N compound-specific multipliers
310 were kept consistent with ratios determined by Mobile et al. (2012) as $0.95 < k_{0,TCE}^N < 1$ and $0.8 <$
311 $k_{0,PCE}^N < 0.95$.

312
313 The posterior uncertainty of NAPL mass was quantified with the iterative ensemble smoother
314 PESTPP-iES (White et al., 2020). PESTPP-iES undertakes Monte-Carlo sampling of parameter
315 uncertainty bounds generating ensembles which are upgraded with the Gauss-Levenberg-
316 Marquardt (GLM) optimization algorithm. Rather than fitting simulation results to data, PESTPP-
317 iES can generate observation ensembles considering a multi-gaussian distribution of measurement
318 noise (ϵ) (White, 2018). Here, σ_ϵ was defined as 5% of measured values. This stochastic approach
319 was used for history-matching of (i) MLS data only, and (ii) both MLS and MD data, quantifying
320 the posterior uncertainty of parameters (C_0^N , k_0^N) and predictions (M_0^N). In the following sections,
321 Model A = optimized with PEST_ HP using MLS data only, Model B = optimized with PESTPP-
322 iES using MLS data only, and Model C = optimized with PESTPP-iES combining MLS and MD
323 data.

324
325 Parameter bounds were used to define 95% confidence intervals of multi-gaussian prior probability
326 distributions (PDF) of model parameters, assuming statistically-uncorrelated NAPL zones. The
327 upper C_0^N bounds for NAPL zones were reduced from 25% to 15% S_0^N and initial parameter values
328 were set from PEST_ HP results. Prior to parameter upgrading, PESTPP-iES undertakes a prior-
329 based Monte Carlo analysis to detect "prior-data conflicts" (PDC), which are measurements that
330 cannot be simulated with the structural and parametrization design of the model (White et al.,
331 2020, 2021). All PDCs flagged by PESTPP-iES were removed to eliminate history-matching
332 induced bias, which would otherwise produce erroneous parameter values compensating for model
333 defects (Doherty, 2015). Moreover, PESTPP-iES tracks the evolution of a "base realization"
334 during the optimization process, corresponding to the initial parameter realization upgraded

335 without ϵ ensembles. Estimates of NAPL mass removed by natural dissolution produced by Model
336 A were compared to estimates generated with the posterior base realization of Models B and C.
337

338 2.4. Data-Worth Analysis

339
340 First-order second-moment (FOSM) analysis was used to evaluate data worth for reducing the
341 uncertainty of model parameters and predictions through history-matching. The GENLINPRED
342 and PREDUNC utilities from the PEST software suite (Watermark Numerical Computing, 2018)
343 were used for FOSM analyses. The primary assumption in FOSM analysis is model linearization
344 expressed as:
345

$$346 \quad \mathbf{h} = \mathbf{Zk} + \boldsymbol{\epsilon} \quad (6)$$

347 which states that a vector of measurements of system state \mathbf{h} (aqueous-phase concentrations)
348 equals the action of the model \mathbf{Z} (Jacobian sensitivity matrix weighted by σ_{ϵ}^{-1}) on a vector of
349 parameters \mathbf{k} plus a vector of measurement noise $\boldsymbol{\epsilon}$ (Doherty, 2015). In this case, σ_{ϵ}^{-1} was calculated
350 on the basis of misfit between measurements and model outputs using the PEST-based utility
351 PWTADJ2 (Watermark Numerical Computing, 2018) after history-matching. GENLINPRED and
352 PREDUNC calculate the posterior uncertainty variance of model parameters through covariance
353 propagation:

$$354 \quad \mathbf{C}'(\mathbf{k}) = \mathbf{C}(\mathbf{k}) - \mathbf{C}(\mathbf{k})\mathbf{Z}^t[\mathbf{Z}\mathbf{C}(\mathbf{k})\mathbf{Z}^t + \mathbf{C}(\boldsymbol{\epsilon})]^{-1}\mathbf{Z}\mathbf{C}(\mathbf{k}) \quad (7)$$

355 where the posterior covariance matrix $\mathbf{C}'(\mathbf{k})$ is obtained through history-matching (Doherty, 2015).
356 In this case, the prior covariance matrix $\mathbf{C}(\mathbf{k})$ is diagonal with no spatial correlations between
357 NAPL zones. The estimated initial NAPL mass, a SEAM3D output, was treated as a linearized
358 model prediction:

$$360 \quad s = \mathbf{y}^t \mathbf{k} \quad (8)$$

361 where \mathbf{y} is a vector of sensitivities of s with respect to \mathbf{k} . The prior (σ_s^2) and posterior ($\sigma'_s{}^2$)
362 uncertainty variances of s were calculated as:
363

$$364 \quad \sigma_s^2 = \mathbf{y}^t \mathbf{C}(\mathbf{k}) \mathbf{y} \quad (9)$$

$$365 \quad \sigma'_s{}^2 = \mathbf{y}^t \mathbf{C}'(\mathbf{k}) \mathbf{y} \quad (10)$$

366 The worth of individual MLS ports for reducing prior parameter uncertainties was calculated with
367 model A, whereas the worth of compound-specific MLS and MD datasets was calculated with the
368 posterior base realization of model C. Model C was also used to quantify the worth of individual
369 MLS and MD measurements and to quantify the relative uncertainty variance reduction (RUVR)
370 of each parameter (i), defined as:
371

$$372 \quad RUVR_i = 1 - \frac{\sigma'_i{}^2}{\sigma_i^2} \quad (11)$$

374 where σ_i^2 are prior parameter variances encapsulated in $C(\mathbf{k})$ and σ'_i^2 are posterior parameter
375 variances extracted from $C'(\mathbf{k})$ (Doherty, 2015).

376

377 3. Results and Discussion

378

379 3.1. Parameter Estimation and Uncertainty Quantification

380

381 A comparison of estimated NAPL mass and k_0^N multipliers is presented in Table 1. In general, the
382 known NAPL mass (7.7 Kg) was underestimated (6.4 – 7.2 Kg) when history-matching to MLS
383 data only, particularly with gradient-based optimization (PEST_HP). An improvement in the
384 accuracy of mass estimation with PESTPP-iES was achieved through the removal of 206 PDC
385 values flagged by PESTPP-iES (Figures 2, 3, and 4) comprising 13% of the MLS constraints.
386 Likewise, 10 PDCs (Figure 5) representing 13% of the MD dataset were also flagged by PESTPP-
387 iES and removed for history-matching. Including the MD constraints resulted in an excellent
388 agreement with the known initial source mass, which was encompassed by 95% confidence limits
389 (Table 1). Prior-based Monte Carlo results suggested that emphasizing early peak concentrations
390 for history-matching can result in underestimation of NAPL mass and overestimation (bias) of k_0^N
391 values, leading to underestimation of source dissipation timeframes. In this case, removing PDC
392 values, rather than modifying the model design, was sufficient to accurately estimate NAPL mass
393 and constrain mass transfer coefficients.

394

395

Table 1. Model-estimated DNAPL mass and k_0^N multipliers.

Parameter/Prediction	PEST_HP (A: MLS)	PESTPP-iES (B: MLS)	PESTPP-iES (C: MLS and MD)
Mass (Kg)	6.367	$\mu = 7.187$ $\sigma = 0.120$	$\mu = 7.626$ $\sigma = 0.110$
$k_{0,TCE}^N$ (d ⁻¹)	1.00	$\mu = 0.99$ $\sigma = 0.006$	$\mu = 0.99$ $\sigma = 0.005$
$k_{0,PCE}^N$ (d ⁻¹)	0.95	$\mu = 0.93$ $\sigma = 0.018$	$\mu = 0.88$ $\sigma = 0.017$

396 μ = mean estimated value. σ = standard deviation of estimated parameters (k_0^N) and predictions (Mass).
397
398
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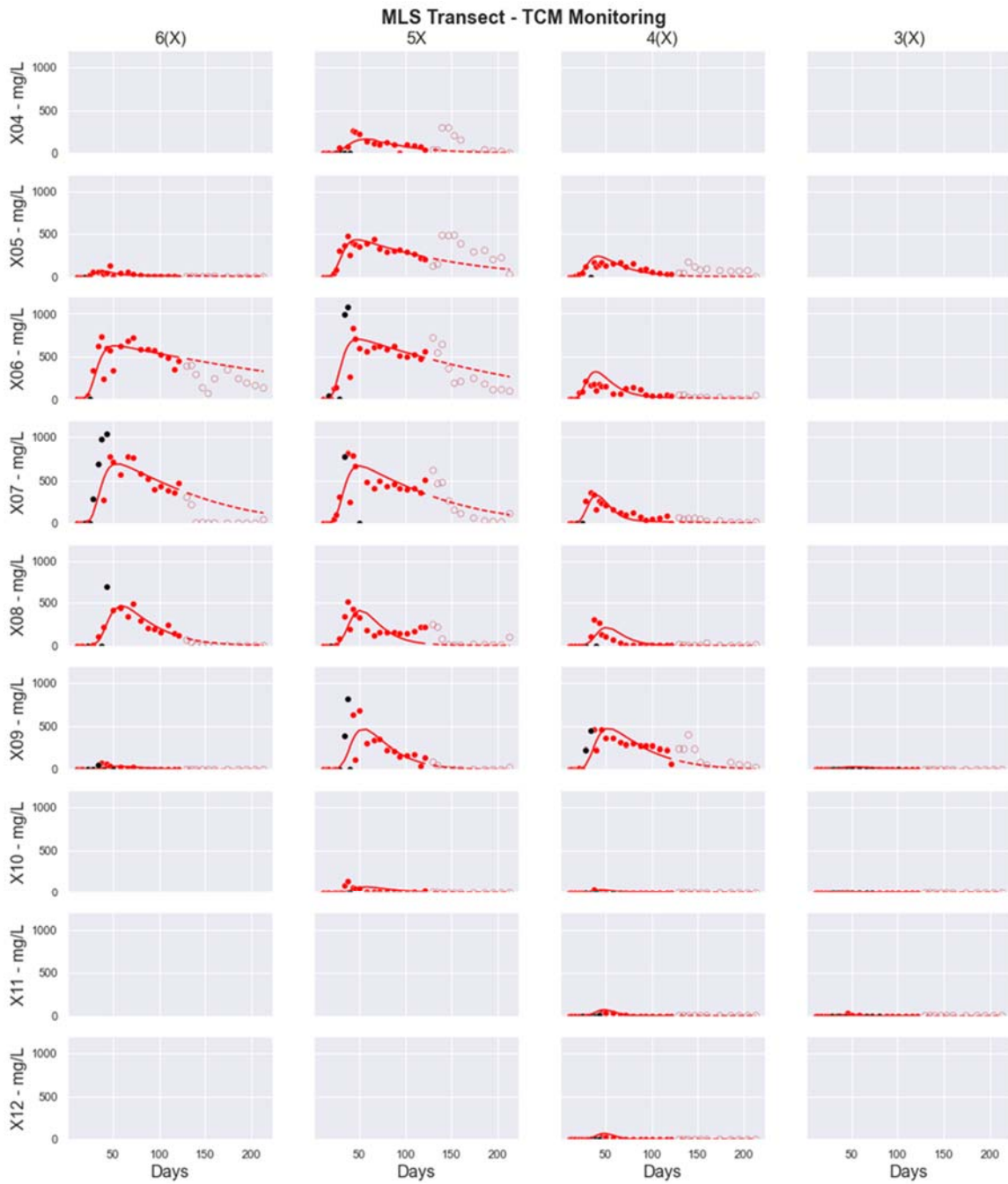
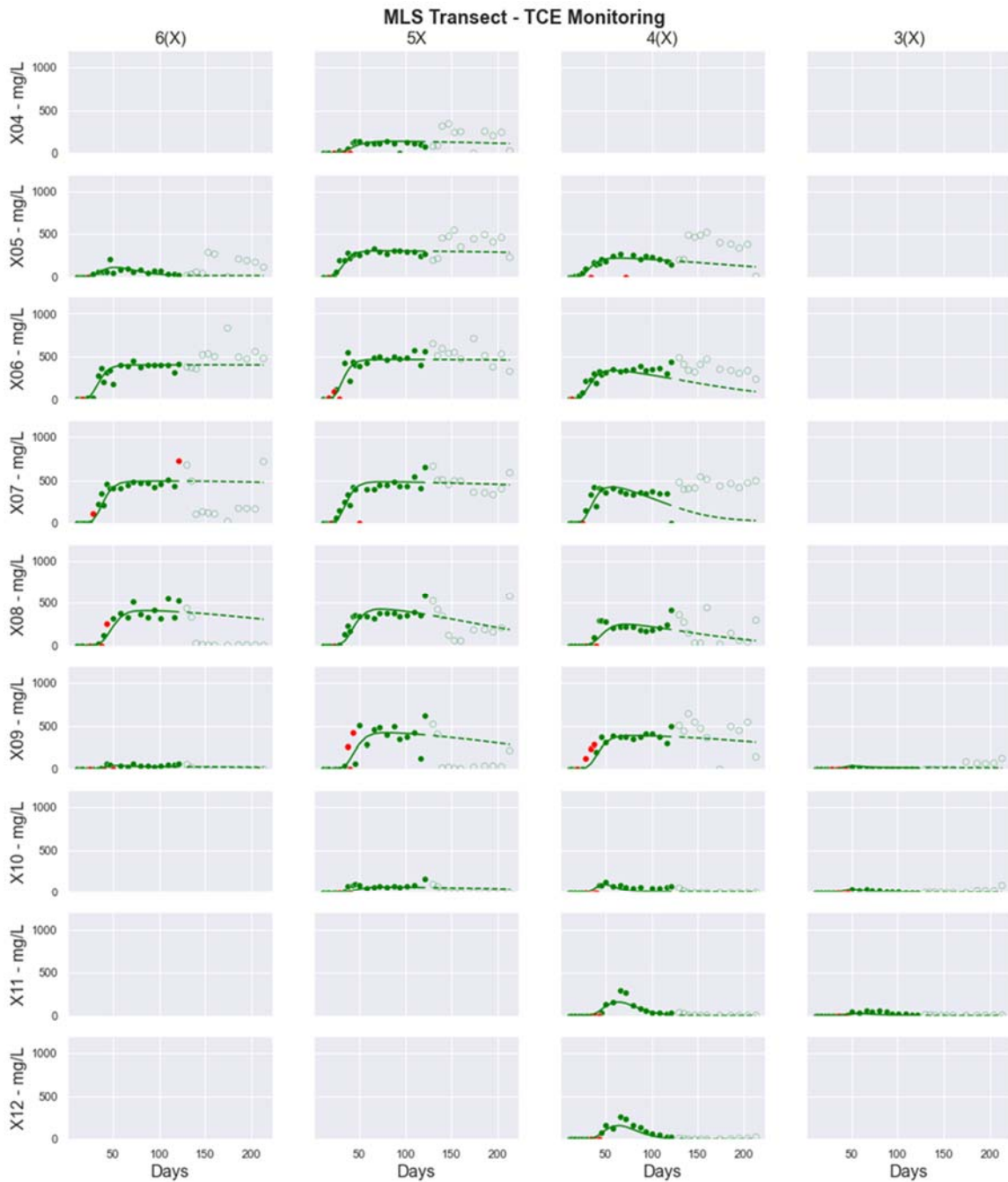


Figure 2. Comparison of measured (circles) and simulated (lines) aqueous-phase TCM concentrations at the MLS fence. Simulation results correspond to the posterior base realization of model C, including MLS and MD constraints. Empty circles correspond to concentrations ignored for history-matching because of significant measurement noise induced by water table fluctuations after 130 days of monitoring. The dashed lines correspond to simulated values beyond 130 days, informed by MD data exclusively. Black-filled circles are prior-data conflicts removed from the history-matching process to avoid parameter bias.

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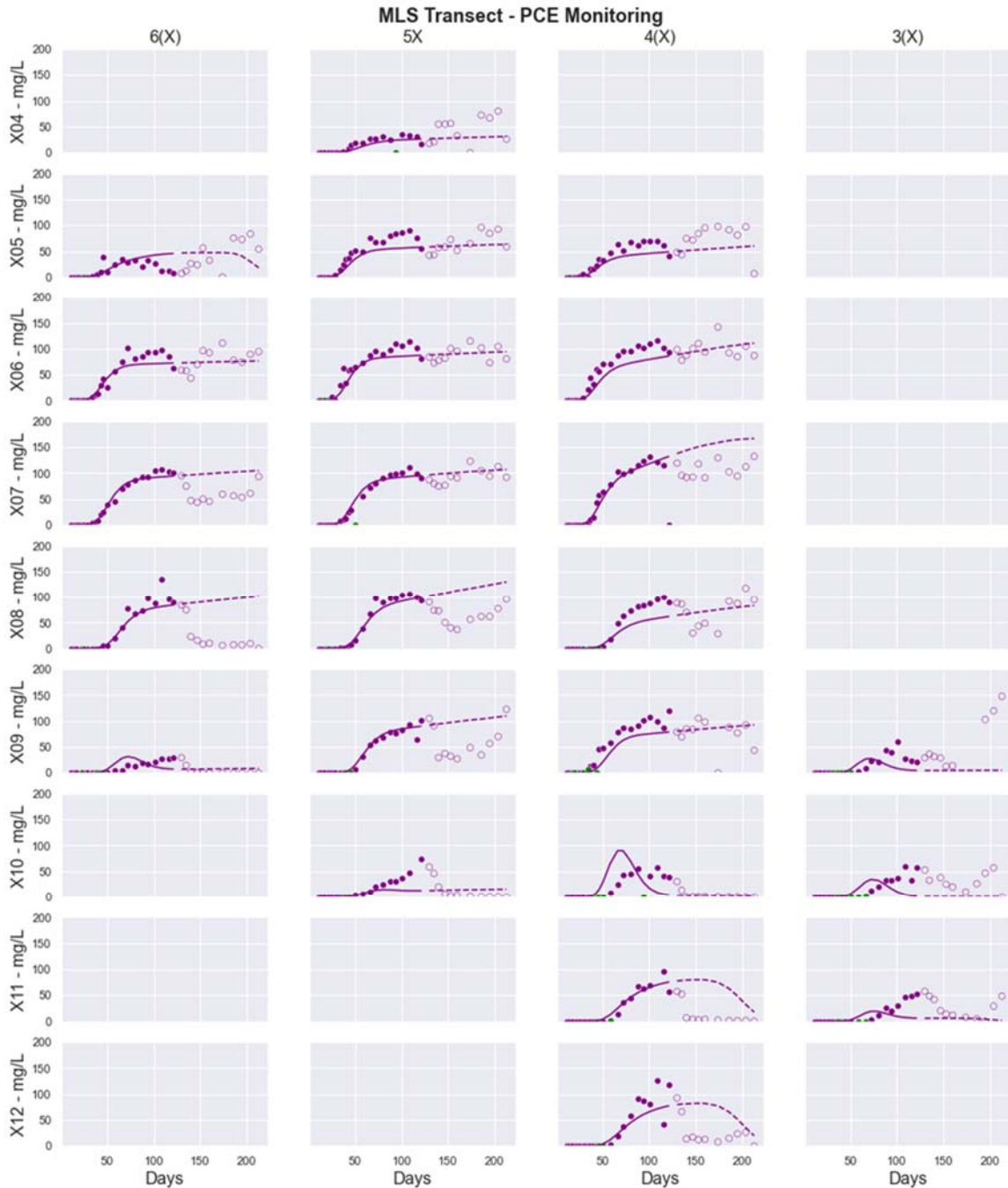
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Figure 3. Comparison of measured (circles) and simulated (lines) aqueous-phase TCE concentrations at the MLS fence. Simulation results correspond to the posterior base realization of model C, including MLS and MD constraints. Empty circles correspond to concentrations ignored for history-matching because of significant measurement noise induced by water table fluctuations after 130 days of monitoring. The dashed lines correspond to simulated values beyond 130 days, informed by MD data exclusively. Red-filled circles are prior-data conflicts removed from the history-matching process to avoid parameter bias.



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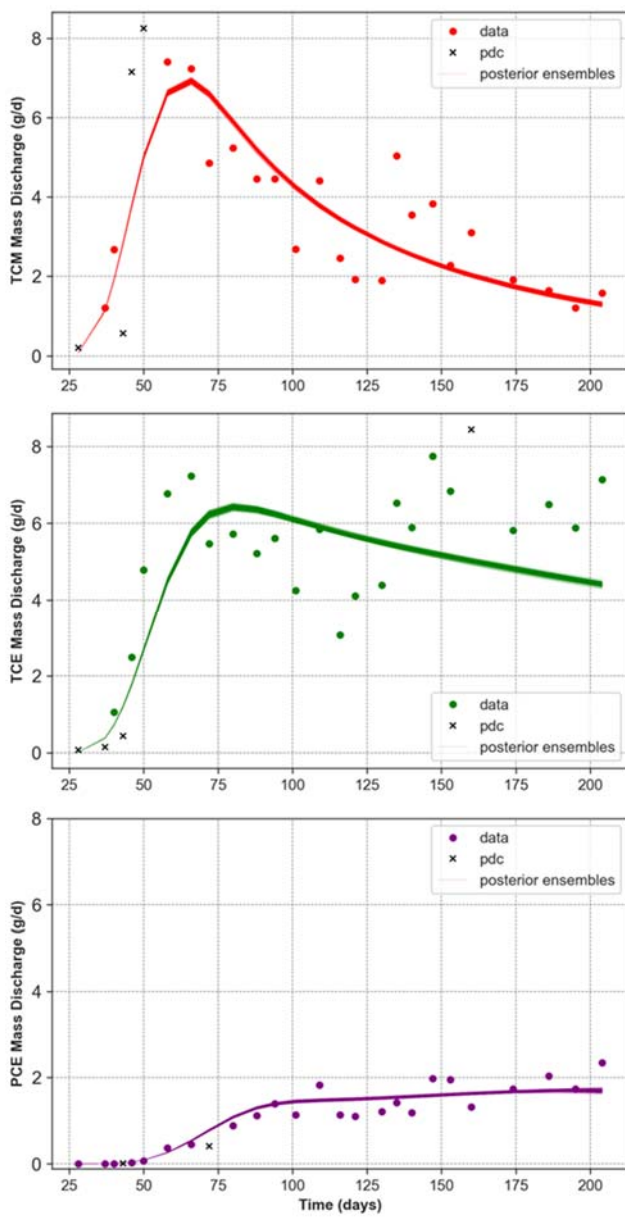
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Figure 4. Comparison of measured (circles) and simulated (lines) aqueous-phase PCE concentrations at the MLS fence. Simulation results correspond to the posterior base realization of model C, including MLS and MD constraints. Empty circles correspond to concentrations ignored for history-matching because of significant measurement noise induced by water table fluctuations after 130 days of monitoring. The dashed lines correspond to simulated values beyond 130 days, informed by MD data exclusively. Green-filled circles are prior-data conflicts removed from the history-matching process to avoid parameter bias.

Prior-data conflicts pertaining to each dissolved NAPL component were detected at similar locations along the MLS and MD profiles (Figures 2 through 5). Most PDCs corresponded to initial TCM concentration peaks, some were detected along TCE breakthrough, and a few before PCE breakthrough. This may have been associated to propagation rates of component-specific mass

432 transfer zones in the NAPL as the source architecture developed. In theory, the dissolution process
 433 of NAPL mixtures has been described as a chromatographic process, where component-specific
 434 mass transfer zones propagate at different velocities through NAPL accumulations as a function
 435 of their local solubilities and their length along the principal flow direction (Geller & Hunt, 1993;
 436 Soerens et al., 1998). In practice, the numerical discretization of NAPL zones along the flow
 437 direction may influence the estimation of k_0^N values, as grid-scale concentration gradients
 438 (Equation 1) would also regulate the sequential mass transfer process for any prescribed q (Darcy
 439 velocity) and $\alpha_{L,T}$ (dispersivity) values (Falta, 2003; Frind et al., 1999; Hunt & Sitar, 1988). In this
 440 work, the unknown source zone architecture along the flow direction may have placed additional
 441 uncertainties on q , $\alpha_{L,T}$, and $k_{i,0}^N$ parameters, which regulate overall NAPL mass transfer at the grid
 442 scale (Rivett & Feenstra, 2005).

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Figure 5. Posterior ensembles of MD profiles generated with model C.

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447 Table 1 indicates that both $k_{i,0}^N$ multipliers estimated with Model A showed a tendency to reach
 448 their upper bounds. Only the values of $k_{PCE,0}^N$ estimated with PESTPP-iES were constrained within
 449 their prior uncertainty bounds (Table 1). In addition to the uncertain effects of grid scale on
 450 potential parameter bias, multistage NAPL dissolution below the MLS scale may have impacted
 451 posterior results. For example, inspecting the TCM signature of ports 506 and 507 in Figure 2
 452 suggested two slopes of declining concentrations before 130 days. An initially steep slope between
 453 the concentration peaks through day 50, followed by a more gradual slope through day 130,
 454 suggested heterogeneity of NAPL architecture and dissolution below the MLS scale. Despite
 455 removing PDCs, sub-grid-scale multistage NAPL dissolution cannot be adequately simulated with
 456 a single parameter set (C_0^N , k_0^N) per NAPL zone. This explained why model A produced lower
 457 mass estimates with an increased propensity for biasing (overestimating) $k_{TCM,0}^N$ parameters (e.g.,
 458 port 608 in Table 2), whereas model B (PDC targets removed) produced $k_{TCM,0}^N$ values consistent
 459 with model C (Table 2). Although these results suggested that a dual-domain approach may have
 460 better captured TCM profiles at MLS ports, removing PDCs to estimate a single-domain
 461 parameters accurately constrained NAPL mass and a consistent range of $k_{0,TCM}^N$ values.

462

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Table 2. Distribution of estimated DNAPL mass and mass transfer coefficients

NAPL Zone	MLS Port	$k_{TCM,0}^N$ (day ⁻¹)			NAPL Mass (Kg)		
		A	B	C	A	B	C
4	404	0.037	0.041	0.041	0.14	0.14	0.15
5.1	605	0.038	0.029	0.037	0.02	0.02	0.01
5.2	505	0.102	0.104	0.106	0.59	0.59	0.64
5.3	405	0.050	0.061	0.067	0.12	0.13	0.13
6.1	606	0.174	0.171	0.175	1.23	1.63	1.79
6.2	506	0.252	0.226	0.236	1.16	1.32	1.43
6.3	406	0.147	0.138	0.131	0.10	0.11	0.13
7.1	607	0.735	0.282	0.303	0.79	0.93	0.96
7.2	507	0.183	0.231	0.251	0.79	0.81	0.85
7.3	407	0.303	0.255	0.245	0.11	0.11	0.11
8.1	608	3.259	0.220	0.202	0.33	0.35	0.36
8.2	508	0.227	0.254	0.204	0.15	0.17	0.20
8.3	408	0.108	0.115	0.082	0.07	0.07	0.09
9.1	609	0.385	0.019	0.048	0.00	0.00	0.00
9.2	509	0.248	0.196	0.216	0.29	0.30	0.31
9.3	409	0.314	0.184	0.171	0.35	0.38	0.40
9.4	309	0.010	0.010	0.066	0.01	0.00	0.00
10.1	510	0.024	0.033	0.016	0.03	0.02	0.04
10.2	410	0.015	0.068	0.534	0.01	0.01	0.01
10.3	310	0.010	0.015	0.049	0.01	0.00	0.00
11.1	411	0.056	0.097	0.081	0.02	0.03	0.02
11.2	311	0.010	0.017	0.084	0.02	0.01	0.00
12	412	0.077	0.109	0.083	0.02	0.03	0.02
Total NAPL Mass (Kg)					6.4	7.2	7.6

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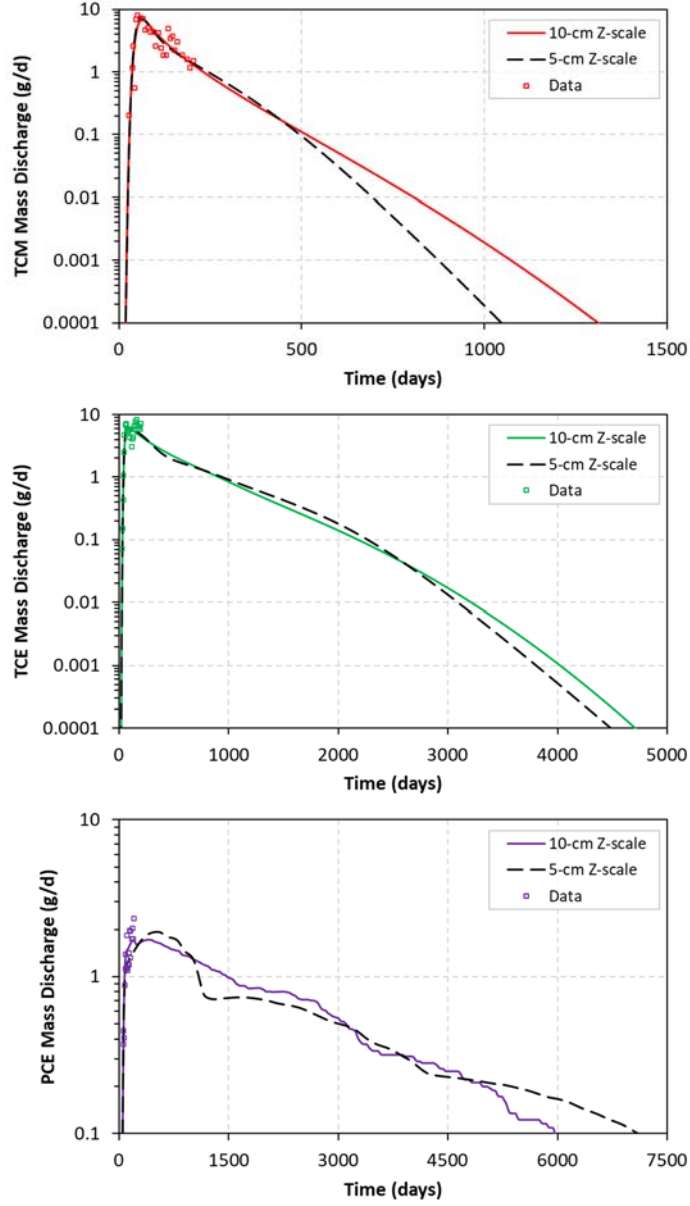
A (PEST_HP, MLS only), B (PESTPP-iES, MLS only), C (PESTPP-iES, MLS and MD)

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466 Long-term projections of MD using the posterior base realization of Model C (Table 2) are
467 compared to the model calibrated by Mobile et al. (2012) in Figure 6. In both cases, multistage
468 dissolution profiles emerged from the spatial distribution of NAPL mass and dissolution rates, as
469 MD projections were produced with $\beta = 0$ (Equation 3). The mass transfer rates shown in Table 2
470 encompassed the same order-of-magnitude range reported in Mobile et al. (2012), although the
471 values determined in this study were lower by ~50% on average, reflecting the grid-scale
472 dependence of estimated k_0^N values. Specifically, the grid-block size in Mobile et al. (2012) of 500
473 cm³ was also 50% smaller than the 1000 cm³ scale used in this study. Furthermore, stochastic
474 model optimizations with more adjustable NAPL zones than available MLS ports, using 5-cm
475 thick layers, resulted in mass overestimation by orders of magnitude and inconsistent k_0^N and C_0^N
476 distributions (results not shown).

477
478 **3.2. FOSM-Analysis Results**
479

480 The worth of monitoring datasets for reducing prior uncertainty variance of NAPL mass is shown
481 in Figure 7. This figure highlights the importance of TCM data for constraining NAPL mass in
482 contrast to the negligible worth in the monitoring profiles of other dissolved components.
483 Likewise, Figure 7 shows the increase in data worth for reducing prior uncertainty of NAPL mass
484 by removing PDCs. Although the short-term TCM MLS signatures (< 130 days) alone constrained
485 the prior uncertainty of NAPL mass by ~100%, adding the complete TCM MD signature (220
486 days) improved mass estimates by spreading ϵ induced by water table fluctuations across all NAPL
487 zones. Furthermore, Table 3 indicates a decrease in the worth of MLS ports commensurate with
488 the S_0^N of their corresponding upgradient NAPL zones. This apparent correlation reflected the
489 similarity between the TCM dissolution profile of each MLS port and the TCM MD profile,
490 emphasizing the indirect value of multilevel monitoring for characterizing NAPL distribution and
491 reducing the uncertainty of source depletion rates.

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Figure 6. Comparisons of long-term projections of MD profiles generated with the base parameter realization of model C (continuous lines) and the modeling results of Mobile et al. (2012) (dashed lines). All projections were generated with $\beta = 0$. In both cases, multistage behavior of NAPL depletion emerged from the NAPL architecture, which was constrained by the known mass and the post-experiment source footprint in Mobile et al. (2012). Small differences in long-term projections of source depletion emphasized the importance of constraining the source mass.

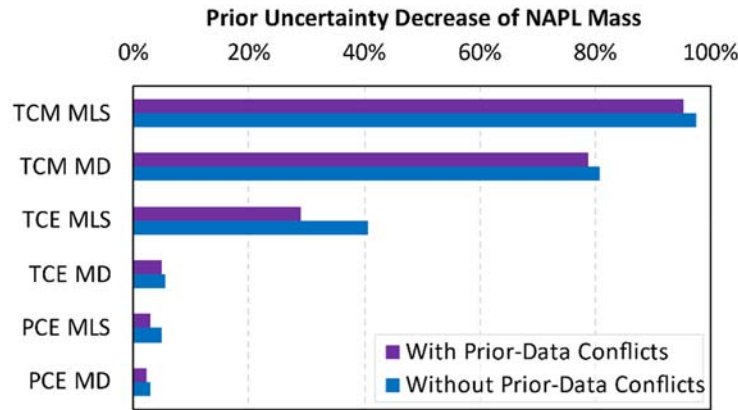


Figure 7. Percent worth of monitoring datasets for reducing the prior uncertainty of initial source mass.

Table 3. Distribution of S_0^N and worth of MLS ports for reducing prior uncertainty of NAPL mass.

NAPL Zone	MLS Port	Prior Uncertainty Variance Decrease	S_0^N (%)	S_0^N (%)	S_0^N (%)
		A	B	C	
6.1	606	54.0%	9.7%	12.8%	14.0%
6.2	506	42.6%	9.1%	10.4%	11.2%
7.2	507	19.0%	6.2%	6.4%	6.7%
7.1	607	15.7%	6.2%	7.3%	7.5%
5.2	505	5.1%	4.6%	4.6%	5.1%
8.1	608	4.0%	2.6%	2.8%	2.8%
6.3	406	3.2%	0.8%	0.9%	1.0%
7.3	407	2.7%	0.9%	0.9%	0.9%
8.2	508	2.6%	1.2%	1.3%	1.6%
9.3	409	2.4%	2.8%	3.0%	3.1%
9.2	509	1.8%	2.3%	2.4%	2.4%
5.1	605	1.5%	0.1%	0.1%	0.1%
9.1	609	0.7%	0.0%	0.0%	0.0%
9.4	309	0.6%	0.1%	0.0%	0.0%
5.3	405	0.6%	1.0%	1.0%	1.0%
8.3	408	0.3%	0.5%	0.5%	0.7%
10.2	410	0.0%	0.1%	0.1%	0.1%
10.1	510	0.0%	0.2%	0.2%	0.3%
12	412	0.0%	0.1%	0.3%	0.2%
11.1	411	0.0%	0.1%	0.2%	0.2%
11.2	311	0.0%	0.1%	0.1%	0.0%
10.3	310	0.0%	0.1%	0.0%	0.0%
4	404	0.0%	1.1%	1.1%	1.1%

A (PEST_HP, MLS only), B (PESTPP-iES, MLS only), C (PESTPP-iES, MLS and MD)

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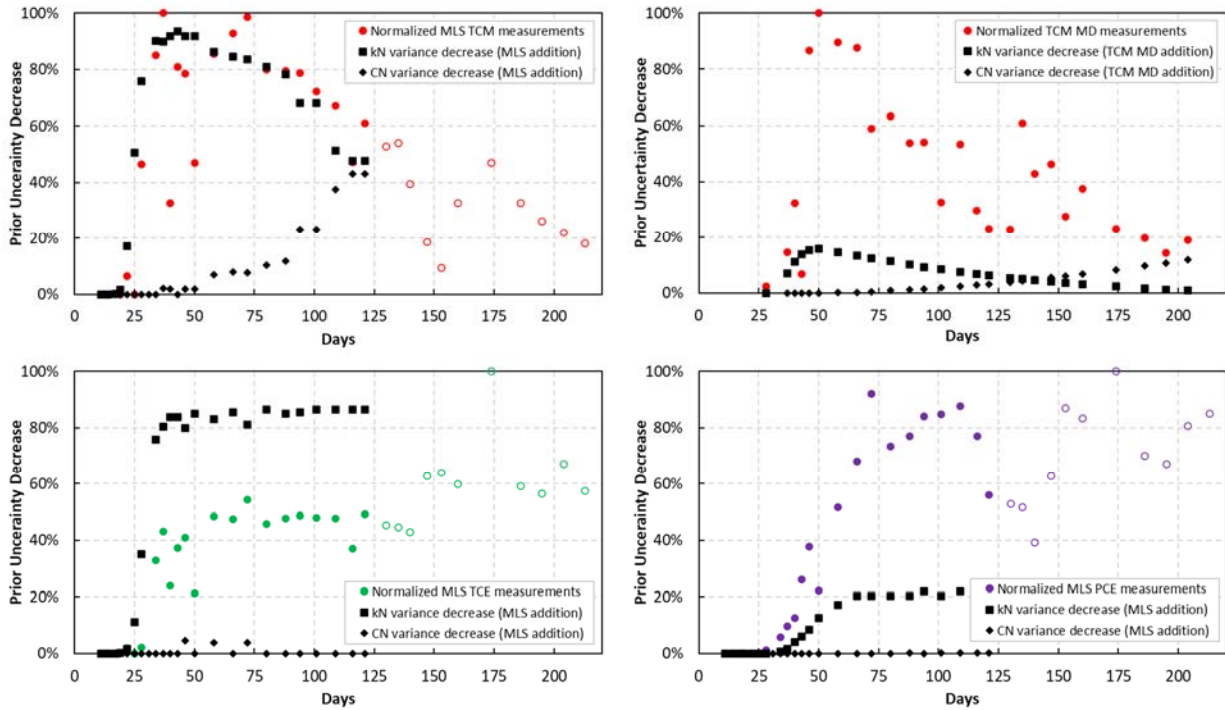
An example of the worth of individual MLS and MD measurements for reducing the prior uncertainty of NAPL-zone parameters is presented in Figure 8. These results indicated opposite trends in the worth of aqueous-phase concentrations for estimating C_0^N and k_0^N . Maximum

510 concentrations constrained k_0^N parameters, while declining concentrations constrained C_0^N
 511 parameters accounting for NAPL mass. This explained why only TCM data significantly reduced
 512 the prior uncertainty of NAPL mass, as all MLS ports showed declining TCM concentrations
 513 (Figure 2). Similarly, ~30% (with PDCs) and ~40% (without PDCs) reductions in the prior
 514 uncertainty of NAPL mass by TCE MLS data (Figure 7) was attributed to MLS ports with
 515 approximately more than 100 g of NAPL mass (e.g., port 508 in Table 2 and Figure 3) and
 516 declining TCE concentrations. Conversely, Table 4 indicates parameters with a low RUVR
 517 corresponding to deeper NAPL zones accounting for less than 1% of total NAPL mass. The low
 518 RUVR values in Table 4 were also caused by narrower prior uncertainty bounds compared to those
 519 of other NAPL-zone parameters. As shown in Figure 8, NAPL zones harboring most of the NAPL
 520 mass (e.g., 6.1 and 6.2 in Table 3) also benefited from additional C_0^N uncertainty reductions by
 521 TCM MD data, highlighting the contribution of those zones to the overall source depletion rates.
 522

523 **Table 4.** Source zone parameters with lower than 80% prior uncertainty variance reduction.

MLS Port	NAPL Zone	RUVR (with PDC)		RUVR (without PDC)	
		C_0^N	$k_{TCM,0}^N$	C_0^N	$k_{TCM,0}^N$
605	5.1	65%	> 80%	79%	> 80%
609	9.1	11%	13%	18%	22%
309	9.4	11%	5%	17%	8%
510	10.1	50%	> 80%	67%	> 80%
410	10.2	> 80%	17%	> 80%	18%
310	10.3	12%	15%	18%	15%
311	11.2	7%	1%	28%	1%
NA	$k_{TCE,0}^N$	NA	5%	NA	8%
NA	$k_{PCE,0}^N$	NA	18%	NA	30%

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 525



526
 527 **Figure 8.** Percent worth of individual aqueous-phase concentrations (MLS port 606 and TCM MD)
 528 for reducing the prior uncertainty variance of C_0^N and $k_{TCM,0}^N$ of NAPL zone 6.1. Although the same trend of added value by individual
 529 measurements was determined for all ports, only NAPL zones containing most of the source mass benefited from
 530 additional uncertainty reductions by TCM MD data. In turn, the correspondence between the TCM MD profile with
 531 individual MLS ports emphasized the value of multilevel monitoring for estimating NAPL architecture.
 532

533 Except for parameters listed in Table 4, history-matching reduced the prior uncertainty of all
 534 NAPL parameters by up to 100%. In contrast to C_0^N parameters constrained by TCM data
 535 exclusively, maximum TCE and PCE concentrations also constrained $k_{0,TCM}^N$ parameters (Figure
 536 7). However, as indicated in Table 4, the prior uncertainty of $k_{0,TCE}^N$ and $k_{0,PCE}^N$ (global $k_{0,TCM}^N$
 537 multipliers) was not reduced, partially because of their narrow prior uncertainty bounds. Yet the
 538 small RUVR of these mass transfer parameters was driven by their corresponding MLS datasets
 539 (results not shown). Moreover, while the mean values of both multipliers (Table 1) were in close
 540 agreement with those estimated by Mobile et al. (2012), as $k_{TCE,0}^N = 0.96$ and $k_{0,PCE}^N = 0.85$, FOSM
 541 analysis with Models A, B, and C suggested that $k_{TCE,0}^N$ could take a value greater than 1, which
 542 would not be consistent with previous findings (Mobile et al. 2012) or with mass transfer
 543 correlations with component diffusivities (Imhoff et al., 1993; Powers et al., 1992, 1994). As
 544 previously discussed, possible explanations for remaining uncertainties on $k_{TCE,0}^N$ and $k_{PCE,0}^N$ could
 545 include sub-grid-scale NAPL dissolution behavior, noticeable primarily in TCM MLS data, and/or
 546 the influence of grid scale on concentration gradients and α_{LT} (transverse and longitudinal).
 547 Transverse dispersion has been shown to regulate mass transfer rates from DNAPL pools (Hunt &
 548 Sitar, 1988, Stewart et al., 2021), requiring an ultrafine grid scale for accurate numerical
 549 simulations of DNAPL dissolution (Falta, 2003).

550 3.3. Analysis of DNAPL Mass Depletion

551 Table 5 presents mass balance results of NAPL mass removed by natural dissolution calculated
 552 using all models. The percent reductions of initial mass were calculated using the known initial
 553

composition values. Despite differences in the estimated source zone architectures, Model C produced nearly identical results as Mobile et al. (2012), emphasizing the importance of constraining NAPL mass for estimating source depletion rates. Conversely, Model A resulted in a 40% reduction of initial NAPL mass, almost doubling Model C results. Model C also indicated a source persistence at the end of the natural dissolution period ~4 and ~2 times lower than indicated by Models A and B, respectively, using PCE as reference. These results reflected the advantage of implementing prior Monte Carlo analyses to understand model deficiencies in relation to the observed system behavior. Additionally, Table 2 indicates the amount of NAPL mass eliminated by methanol remediation, calculated by subtracting the NAPL mass remaining in the soil estimated by Broholm et al. (1999) after conclusion of the experiment, from the remaining mass after 220 days estimated with Model C. Differences in the methanol calculations were linked to post-experiment mass estimated by Broholm et al. (1999) from C_i^N values assuming different S^N values. The obvious impact that such differences would have on remedial designs at hazardous waste sites highlighted difficulties in measuring S^N directly, even by soil confirmatory sampling. These results suggested value in the indirect source characterization method undertaken in this study to estimate and reduce the uncertainty of site-specific mass-transfer parameters, which is critical for effective, risk-based remedial optimizations.

Table 5. Mass of NAPL removed.

Data Source	NAPL Mass Removed (kg)				Initial NAPL Mass Reduction (%)			
	TCM	TCE	PCE	Total	TCM	TCE	PCE	Total
Initial (injected)	0.74	2.92	4.04	7.70	0	0	0	0
Model A	0.67	1.46	0.97	3.10	91	50	24	40
Model B	0.63	1.10	0.55	2.28	85	38	14	30
Model C	0.61	0.92	0.25	1.78	82	32	6	23
Mobile et al. (2012)	0.59	0.91	0.24	1.74	80	31	6	23
Methanol flush ⁽¹⁾	0.13	1.47	1.88	3.49	17	50	47	45
Methanol flush ⁽²⁾	0.13	1.18	0.80	2.11	17	40	20	27

Mass removed by methanol flushing was calculated by subtracting post-experimental NAPL mass remaining in soil estimated by Broholm et al. (1999) with C_i^N values assuming ⁽¹⁾ a homogeneous 3.6% S^N in all excavation layers (Figure S1) and ⁽²⁾ assuming 20% S^N in excavation layer 2 (where a DNAPL pool was observed), from the remaining NAPL mass on day 220 estimated with the posterior base realization of Model C. The percent reductions of initial NAPL mass were calculated with respect to the known initial composition of the mixture.

4. Conclusions

This study demonstrated the worth of high-resolution monitoring and inverse numerical modeling for characterizing a DNAPL source zone. The accuracy of estimated NAPL mass was tied to the depleting signature of MLS and MD aqueous-phase concentrations of the most soluble NAPL component and least by volume (TCM). At contaminated sites, decreasing concentrations may not reflect final NAPL dissolution stages, which could bias estimated parameters and long-term projections of source depletion. The impact of multiscale heterogeneity of NAPL architecture and dissolution on the uncertainty of model parameters was investigated with prior-based Monte Carlo analyses, where PDCs highlighted model limitations for representing sub-grid-scale mass transfer processes. Hence, multiscale heterogeneity of NAPL architecture and dissolution not captured in available monitoring profiles could limit model confidence for remedial-decision making at sites with large and architecturally complex source zones. These situations may benefit from the field

593 test proposed by Mobile et al. (2016) to determine mass transfer rate coefficients in situ. This test
594 would induce breakthrough of nonequilibrium concentrations through forced hydraulic gradients
595 and flushing in the source zone, generating monitoring profiles suitable for the inverse modeling
596 techniques applied in this study.

597
598 Breakthrough data collected from the mass transfer test described in Mobile et al. (2016) would
599 also be useful for allocating grid-scale NAPL zones, similar to the model parameterization guided
600 by MLS data in this study. In contrast to the simplified aquifer parameters in this work, spatially-
601 correlated hydraulic and transport properties can be characterized by geostatistical methods
602 coupled with numerical modeling for management support of source zones in heterogeneous
603 aquifers. Also, high-resolution NAPL delineation with MIP and LIF tools may further reduce
604 uncertainties on source architecture, including residual saturations estimated with inverse
605 numerical modeling. Furthermore, sites where natural attenuation mechanisms are significant may
606 benefit from several monitoring transects along the flow direction to distinguish attenuation
607 capacity from NAPL dissolution rates.

608
609 Uncertainty analyses confirmed an inability to estimate the β depletion exponent for any NAPL
610 zone despite declining TCM concentrations measured at all MLS ports. Thus, predictive
611 timeframes of source mass depletion should include a variability range for β parameters with
612 ensemble realizations, representing the transient nature of NAPL mass transfer rates in a stochastic
613 manner. Several advantages of parameter optimization and uncertainty quantification with model
614 ensembles were also demonstrated in this investigation. For example, while single parameter sets
615 per each NAPL zone could not represent sub-grid-scale multistage dissolution profiles, removing
616 early-stage TCM peak concentrations reduced the propensity for biasing mass transfer rates and
617 improved the accuracy of NAPL mass estimation. Accuracy of mass estimation was also attained
618 through parameter parsimony, as estimating NAPL-zone parameters without directly upgradient
619 MLS ports produced inconsistent and erroneous results. The importance of accurately constraining
620 the source mass was emphasized in a similar projection of source depletion compared to Mobile
621 et al. (2012), where multistage NAPL dissolution behavior in both models emerged from MLS
622 constraints.

623
624 Prior-based Monte Carlo and FOSM analyses suggested that simulating interphase mass transfer
625 from NAPL mixtures may be influenced by grid scale, despite incorporating adjustable compound-
626 specific mass transfer rate coefficients. Specifically, FOSM results indicated no prior uncertainty
627 reductions on the global multipliers of mass transfer rate coefficients, while $k_{TCE,\theta}^N$ showed a
628 tendency to exceed its upper uncertainty limit in all models, potentially biasing source dissipation
629 timeframes. Although these results did not prevent a reasonable estimation of initial source mass
630 and consistent ranges of mass transfer rates, further research is required to investigate the impact
631 of grid scale on dispersivity and mass transfer rate coefficients describing multicomponent NAPL
632 dissolution of source zones with heterogeneous architectures (i.e., comprised by ganglia- and pool-
633 dominated accumulations of NAPL mass). As demonstrated, combining mass discharge/flux rates
634 with high-resolution monitoring can improve history-matching of noisy data, where ensemble-
635 based parameter estimation considering measurement noise can reduce parameter bias without
636 resorting to a more complex simulation of multiple subsurface processes, supporting the indirect
637 characterization of NAPL source zones.

638

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640
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642 Program (ESTCP) under Project ER19-5223. The content of this manuscript has not been subject
643 to agency review and does not necessarily represent the view of the sponsoring agency.

644
645 **Data Availability Statement**

646
647 The Groundwater Modeling System (GMS) software hosting the MODFLOW2000 and SEAM3D
648 programs used in this study is available through Aquaveo at
649 <https://www.aquaveo.com/software/gms-groundwater-modeling-system-introduction>. The
650 PWTADJ2, PREDUNC, GENLINPRED, and PEST_HP programs are available at
651 <https://pesthomepage.org/programs>. The PESTPP-iES software is available at
652 <https://www.usgs.gov/software/pest-software-suite-parameter-estimation-uncertainty-analysis-management-optimization-and> (version 5.1.6 was used and the source code is available on
653 <https://github.com/usgs/pestpp/releases/tag/5.1.6>). Figures 2 through 4 were produced with the
654 Matplotlib (<https://matplotlib.org/>) version 3.5.1 and Seaborn (<https://seaborn.pydata.org/>) version
655 0.11.2 libraries using the Python programming language. Aqueous-phase concentration data will
656 be archived in an online repository maintained by Virginia Tech with a unique DOI number.
657

658
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