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Tracking of injected and resident (previously injected) bacterial cells in groundwater using ferrographic capture

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Abstract

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8 A high-resolution bacterial tracking technique, ferrographic capture, was used to enumerate fluorescent-stained bacterial 9 cells that were injected into groundwater during a field experiment. The goal of the experiment was to investigate whether 10detachment of previously injected stained resident cells attached to aquifer sediment was enhanced in the presence of the newly 11 injected mobile cells. This injection was an improvement on past experiments in that the attached (resident) cells were stained, 12allowing their concentrations to be enumerated directly by ferrographic capture (upon detachment). Contrary to expectations 13 based on previous experiments, enhanced detachment of stained resident cells did not occur upon the arrival of injected cells. 14Consistent with previous experiments, however, was the observation of ephemeral increases in unstained cell concentrations 15coincident with the arrival of the stained injected cells. The ephemeral pulses of unstained cells were previously speculated to 16represent enhanced detachment of unstained indigenous cells in response to hydrodynamic collision with injected cells. The 17lack of enhanced detachment of stained resident cells in the present experiments indicates that increased concentrations of unstained cells may have occurred by mechanisms other than hydrodynamic collision. Visually observed variations in stain 1819intensity indicated that increased unstained cell concentrations may have resulted from cell division at the low-concentration 20fringe of the injected plume.

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23 Keywords: Ferrographic capture; Bacterial cells; Hydrodynamic collision

24 26 **1. Introduction**

Bacterial attachment and detachment during transport in porous media is a subject of interest to those concerned with filtration for water treatment (Harvey and Garabedian, 1991), pathogen transport in groundwater (Harvey, 1997), and in situ bioaugmentation (Steffan et al., 1999). Bacteria are colloidal, being in the micrometer size range, and despite their greater 33 complexity, their transport is governed by many of the 34mechanisms that govern the transport of mineral 35colloids. Mechanisms controlling colloid detachment 36have been less studied relative to those governing 37 attachment, likely because rates of colloidal detach-38ment tend to be orders of magnitude lower than rates 39of attachment. The large difference in rates of attach-40 ment versus detachment allows the exclusion of 41detachment in models describing the gross transport 42behavior of colloids (e.g. filtration theory). However, 43second-order aspects of colloid transport are con-44

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trolled or influenced by detachment, e.g. bacterial 45tailing following an input pulse (Johnson et al., 46471995; Zhang et al., 2001), and the profile of attached cell concentrations versus distance (Zhang et al., 482001). Among the many processes potentially influ-4950encing colloidal detachment is hydrodynamic collision between mobile and attached colloids, which has 51been observed for microspheres in impinging jet flow 52experiments (Dabros, 1989; Dabros and van de Ven, 531992) and bacteria and microspheres in parallel plate 54chambers (Meinders et al., 1995). 55

That hydrodynamic collision between mobile and 56attached particles can enhance detachment of attached 57particles has been established based on force balance 58calculations (Dabros, 1989; Dabros and van de Ven, 5960 1992), which indicate that hydrodynamic interaction between deposited and flowing particles in stagnation 61 point flow systems causes significant variations of the 62normal force acting on the deposited particles even for 63relatively large particle separations, resulting in the 64 escape of weakly bound particles. Bacteria are rela-6566 tively "soft" particles, which thereby decreases their potential for hydrodynamic collision relative to 67 "hard" mineral colloids. However, after observing 68 69 increased detachment of attached microspheres and attached bacteria in the presence relative to the 70absence of mobile bacteria or microspheres, Meinders 71et al. (1995) concluded that hydrodynamic collision 72operated in their experiments in impinging jet flow 7374systems.

75Tentative observation of enhanced detachment of 76attached cells in the presence of mobile cells was recently reported from a bacterial transport study 77 conducted in Oyster, VA (Johnson et al., 2001). In 78this past study (conducted July 2000), two strains of 7980 bacteria were injected that had been previously isolated from the subsurface of the Oyster site (DeFlaun 81 et al., 1990). The isolates were stained with a vital 82fluorescent stain (Fuller et al., 2000) and were injected 83 into established flow cells at the site. Injected cells 84 were distinguished from resident cells (unstained 85 indigenous cells) on the basis of their internal stain. 86 87 Ephemeral increases in concentrations of unstained bacteria coincident with arrival of the stained bacteria 88 were observed in several wells during the transport 89 experiment. The unstained cells could have originated 90from either the injected cell population or the pop-91 92ulation indigenous to the aquifer (not injected). To

originate from the injected cell population, the 93 unstained cells would need to have been selectively 94concentrated relative to stained cells during transport. 95This could have occurred by division of injected cells 96 (assuming lack of stain transfer to daughter cells), loss 97 of stain by diffusion during transport, and lesser 98 adhesion of unstained relative to stained cells. Cell 99 division and stain loss in samples collected during 100peak breakthrough were insufficient to explain the 101 observed pulses of unstained cells. Standard adhesion 102assays indicated no difference in adhesion of stained 103versus unstained cells. Furthermore, to explain the 104 observed ephemeral dominance of unstained cells, 105selective concentration would need to have occurred 106exclusively on the low-concentration fringes of the 107 bacterial plume. Based on the above observation, it 108was tentatively concluded that the unstained cells 109 originated from the cell population indigenous to the 110aquifer (not injected). Potential mechanisms of 111 appearance of unstained indigenous cells include 112growth or detachment in response to the arrival of 113injected cells, with detachment more likely, given the 114lack of a lag time between arrival of the unstained and 115injected cells. 116

A significant drawback of the previous study was 117reliance on indirect means to quantify unstained cells, 118i.e. unstained cells were quantified by the difference 119between stained and total cells (stained plus un-120stained). In the present study, potential enhanced 121detachment of cells from the site sediment, in 122response to the arrival of mobile injected cells, was 123monitored directly since the attached cells were 124stained. It was determined in July 2001 that previ-125ously injected cells (remaining from the July 2000 126injection) remained intact and visibly stained in the 127site groundwater, and it was assumed that these cells 128also resided on the sediment. In July 2001, the same 129two bacterial strains were injected as in the previous 130year, however, the cells were stained oppositely from 131 the cells injected in the previous year (the July 2000 132injection). Switching the stains allowed for direct 133observation of the stained cells remaining from the 134July 2000 injection (hereafter referred to as stained 135resident cells). 136

Detachment of attached cells by hydrodynamic 137 collision with mobile cells is thought to require 138 generation of sufficient normal force to remove the 139 attached cells. The magnitude of the normal force 140

141 resulting from hydrodynamic collision is directly proportional to the rate of shear due to flow in the 142143system (Dabros and van de Ven, 1992). In experiments described by Dabros (1995) and Dabros and 144van de Ven (1992), the wall shear rates were in the 145range of $100-1000 \text{ s}^{-1}$. In the experiments per-146formed by Meinders et al. (1995), which examined 147rates of microsphere and bacteria detachment, the 148shear rates were ~ 50 s⁻¹. The shear rates in our 149experimental systems are calculated to be about 5-10150151 s^{-1} , based on idealized parabolic pore geometry 152(Bergendahl and Grasso, 2000) and a representative grain size of 300 µm. These calculated values are 5-15315410 times less than those examined by the abovedescribed studies (Dabros, 1995; Dabros and van de 155Ven, 1992; Meinders et al., 1995). However, in our 156system of packed porous media, the calculated values 157of shear likely underestimate the actual values for 158several reasons: (1) the grain size is distributed, 159thereby decreasing the pore throat sizes below those 160161expected for homogenous media that comprised of 162300-µm grains; (2) the grains are not smooth nor spherical, and thereby do not result in the parabolic 163pore geometry idealized for our calculation. 164

165Because the hypothesized collisions are hydrodynamic rather than direct, it may not be appropriate to 166assess the potential rate of hydrodynamic collision 167based on well-known rates of direct collision based on 168filtration theory (e.g. Ryan and Elimelech, 1996). 169However, given the above caveat, the rates of direct 170171 collision (per meter of transport) based on filtration 172theory were 60 and 850 for assumed average grain sizes of 500 and 100 µm, respectively. A range is 173given since the grain size of the natural sediment is 174distributed (the sediment displays an arithmetic mean 175176grain size of 270 μ m). The ranges in collision number obviously do not account for the packing effects of 177distributed grain sizes, but give some idea of the 178potential frequency of bacterial collision with sedi-179ment grain surfaces. The unstained cell concentrations 180in the pulses observed in the previous experiment 181 typically ranged from several hundred to several 182thousand cells per milliliter, which appears to be 183184supportable via collision numbers between 60 and 850 (per mobile cell). However, it must be stressed 185that the utility of the collision number in assessing the 186feasibility of hydrodynamic collision is limited, since 187 188 the "collisions" that are hypothesized to occur between mobile and attached bacteria are hydrodynamic 189 rather than direct. 190

There is no simple means known to the authors of 191accurately determining the values of shear or the 192potential frequency of hydrodynamic collision in a 193packed porous media. Hence, the opportunity of a 194direct test for enhanced detachment of resident cells 195was useful. If enhanced detachment of resident stained 196bacteria had been observed, then further investigation 197 of its causes would have been warranted. In contrast, 198the observed lack of enhanced detachment reported 199 below indicates the need to further consider other 200mechanisms of appearance of the unstained cells. 201Regardless of outcome, the investigation described 202below demonstrates the utility of the high-resolution 203technique that was used for tracking the bacteria. 204

2. Methods

Two bacterial strains were originally isolated from 206the site and were used in this study. DA001 is an 207aerobic adhesion-deficient variant selected using col-208umn assays, and is identified as a Comamonas sp. 209(DeFlaun et al., 1990). DA001 is Gram-negative and 210is $1.2 \times 0.6 \ \mu m$ in size. OY-107 is a facultative iron-211reducing bacterium of the genus Acidovorax that was 212determined to be naturally adhesion deficient upon 213isolation from the South Oyster (SO) site. OY-107 is a 214Gram-negative bacterium of size 1.9×1.0 µm. 215

Bacterial strains were grown by Envirogen (Law-216renceville, NJ) in minimal media supplemented with 217lactate using standard fermentation procedures. 218Strains were harvested by centrifugation and starved 219in at a cell concentration of approximately 10⁹ cells/ 220ml in artificial site groundwater for at least 48 h 221(DeFlaun et al., 2001). Cells were stained during 222preparation by Envirogen using vital fluorescent 223stains. Vital fluorescent stains cause no adverse effect 224to cell culturability (Fuller et al., 2000; Fuller et al., 2252001a,b). DA001 was stained using the green fluo-226rescent stain 5-(and 6)-carboxyfluorescein diacetate 227succinimidyl ester (CFDA/SE). OY-107 was stained 228using the red fluorescent stain 5-(and 6)-carboxyte-229tramethylrhodamine succinimidyl ester (TAMRA/SE). 230Immediately prior to injection, cell suspensions were 231diluted into site groundwater to a nominal concen-232 tration of 1.5e8 cells/ml. 233

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234Bacterial cell counts were monitored using ferrographic capture, which combines the selectivity of 235236immunomagnetic tagging with the high resolution of 237ferrography. For the ferrographic capture analyses, 238polyclonal rabbit antibodies (Rockland Immunochem-239icals, Gilbertsville, PA) raised to whole cells of the 240target bacterial strains were used to tether goat antirabbit-coated paramagnetic beads (50-nm diameter, 241Miltenyi Biotec, Auburn, CA) to the surface of the 242243target cells following sample collection (Zhang and 244Johnson, 1999; Zhang et al., 1999). The bacteriumbead suspension was introduced into a Bio-Ferrograph 245(Guilfoyle, Belmont, MA), which deposited the mag-246247netically tagged bacteria onto a small area on a glass substratum. The bacteria were then enumerated under 248249an epifluorescence microscope. TAMRA/SE- and 250CFDA/SE-stained cells were enumerated using red $(\lambda_{ex} = 510 - 560 \text{ nm}, \text{ dichroic} - 575 \text{ nm}, \lambda_{em} = 590 \text{ nm})$ 251and green ($\lambda_{ex} = 470 - 490$ nm, dichroic—505 nm, 252 $\lambda_{\rm em} = 510 - 530$ nm) filters, respectively. Previous 253254studies have shown that ferrographic capture provides virtually 100% recovery of target cells and yields 255extremely high analytical resolution (~ 20 cells/ml) 256(Zhang and Johnson, 1999; Zhang et al., 1999, 2001; 257258Johnson et al., 2000, 2001).

259The same two bacterial strains were used in the year 2000 and year 2001 injections; however, the 260stains were switched between the two strains for the 261July 2001 experiment relative to the July 2000 experi-262263ment (Table 1). Hence, in groundwater samples, the two different strains were distinguished antigenically, 264265whereas the injection year was distinguished using the internal stain. Unstained (or weakly stained) cells 266were also monitored using anti-DA001 and anti-OY-267107 antibodies that had been conjugated to a green 268269fluorophore, fluorescein isothiocyanate (FITC).

270 Since the FITC-conjugated antibodies provide a 271 visible outline of both unstained and stained target 272 cells, the unstained cell concentrations were deter-273 mined by difference between analyses using FITC-274 conjugated antibodies (stained plus unstained cells)

Table 1 Stains used during the series of experiments		
Year	DA001	OY-107
2000	TAMRA/SE	CFDA/SE
2001	CFDA/SE	TAMRA/SI

and nonconjugated antibodies (stained cells). The 275antibodies may have also captured antigenically sim-276ilar nontarget cells. Although nontarget cells would 277not be enumerated using nonconjugated antibodies (in 278analyses for stained cells), they would be enumerated 279using FITC-conjugated antibodies (in analyses for 280 stained plus unstained cells). Hence, the "unstained" 281cell counts may include both target strains as well as 282antigenically similar cells. Laboratory tests showed 283that the antibodies successfully capture wild-type 284DA001 and wild-type OY-107 cells (where wild type 285refers to initial isolates), indicating that the analyses 286 could indeed be capable of capturing indigenous 287forms of these strains. 288

Since both bacterial strains were potentially present 289in both colors in each groundwater sample, the extent 290of antibody cross-reactivity between the two bacterial 291 strains was determined. Serial dilutions of both bacte-292rial strains collected from their respective injection 293lines were analyzed using the respective nontarget 294 antibody. It was determined that approximately 1% of 295cells from serial dilutions of both bacterial strains 296reacted with their respective nontarget antibody (data 297not shown). However, unstained cell suspensions 298grown and serially diluted in the laboratory showed 299zero cross-reactivity with their nontarget antibody 300 (data not shown), indicating that the apparent 1% 301cross-reactivity observed in the injection line stand-302 ards may have resulted from cross-contamination 303 during sampling of the injection lines. Even 1% 304 apparent cross-reactivity would have insignificant 305 effect on the bulk of our analyses, since cell concen-306 trations examined by ferrographic capture were pri-307 marily in the 10-1000 cells/ml range, yielding a 308maximum of 10 cells/ml nontarget cells in the sam-309ples. However, interpretation of results from samples 310 containing greater than 1000 cells/ml necessarily 311considered potential effects of cross-reactivity. 312

Another potential artifact, given the possibility that 313each of the two strains could be present in each of the 314two colors, was the potential visibility of one stain 315under the filter used for the other stain during enu-316meration. To avoid this problem, filters were switched 317back and forth during enumeration to ensure that the 318cells being counted were exclusively observed under 319the target filter. 320

Samples taken from the injection solutions were 321 collected to serve as standards (following dilution to 322

1000 cells/ml) for ferrographic analysis. These stand-323 ards and blanks consisting of high purity Milli-Q 324 325water were included in each set of eight samples analyzed. Standard concentrations of both bacterial 326 strains were observed to degrade at approximately 1% 327328 per day in all ferrographic analyses (using FITCconjugated and nonconjugated antibodies). The 329 decrease in captured cell number over time is likely 330 due to antigen alteration in the presence of form-331 aldehyde, as has been previously reported (Johnson et 332333 al., 2000).

Flow cytometry (FACScan, Becton-Dickinson 334335Immunocytometry Systems, San Jose, CA) was used to monitor cell concentrations of CFDA/SE-stained 336 DA001 when they were present at concentrations 337 greater than 5000 cells/ml. Although ferrographic 338 capture can analyze cell concentrations above this 339 magnitude following 1:10 or greater dilution, flow 340 cytometry has the advantage of being extremely fast. 341A filter to allow detection of TAMRA/SE-stained OY-342 343 107 was not available on the instrument, and so TAMRA/SE-stained cells were enumerated exclu-344 sively using ferrographic capture. TransFluoSpheres 345(Molecular Probes, Eugene, OR) were added at a 346 347 known concentration (60,000 spheres/ml) to the samples in order to determine volume of sample analyzed. 348Each sample was run for 50 s at the low flow rate 349 setting of the machine, which corresponds to approx-350imately 10 µl/min. 351

352 **3. Field experiments**

The South Oyster (SO) focus area is one of two 353flow cells located at the South Oyster site in Oyster, 354VA on the southern end of the Delmarva Peninsula. 355356 The SO flow cell is located in a surficial aquifer that comprised of unconsolidated to weakly cemented, 357well-sorted, medium- to fine-grained sands and peb-358bly sands. The aquifer is underlain by a fine-grained 359 muddy silt aquitard situated approximately 9 m 360 below ground surface (BGS). A peat layer from 361 362 ~ 2.5 to 4.0 m BGS lies within the granular sands. 363 The SO focus area flow cell is bounded vertically by the peat layer above and the clay aquitard below. 364 The sediment between the peat layer and the aquitard 365 366 is dominantly very fine sand and fine sand (quartz, 367 feldspar, and significant micaceous minerals) with

occasional medium- and coarse-grained sand stringers. The SO flow cell is characterized as suboxic, with dissolved oxygen concentrations in the range of 0.2-1.0 mg/l.

Twenty-four multilevel samplers (MLS) exist in 372the flow cell (Fig. 1) in an array that is oriented with 373 the main axis parallel to natural local groundwater 374flow direction, as inferred from water level observa-375tions prior to site installation. The flow cell is 376 bounded at the up-gradient limit by the injection well 377 SO-B2 and at the down-gradient limits by three 378 extraction wells SO-A3, SO-B3, and SO-C3, which 379 form a line perpendicular to flow about 19.5 m down-380 gradient of SO-B2. 381

Each MLS consisted of a 3-cm-diameter PVC rod 382separated into 12 equally spaced zones that is sepa-383 rated by neoprene baffles. Stiff polyethylene tubing 384 (3.2 mm ID) extended from approximately 0.5 m 385above ground surface to each sampling port. Detailed 386 descriptions of the MLS used at SO focus area can be 387 found elsewhere (Mailloux et al., in press). In this 388 paper, the MLS are denoted by their number and the 389 prefix "SO", e.g. SO-17 denotes South Oyster focus 390 area MLS 17. The sampling port is denoted by 391



Fig. 1. Layout of the South Oyster focus area flow cell. Numbers refer to multilevel samplers (MLS). B2 is the injection MLS. The forced gradient was set 1 week prior to injection by extraction from three wells in a line oriented perpendicular to flow 12 m down-gradient of MLS SO-24.

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392 number (1-12) and the prefix "P", e.g. SO-17 P4 denotes port 4 in MLS 17. The port number increases 393 394with depth in the flow cell.

395 The ambient pore water velocity at the site is 396approximately 0.1 m/day. Forced gradient conditions 397 were established at the site approximately 48 h prior to the start of injection by withdrawing from the three 398 down-gradient extraction wells, SO-A3, SO-B3, and 399 SO-C3, at flow rates of 20, 40, and 20 ml/min, 400 respectively. The resulting average pore water veloc-401 402 ity was approximately 1 m/day.

Injection was conducted in MLS SO-B2, a 4-in. 403PVC well with four vertically discrete injection 404 zones separated by packers. In contrast to the 405previous experiment (July 2000), in which both 406 bacterial strains were injected over the entire depth 407 408 of the flow cell, the July 2001 experiment injected the two bacterial strains over two discreet depths. 409DA001 was injected over the interval of -3.5 to 410 -4.5 m relative to mean sea level (MSL), and OY-411 412 107 was injected over the interval of -5.0 to -6.0 m MSL, both at injection rate of 0.8 l/min. 413The bacterial injection zones were separated by 414 injection of unamended groundwater into the inter-415416 val from -4.5 to -5.0 m MSL at a rate of 0.4 l/ min. Unamended groundwater was also injected 417 above the DA001 injection zone $(-3.0 \text{ to } -3.5 \text{ to$ 418 m MSL) at a rate of 0.4 l/min. The total injection 419rate from SO-B2 was 2.5 l/min. Injection occurred 420 421over a 12-h period. Both bacterial injection solu-422 tions were sampled every 30 min during the injec-423 tion interval.

424 A custom manifold was developed to apply continuous suction to all sampled ports simultaneously 425using a peristaltic pump (Johnson et al., 2000; Mail-426427 loux et al., in press). All sampled ports were pumped 428continuously and equally in order to minimize inducement of a vertical hydraulic gradient between the 429ports. Relative to the previous experiment (July 4302000), the number of samples taken during early 431432breakthrough was increased in the July 2001 experiment to increase the resolution of the early break-433 434through pulses. All bacterial samples were taken using 43550-ml polypropylene centrifuge tubes and then fixed with 1% formaldehyde. Samples were then split into 436smaller aliquots (15 ml), briefly stored on ice, shipped 437 back to the University of Utah on ice (unfrozen), and 438 stored at 4 °C. 439

4. Laboratory experiments

Several laboratory experiments were run to inves-441 tigate potential enhanced detachment of attached cells 442in a controlled system. Each experiment involved a 443loading step, in which a bacterial suspension was 444 introduced to a sediment-packed column. Loading 445was followed by elution until the effluent cell con-446 centration was 100 cells/ml or less. Elution was 447 followed by injection of a cell suspension to deter-448mine whether detachment of previously loaded (resi-449dent) cells was enhanced by arrival of mobile cells. 450

Sediments used in the column experiments were 451obtained from the narrow channel (NC) focus area in 452Oyster, VA. The sediment is primarily composed of 453quartz, feldspar, clays, and iron and aluminum 454hydroxides in decreasing order of abundance (John-455son et al., 2000). A target average pore water velocity 456of 1 m/day for each experiment was maintained 457throughout the duration of the experiment. Flow was 458allowed to equilibrate at the target average pore water 459velocity for at least 1 pore volume (PV) prior to 460 injection of the bacterial solutions. Narrow channel 461artificial groundwater (NCAGW) was used as the 462 aqueous medium in all laboratory transport experi-463 ments (DeFlaun et al., 2001). 464

Strains were grown according to growth protocols 465described earlier (DeFlaun et al., 2001). Following 466 collection, all samples were fixed with 1% formalde-467 hyde. Since the concentrations of attached resident 468 cells in the field experiments were not known, it was 469not clear what attached bacterial concentrations would 470be reflective of field conditions. For this reason, 471 injection duration (0.2-0.5 pore volumes), injection 472 concentration (2e3 to 4e5 cells/ml), and column size 473(2.5 cm diameter by 15 cm length to 7 cm diameter by 474 53 cm length) were varied among the set of column 475experiments to yield resident (attached) cell concen-476trations ranging from about 50 to about 1.5e5 cells per 477 gram of sediment. The concentration of subsequently 478injected bacteria was also varied (2e3 to 4e5 cells/ml). 479

5. Results

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MLS SO-10 and SO-14 were sampled in July 481 2001, prior to the establishment of forced gradient 482conditions at the site, in order to determine the stained 483

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resident cell concentrations (remaining from July 484 2000 injection). Stained resident cells of both strains 485remaining from the July 2000 injection were observed 486in both sampled MLS. TAMRA/SE-stained DA001 487 488concentrations were approximately 50 cells/ml at all depths in SO-10, and were approximately 200 cells/ml 489at all depths in SO-14. CFDA/SE-stained resident OY-490107 cells were observed at low concentrations (aver-491aging ~ 20 cells/ml) at all depths in SO-10 and low 492concentrations (averaging ~ 30 cells/ml) at all depths 493in SO-14. Laboratory blanks showed no cells, proving 494that the cells did not derive from contamination during 495analysis. Tubing in all MLS was removed and 496replaced prior to monitoring for the year 2001 injec-497tion, eliminating the possibility that these cells had 498remained in the tubing during the year that passed 499between experiments. 500

501 No enhanced detachment of stained resident cells 502 was observed to occur with the arrival of injected cells 503 during the July 2001 injection. This is illustrated in 504 several sampling ports in various MLS, including SO-505 T2 P12, SO-17 P10, SO-24 P8, and SO-24 P12 (Fig. 2), where breakthrough of injected DA001 at 70 (SO-506T2 P12), 35 (SO-17 P10), and 160 (SO-24 P8 and 507P12) elapsed hours was not accompanied by increases 508in stained resident cell concentrations. SO-T2 P12 509shows an example of the background concentrations 510of stained resident cells in samples taken prior to 511breakthrough of injected cells. The background con-512centrations of stained resident cells were not observed 513in all of the MLS, including three of the four MLS 514shown in Fig. 2. Results from four other monitored 515MLS (not shown) also showed a lack of enhanced 516detachment of stained resident cells regardless of 517presence of stained resident cells, indicating that 518enhanced detachment of stained resident cells did 519not occur in response to breakthrough of injected 520cells. 521

It should be noted that the analytical resolution of 522 the analysis allowed enumeration of extremely low 523 cell concentrations during initial breakthrough, and 524 that variations among concentrations during initial 525 breakthrough represent a combination of transport 526 effects due to aquifer heterogeneity, sampling errors, 527



Fig. 2. Results from July 2001 field experiment showing breakthrough of injected DA001 and OY-107. Detachment of resident DA001 and OY-107 cells was not enhanced by breakthrough of the injected cells. Resident DA001 was observed in the aqueous phase before, during, and after extraction (e.g. SO-T2 P12).

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and analytical error. In addition to high resolution, the method provided sufficient flexibility to allow simultaneous monitoring of two strains each of two different colors (each of two different injection times) as well as their unstained counterparts in combination with any antigenically similar strains.

Unstained cell concentrations showed ephemeral 534increases coincident with the arrival of stained in-535jected cells, similarly to increases previously reported 536from the July 2000 injection (Johnson et al., 2001). 537 538MLS SO-17 P2 (Fig. 3) shows, in log (left) and linear (right) format, ephemeral increases of total 539(injected plus unstained) cells. These ephemeral 540increases were a factor of three to four times greater 541than injected cell concentrations in pulses centered at 54270, 98, and 120 elapsed hours. MLS SO-17 P4 (Fig. 5433) shows ephemeral increases of total cells, a factor 544of three to four times greater than injected cell 545concentrations in pulses centered at 106 and 145 546elapsed hours. 547

548 The unstained cells were observed only in 549 limited locations throughout the site and were exclusively observed in relatively shallow MLS 550ports (port 4 and above). No significant increases 551in total relative to injected cell concentrations were 552observed in the majority of ports analyzed, as 553shown in MLS SO-17 P6, SO-24 P8, and SO-24 554P10 (Fig. 4), where stained and total cell concen-555trations increased simultaneously with negligible 556differences (less than a factor of two) upon arrival 557of injected cells. 558

Results from the laboratory experiments also 559indicate that no enhanced detachment of resident 560cells occurred in response to the arrival of injected 561cells. Results from a representative experiment are 562shown in Fig. 5. One pore volume of stained DA001 563(2e5 cells/ml) was injected into the column (0-2)564elapsed hours), resulting in an effluent pulse from 2 565to 5 elapsed hours. Extended tailing of low concen-566trations of stained DA001 was observed during 567elution with NCAGW. In this column experiment, 568two separate reservoirs were used for delivery of the 569bacterial suspension and NCAGW, and switching 570between the two reservoirs required the use of a 571



Fig. 3. Results from July 2001 field experiment showing an ephemeral pulse of unstained cells (difference between total and stained cell concentrations) coincident with the breakthrough of injected cells. The peak unstained cell concentrations were more than a factor of two greater than the concentrations of injected cells in several ports. This result was observed only in the shallow ports.



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Fig. 4. Results from the July 2001 field experiment highlighting the lack of significant unstained cell concentrations (difference between total and stained cells) in the majority of the MLS ports sampled at the site.

three-way valve. Unlike the field experiment, switch-572573ing between reservoirs during the laboratory experi-574ment may have caused a sudden increase in hydrodynamic shear in the column. To assess the 575magnitude of enhanced detachment due to hydro-576dynamic shear, the three-way valve was quickly 577 switched between reservoirs at about 24 elapsed 578hours (Fig. 5). 579

Following the valve switch, a pulse injection of 3.5e5 cells/ml of unstained DA001 cells was then introduced into the column beginning at ~ 29 elapsed hours. The effluent pulse of stained resident cells at 24–28 elapsed hours ($\sim 1e3$ cells/ml) was associated with the valve switch. The effluent pulse 585of stained resident cells at 30-33 elapsed hours (1e2 586cells/ml) was associated with the breakthrough of 587 unstained DA001 cells. The fact that greater 588enhanced detachment of resident cells occurred dur-589ing the valve switch alone relative to the valve 590switch plus introduction of mobile cells indicates 591that the observed enhanced detachment of the resi-592dent cells in the laboratory experiment may have 593 resulted from hydrodynamic shear associated with 594the valve switch, and cannot be attributed to colli-595sion with mobile cells. It should be noted that 596 subsequent experiments conducted in a system that 597

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Fig. 5. Representative results from the laboratory column experiments showing a lack of enhanced detachment of stained resident cells. The first two vertical bars represent the time of the injection pulse for stained DA001. The third vertical bar represents the time of the valve switch. The fourth and fifth vertical bars represent the time of the injection pulse of unstained DA001 cells.

598 avoided valve switching also lacked enhanced de-599 tachment of resident cells.

600 6. Discussion

Enhanced detachment of stained resident cells in 601 602 response to arrival of injected cells was not observed in the July 2001 field injection, nor was it observed in 603 any of the laboratory experiments. Lack of enhanced 604 detachment of stained resident cells may indicate that 605the concentrations of stained resident (attached) cells 606 607 were insufficient to support significant enhanced detachment in the presence of mobile cells, assuming 608 that greater attached cell concentrations increase the 609 magnitude of enhanced detachment. The adhesion-610 611 deficient variants used in the experiments may have 612 yielded low attached cell concentrations relative to their indigenous counterparts, although the attached 613concentrations of the latter are unknown. 614

Alternatively, the results may indicate that the 615 ephemeral pulses of unstained cells observed during 616 the field experiments represent a process other than 617 enhanced detachment. Mechanisms, other than 618 619 detachment, that may explain the ephemeral increases in unstained cell concentrations (coincident with the 620 initial breakthrough of injected cells) include prefer-621 622 ential stain loss from the injected cell population 623residing in the low-concentration fringe of the plume (Johnson et al., 2001). Stain loss by diffusion from 624 cells could occur preferentially from cells residing at 625 the low-concentration fringe of the injected plume. 626 However, Fig. 6 shows that CFDA/SE-stained DA001 627 monitored using FITC-conjugated antibodies (total 628cells) and nonconjugated antibodies (stained cells) 629 degraded at very similar rates, indicating that stain 630 loss over time was negligible. 631

Notably, injection-zone samples taken during 632 injection showed concentrations of 1.3e8 and 1.2e8 633cells/ml for DA001 and OY-107, respectively, ac-634 cording to ferrographic capture using FITC-conju-635gated antibodies. This result agreed well with flow 636 cytometry results for samples collected from the 637 injection tanks, which showed DA001 concentrations 638 of 1.5e8 cells/ml. Surprisingly, concentrations of 639both strains determined ferrographically using non-640 conjugated antibodies were about ~ 55% of those 641 determined by the other methods. This latter result 642 was corroborated by direct counts on filters (Mark 643 Fuller, Envirogen, personal communication). These 644 results indicate that nearly 50% of the injected cells 645were weakly stained. The weakly stained cells in the 646injection solution may represent inefficiency in the 647staining procedure (Mark Fuller, Envirogen, personal 648 communication). 649

The presence of unstained cells in the injected 650 solution does not affect the analysis presented 651 regarding the detachment of stained resident cells. 652 The presence of unstained cells in the injection 653



Fig. 6. CFDA/SE-stained DA001 standard results observed using FITC-conjugated antibodies (total DA001) and nonconjugated antibodies (stained DA001) to DA001 cells.

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654solution does, however, explain the persistent factorof-two difference between total and unstained cell 655 656 concentrations observed in nearly all ports when cell concentrations became significant (e.g. greater than 657 100 cells/ml in Figs. 3 and 4), as well as the factor-658659 of-two difference between standards monitored using FITC-conjugated versus nonconjugated antibodies 660 (Fig. 6). The presence of unstained cells in the 661 injection solution does not, however, explain the 662 ephemeral pulses of unstained cells yielding ratios 663 of total to injected cell concentrations of three to 664 four (Fig. 3). 665

Stain loss due to cell division at the low-concen-666 tration fringe of the plume of injected bacteria repre-667 sents another potential means of generating the 668 ephemeral pulses of unstained cells. Recall that the 669 ephemeral pulses of unstained cells were observed in 670 relatively shallow ports. Indeed, cells in samples 671collected from shallow ports exhibited a polar distri-672 bution of stain that was suggestive of stain loss due to 673 674 cell division, in which the daughter cells each retained 675 stain in the portion of the cell that was originally part of the parent cell. Fig. 7 shows two images of polar 676 stained cells (left) and normal fully stained cells 677 678 (right). The cells that exhibited a polar distribution of stain would easily be missed during enumeration 679 following ferrographic capture by nonconjugated anti-680 bodies due to their relatively weak illumination. It is 681 therefore possible that the ephemeral pulses of 682 unstained cells coincident with the injected cells 683 represent stain loss due to cell division in the low-684 685 concentration fringe of the plume in the shallow portion of the flow cell. 686

Cell division in the low-concentration fringe of the 687 plume could potentially have been driven by a favor-688 able combination of electron acceptor and donor 689 exclusively at the interface of the injected and ambient 690 groundwater in the shallow portion of the aquifer. 691 Differences in the chemistries of the injected versus 692 ambient groundwater, if any, are expected to have 693 been subtle, since the injected groundwater was 694extracted from the site at a location just outside the 695 flow cell. The limited groundwater chemistry data 696 measured during the field injection indicate no sig-697 nificant variation with depth for the measured param-698 eters (including dissolved oxygen, dissolved organic 699 carbon, ionic strength, and pH). The limited data do 700 suggest, however, that dissolved ($< 0.2 \mu m$) organic 701 carbon concentrations were slightly higher in the 702ambient relative to the injected groundwater (by 1-703 10 mg/l), potentially supporting growth at the inter-704 face. However, the temporal and spatial sparseness of 705 the chemical data prevent rigorous identification of 706 potential chemical drivers for cell division at the 707 interface between ambient and injected groundwater, 708 and so this determination is beyond the capabilities of 709the present study. 710

This investigation highlights the resolution of fer-711 rographic capture, which clearly showed that 712enhanced detachment of stained resident cells did 713not occur in response to breakthrough of injected cells 714in our system. This result weakens the original 715hypothesis that hydrodynamic collision between 716 mobile and attached bacteria resulted in the observed 717 ephemeral domination by unstained cells during initial 718 breakthrough. 719



Fig. 7. Images of DA001 cells from the July 2001 field experiment showing polar distribution of stain in relatively dim cells, suggestive of cell division (a). Staining normally results in homogeneously bright cells (b).

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720 7. Uncited reference

721 Hendry et al., 1999

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