UC Santa Barbara

UC Santa Barbara Previously Published Works

Title

Investigation of Groundwater Flow Variations near a Recharge Pond with Repeat Deliberate Tracer Experiments

Permalink

https://escholarship.org/uc/item/79p8j31x

Journal

Water, 6(6)

ISSN

2073-4441

Authors

Clark, Jordan Morrissey, Sheila Dadakis, Jason et al.

Publication Date

2014-06-17

DOI

10.3390/w6061826

Peer reviewed



Article

Investigation of Groundwater Flow Variations near a Recharge Pond with Repeat Deliberate Tracer Experiments

Jordan F Clark 1,*, Sheila Morrissey 1, Jason Dadakis 2, Adam Hutchinson 2 and Roy Herndon 2

- Department of Earth Science, University of California, Santa Barbara, CA 93106, USA; E-Mail: skmbruin@gmail.com
- Orange County Water District, Fountain Valley, CA 92728, USA; E-Mails: JDadakis@ocwd.com (J.D.); AHutchinson@ocwd.com (A.H.); RHerndon@ocwd.com (R.H.)
- * Author to whom correspondence should be addressed; E-Mail: jfclark@geol.ucsb.edu; Tel.: +1-805-893-7838; Fax: +1-805-893-2314.

Received: 25 January 2014; in revised form: 30 May 2014 / Accepted: 3 June 2014 /

Published: 17 June 2014

Abstract: Determining hydraulic connections and travel times between recharge facilities and production wells has become increasingly important for permitting and operating managed aquifer recharge (MAR) sites, a water supply strategy that transfers surface water into aquifers for storage and later extraction. This knowledge is critical for examining water quality changes and assessing the potential for future contamination. Deliberate tracer experiments are the best method for determining travel times and identifying preferential flow paths between recharge sites over the time scales of weeks to a few years. This paper compares the results of two deliberate tracer experiments at Kraemer Basin, Orange County, CA, USA. Results from the first experiment, which was conducted in October 1998, showed that a region of highly transmissive sedimentary material extends down gradient from the basin for more than 3 km [1]. Mean groundwater velocities were determined to be approximately 2 km/year in this region based on the arrival time of the tracer center of mass. A second experiment was initiated in January 2008 to determine if travel times from this basin to monitoring and production wells changed during the past decade in response to new recharge conditions. Results indicate that flow near Kraemer Basin was stable, and travel times to most wells determined during both experiments agree within the experimental uncertainty.

Keywords: managed aquifer recharge (MAR); groundwater flow and management; geochemical tracers; Orange County Water District (CA, USA)

1. Introduction

Groundwater has been a primary source of potable and irrigation water for centuries. During the last 50 years, the soaring demand for freshwater has placed unprecedented stresses upon many aquifers throughout the world; many aquifers are now in overdraft. The projected growth in population combined with uncertainties associated with a changing climate will only aggravate this problem. A cost-effective advancement in groundwater/surface water management aimed at augmenting local water supplies is managed aquifer recharge (MAR), the practice of artificially recharging imported surface water, reclaimed (recycled) wastewater, or storm runoff into aquifers for storage and later extraction [2–4].

For a MAR operation to be successful, a few challenges must be overcome. First, a source of recharge water must be found. Second, facilities, which can rapidly transfer water into aquifers, must be engineered. These include injection wells and spreading basins (recharge ponds), with both requiring periodic removal of accumulated clogging material to maintain recharge rates. Third, because the available water for the recharge operation can be of lesser quality than the local groundwater supply, there is the potential to degrade the existing aquifer. The introduction of salts, pathogens, disinfection by-products, and trace organic compounds such as pharmaceuticals, is a concern, especially when urban runoff or reclaimed wastewater is a large component of the source water for the operation [4]. Lesser quality sources can become a larger portion of the recharge water at MAR operations because the availability of higher quality water such as imported water from remote watersheds is limited and may shrink due to shifts in climate and the diversion of this water to other uses such as maintaining riparian ecosystems (i.e., loss of California State Water Project supplies to the delta smelt). Even in cases where the recharge water may be of a higher quality, it may have a different geochemical character (e.g., redox state) than the ambient groundwater and the potential exists for mobilization of intrinsic constituents, such as arsenic. For these reasons, it is vital to understand the fate and transport of potential contaminants near MAR sites. Only from this understanding can cost effective and appropriate regulations be developed.

Recent water quality studies near MAR operations have shown that many potential contaminants such as dissolved organic carbon (DOC), nitrate, some pharmaceuticals, and most pathogens are naturally removed or become inactive with time and distance in the subsurface (e.g., [4–9]). These water quality improvements, known as soil-aquifer treatment (SAT), are considered one of the benefits of MAR [3,4] and are generally observed soon after recharge, near the facility.

Despite these studies, water quality concerns still remain the focus of many regulations and an obstacle for the permitting of new MAR operations that include a reuse component (*i.e.*, reclaimed wastewater). For instance, MAR operations in California, USA, must conform to the state Drinking Water Program's Groundwater Recharge Reuse Regulations, which require a subsurface retention time prior to its extraction for a potable supply that varies based on the level of pre-recharge treatment, in

addition to a number of source water controls and how the travel times is being assessed [10]. The required retention time varies from two to six months and must be verified using a tracer study.

1.1. Travel Time Estimates

Development of field methodologies to evaluate flow near MAR facilities is critical for the effective management of these operations where water quality assessment is warranted. The California Drinking Water Program's Groundwater Recharge Reuse Regulations place a greater level of confidence on tracer data than on numerical models or Darcy flow calculations (see Table 2 in reference [10]). Geochemical techniques provide a fundamental approach for investigating travel times, flow paths, recharge rates, and dispersivity in groundwater. Intrinsic environmental tracers such as Tritium/³He (T/³He) dating (e.g., [11,12]) are ideal for examining flow over spatial scales of kilometers and temporal scales of years to decades near MAR operations [1,13,14]. Other environmental tracer dating techniques such as chlorofluorocarbons (CFCs) or sulfur hexafluoride (SF₆) are likely to be unreliable near MAR operations because non-atmospheric sources of these tracers may complicate the interpretation of apparent ages, especially when chlorinated reclaimed or potable water is being recharged (e.g., [15–17]).

Because of the typical uncertainty associated with geochemical dating techniques (generally no better than ± 2 years), these methods are not well suited for examining short-term transport specified by the Recharge Reuse Regulations in the state of California [14]. Deliberate (*i.e.*, injected or applied) tracer experiments using gases such as SF₆ or noble gases (*i.e.*, He or Xe isotopes) have become acceptable methods for evaluating transport from recharge locations to wells over periods of weeks to a few years [1,10,18,19]. There is a long history of using SF₆ as a deliberate tracer starting with atmospheric experiments and expanding into aqueous systems initially by the gas exchange and oceanographic communities [20–23]. SF₆ can also be used as an intrinsic or environmental tracer for dating water in a similar fashion as CFCs [17].

The scale of deliberate tracer experiments is defined by the quantity of water that can be "tagged" and the signal to noise ratio of the tracer being used. The three factors that often limit an experiment's scale are: (1) identification of a tracer that does not adversely impact potable aquifers; (2) the cost of tracer; and (3) the ability to introduce a sufficient amount of tracer without significantly changing the buoyancy (or density) of the tagged water. The cost of the tracer can be a particular problem when large volumes of water (> 10^5 m³ or >80 AF) need to be tagged, as is often the case near MAR operations. As shown by the oceanographic community (e.g., [20]), SF₆ can be used economically to tag large volumes of water.

There are cost advantages of using SF₆ over noble gas isotopes in terms of analysis; more SF₆ samples can be analyzed over a given period of time on less expensive equipment. However, it is a strong greenhouse gas and its emission is regulated in California. SF₆ is a synthetic gas used primarily in the electrical industry as a gas insulator and has been used as a tracer in the atmosphere and natural waters for more than two decades. It is an ideal tracer for the following reasons: (1) SF₆ is nontoxic [24] and permission has been granted to use it as a tracer in potable aquifers in the southwestern USA and South Australia; (2) background concentrations of SF₆ in natural waters are extremely low (<0.05 pmol/L; 1 pmol = 10^{-12} mole) because of its low solubility and low atmospheric mixing ratio (*ca.* 8 pptv in 2014); (3) it can be measured precisely in water (±5% or better) over a concentration

range of eight orders of magnitude (0.01 fmol/L to 1 nmol/L) using a gas chromatograph equipped with an electron capture detector [25]; and (4) laboratory scale experiments have shown that breakthrough curves of SF_6 and bromide are identical in saturated porous media which contains either high contents of organic material or clays, demonstrating that SF_6 is not retarded (has a retardation factor of 1.0) [26,27].

SF₆ differs from ionic and dye tracers in that it is a gas and is easily lost from solution across the air—water interface to the atmosphere. It has often been used as a tracer for reaeration in surface water [21–23,28]. Thus, it is important to monitor the SF₆ concentrations in the recharge water to determine the amount of tracer lost due to gas exchange. This is especially true if the artificial recharge is taking place in a shallow river or spreading pond. Insufficient monitoring of the spatial variability of the tracer concentrations in the "spiked" surface water can lead to erroneous interpretations of travel times. Furthermore, laboratory column experiments have shown that SF₆ transport is slowed (retarded) when trapped air is contained within the porous media [29,30]. While there has been some evidence of gas tracer loss during recharge when a significant vadose zone is present, experiments conducted at three sites in California with a long history of fairly continuous recharge (OCWD, Orange County; Montebello Forebay, LA County; and El Rio, Ventura County) have shown that gas loss during percolation from spreading ponds is manageable if the tracer is introduced properly [1,13,16,18].

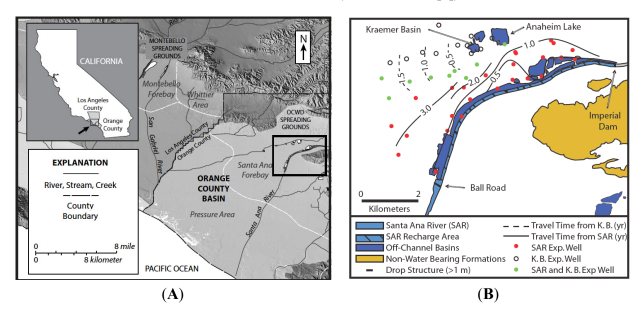
Fundamental questions concerning deliberate tracer experiments at MAR sites include: How general are the results? Can the results of an experiment performed several years in the past be used in the future? These questions arise because groundwater flow and travel times, e.g., to particular wells, should reflect hydrologic conditions at the time of the experiment such as recharge rates both at the MAR operation and surrounding area, pumping rates of nearby wells, and the regional hydraulic gradient. All of these are likely to change over time scales of months to years and imply that deliberate tracer results may not be similar year to year. For instance, long-term changes in the demographics and economics of an area could lead to changes in the locations and extraction/recharge rates from facilities. Furthermore, in areas with a Mediterranean climate such as Southern California, it is likely that recharge and groundwater production are out of phase, with the wet season having more recharge and less production. This should create seasonal variations in groundwater flow that should affect travel times, especially at shorter time scales (*i.e.*, months).

In this paper, we discuss the results of two deliberate tracer experiments conducted at a spreading basin in Southern California that is operated by the Orange County Water District (OCWD), ten years apart to assess if travel times changed significantly over a decade.

1.2. Field Site

A 9-km section of the Santa Ana River (SAR) and a series of spreading basins, including Kraemer Basin and Anaheim Lake, located near Anaheim, California USA are used by the OCWD as principal recharge locations for the Orange County groundwater basin (Figures 1 and 2). For more than 75 years, OCWD has been actively replenishing and managing the groundwater basin that supplies about 70% of the total water demand of approximately 2.4 million people and has been actively managing and replenishing the local groundwater basin. Currently, it operates more than 400 hectares of surface spreading facilities and recharges approximately 3.5×10^8 m³ (2.8×10^5 AF; 10-year average between 2000 and 2010) per year [31].

Figure 1. Maps of the Orange County managed aquifer recharge (MAR) facilities. (A) Topographic map of the field area including the location of Orange County Water District (OCWD) recharge basins based on data from the United States Geological Survey (USGS) 10-m National Elevation Dataset [32]; (B) Recharge occurs from the basins and from the Santa Ana River (SAR) between the Imperial Dam and Ball Road. The first arrival times of tracers determined in 1998 are contoured (modified from [1]).



In January 2008, OCWD completed and began operating the Groundwater Replenishment System (GWRS), which is the world's largest wastewater purification system for indirect potable reuse [33]. The system produces up to 2.43×10^5 m³ (197 AF) of high quality reclaimed water each day using a three-step advanced treatment process that consists of microfiltration, reverse osmosis, and ultraviolet light disinfection with hydrogen peroxide advanced oxidation. A variable portion of the reclaimed water is pumped about 20 km to the MAR facilities and is recharged through Kraemer and other permitted nearby basins to replenish the groundwater.

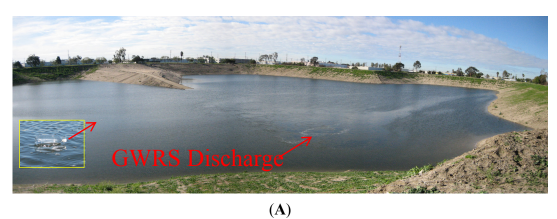
Detailed investigations of the groundwater flow were conducted in the late 1990s using a variety of techniques including T/3He dating and deliberate tracer experiments at Anaheim Lake, Kraemer Basin, and the SAR [1,34]. The January 08 SF₆ experiment was required by state regulators because of the use of Kraemer Basin for GWRS water recharge and to determine if travel times near Kraemer Basin were similar to those determined a decade earlier during the October 1998 Xe isotope tracer experiment.

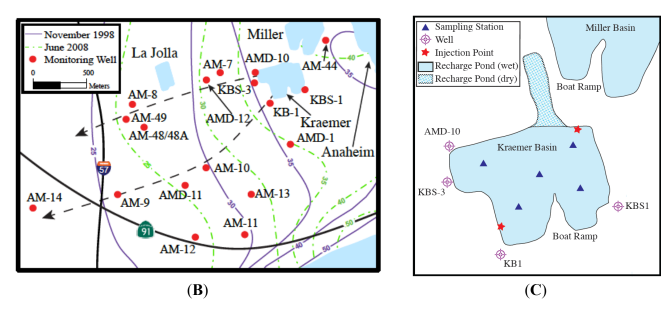
2. Materials and Methods

The methodology of Clark *et al.* [1] was used during the January 2008 Kraemer Basin deliberate tracer experiment and is outlined below. At the time of the tracer injection, the basin contained approximately $1.4 \times 10 \text{ m}^5$ (2000 AF) of water. The volume increased during the experiment and by the end, the pond contained about $2.1 \times 10 \text{ m}^5$ (3000 AF). The study began on 17 January 2008 (defined as day 0) when 99.8% pure SF₆ gas tracer was carefully injected into Kraemer Basin over a period of about 1 hour by bubbling the tracer through a submerged diffusion stone at a rate of about 40 mL/min at two locations $\sim 10 \text{ m}$ offshore (Figure 2C). This injection technique was repeated twice more, on day 8 and 11. During each injection, SF₆ formed a rising bubble plume that only

partially dissolved in the water column (bubbles could be seen bursting at the surface); therefore, an unknown quantity of tracer was released into the recharge water. Previously, it was estimated that less than 5% of the bubbled gas dissolves [14,18,19,34].

Figure 2. Maps of the Orange County MAR facilities. (**A**) Photograph of the basin looking from the northern shore to the south towards the boat ramp. The inserted photo shows one of the fixed buoys; (**B**) Detailed map of the January 2008 field area showing the sampled wells and principle recharge areas. At the start of the experiment, neither Miller nor La Jolla Basins were recharging the aquifer. For reference, the north-south 57 freeway and the east-west 91 highway have been included. The black dashed arrows represent the northern and southern flow lines, the solid purple and dashed/dotted green contours represent, respectively, the and November 1998 and June 2008 piezometric surfaces; (**C**) Detail of Kraemer Basin at the time of the 2008 tracer injection showing where tracer was introduced and sampled.





Because it is vital to know the initial dissolved tracer concentration, SF₆ surveys of Kraemer Basin water were conducted on days 1, 3, 5, 8, 9, 11, 12, 14, 18, and 22. During each survey, surface (~1 m below the pond's surface) and bottom (~1 m above sediment) samples were collected from five stations evenly distributed throughout the basin and marked with fixed buoys (Figure 2A,C). Approximately 2–3 mL samples were collected in VacutainersTM in triplicate for storage and later

analysis. VacutainersTM are convenient storage and reaction containers that are commercially available. Surveys were conducted until the mean SF₆ concentration decreased to approximately the detection limit. Following the tracer injection, well samples were collected for a period of one year by personnel from OCWD in VacutainersTM and sent to UCSB for analyses, which generally occurred within two weeks of collection. The frequency of sampling was adjusted for each well based on the results of recent sampling.

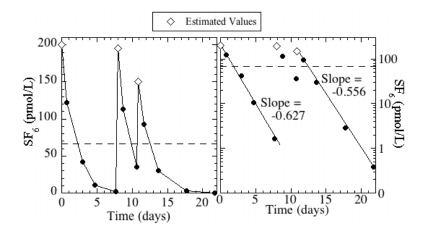
All SF₆ samples were analyzed using a headspace method similar to that described by [1]. In the field, pre-weighed 10 mL VacutainersTM were partially filled (2–5 mL of water). In the laboratory, these containers were weighed (to determine the sample weight) and carefully filled with ultra-high purity nitrogen gas (so that the final pressure was equal to about 1 atmosphere). After a brief shaking to mix the headspace, this gas was injected through a column of $Mg(ClO_4)_2$ (to remove water vapor) into a small sample loop of known volume (about 1.5 mL). Subsequently, the gas in the sample loop was flushed into a gas chromatograph equipped with an electron capture detector with ultra-high purity nitrogen carrier gas. SF₆ was separated from other gases with a molecular sieve 5a column held at room temperature. The detector response was determined by running gas standards purchased from and certified by Scott-Marrin, Inc. (Riverside, CA, USA). The detection limit of this method is about 0.05 pmol/L, three orders of magnitude lower than the mean pond concentration (see below). Error on duplicate VacutainerTM measurements was typically better than $\pm 10\%$ but not as good as the $\pm 5\%$ reported for the syringe headspace method [25].

At each well, arrival times of tracer can be determined by evaluating breakthrough curves, which are plots of concentration *versus* time. In homogenous aquifers—sand boxes, the initial and mean arrival (center of mass or COM) times of tracer at narrow-screened monitoring wells represent, respectively, the fastest and mean flow paths in the aquifer. In heterogeneous aquifers with preferential flow paths tracer breakthrough curves are more complicated, often showing multiple peaks [1,14]. Tailing is also evident on breakthrough curves and can represent the tracer reaching the well by slower flow paths, back diffusion out of lower permeability strata, non-ideal tracer input function, or retardation, which in the case of gas tracers can be due to trapped gas [29,30]. As discussed by Becker *et al.* [35], sampling biases can result in moderate and hard to quantify travel time errors. These biases are often caused by infrequent sampling due to budget limitation.

3. Results and Discussion

Mean basin concentrations of SF₆ tracer determined for each survey ranged between about 0.4 pmol/L (day 22) and 120 pmol/L (day 9); the daily infiltration rate varied between 1.7 m³/s (day 20) and 2.2 m³/s (days 6 and 7) and averaged 1.95 ± 0.15 m³/s (68.9 ± 5.3 cfs). The basin concentrations were the highest following the injections and decreased exponentially due to recharge and gas loss across the air-water interface (Figure 3). The rate of loss was slightly greater during the first week than during the second, most likely due to the progressive deepening of the basin as it was filling with recharge water. This would increase the mean residence time of water in the basin and decrease the gas exchange loss.

Figure 3. SF₆ concentrations in Kraemer Basin during the injections and subsequent monitoring periods. Injections occurred on days 0, 8, and 11; estimated basin concentrations are plotted for these days. The dashed line represents the 14-day flow weighted mean concentration (66 pmol/L) during the defined injection period.



The tracer injection period was defined as the first 14 days (between January 17 and 31) when 96% of the total mass injected percolated into the ground and the flow-weighted mean SF₆ concentration was 66 pmol/L. It is important to note that the basin concentration decreased below 10% of the mean for 2 days (days 6 and 7) in the middle of the injection period (Figure 3). This complexity in the basin concentration is apparent in the breakthrough at monitoring well KBS-3/1, which displays two peaks (Figure 4B).

Although the water level contours represent different seasons and therefore different water use times in terms of the domestic landscape irrigation cycle, they are sub-parallel indicating that the location of recharge and pumping remained similar between experiments (Figure 2B). However, the hydraulic gradient west-southwest of Kraemer Basin was ~50% steeper in November 1998 (~0.0064) than June 2008 (~0.0042). Furthermore, the absolute elevation of the piezometric surface was about 5 m higher in June. These changes most likely reflect seasonality rather than long-term use patterns. The June survey occurred near the beginning of heavy domestic landscape irrigation while the November survey reflects water levels at the end of this irrigation period.

Groundwater samples were collected at 28 monitoring points (at 17 well sites, three of which were multi-level: AMD-10, AMD-11, AMD-12; Figure 2B), with sufficient frequency to construct tracer breakthrough curves (Figure 4). Tracer was detected at twelve wells down gradient (to the west) of Kraemer Basin (Table 1). As was observed during the October 1998 Xe isotope tracer study [1,33], detections progress systematically to the west along two flow paths that follow the local hydraulic gradient: the northern path of KBS-3/1, AM-7, AMD-12/1, AM-48, and AM-8; and the southern path of KBS-1/1, KB-1, AMD-11/1, AM-10, AM-9, and AM-14 (Figure 2B). The one exception is the tracer's first arrival was essentially the same at the deeper screened AMD-11/1 and AM-10 even though AMD-11/1 was more distant. This implies that depth as well as distance is important when considering travel times and that the first arrival is a difficult parameter to interpret. Interestingly, the arrival of the COM followed the distance trend with it arriving at AM-10 prior to AMD-11/1.

Figure 4. Breakthrough curves at representative wells. Wells found along, respectively, the southern (**A**, **C**, **E**) and northern (**B**, **D**, **F**) flow paths are displayed on the left and right. The mean pond concentration was 66 pmol/L and non-detections (<0.05 pmol/L) have been plotted as "0 pmol/L". Please note that the scale on the y-axes (concentration) differs.

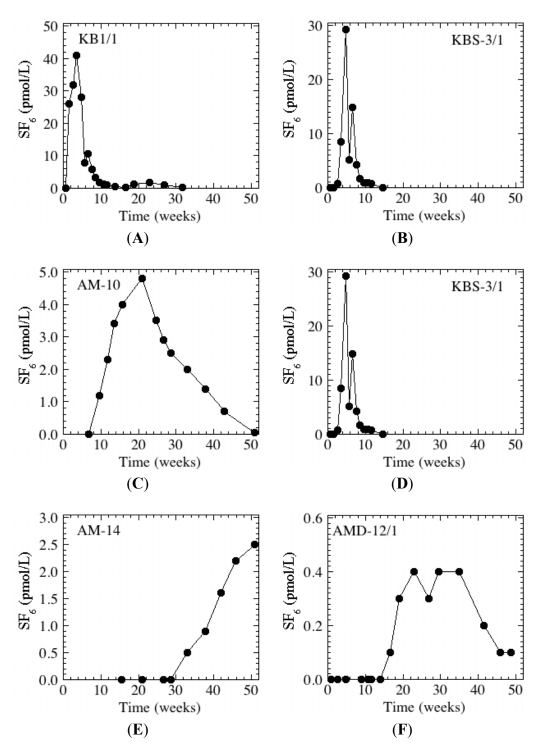


Table 1. Summary of travel times during the Oct-98 136 Xe and Jan-08 SF₆ tracer experiments from Kraemer Basin. The Oct-98 Xe isotope data is from [33]. The distance was measured from the basin's shoreline directly to the well. This is the shortest distance and does not necessarily represent the flow path length. Travel times are in weeks.

Well	Distance (m)	Screen Interval (m msl)*	Jan-08 SF ₆			Oct-98 ¹³⁶ Xe		
			First Detect	Peak	COM	First Detect	Peak	COM
Northern Flow Path								
KBS-3/1	<100	44 to 41	2.7	4.7	5.6	<1.4	<1.4	†
AM-7	130	1 to −4	10.6	22.9	25.4	8.3	15.1	15.3
AMD-12/1	525	−36 to −42	16.6	22.9	30.5	_	_	
AM-48	1250	−20 to −29	<16.6	18.7	25.7		_	
AM-8	1250	−26 to −31	20.9	_	37.1 †	17.7	23.0	38.3
Southern Flow Path								
KBS-1/1	<100	4 to 1	<1.6	<1.6	†	1.4	1.4	
KB1	<100	13 to 7	1.6	3.6	6.0	2.6	2.3	3.7
AM-10	1000	−3 to −8	6.6	20.9	23.8	15.1	23.0	28.4
AMD-11/1	1260	−91 to −97	6.7	29.1	>28*			
AM-9	1840	−26 to −31	15.7	26.6	33.8	26.4	39.7	50.6
AM-14	2630	−32 to −38	37.7	37.7	†	45.8		67.9

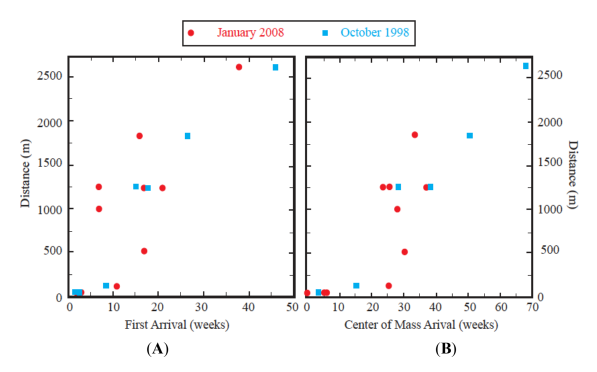
Notes: * Depth is measured relative to mean sea level (msl). The ground surface elevation of Kraemer Basin is about 68 m and the basin floor has an elevation of about 53 m. During the tracer injection period, the surface of the water increased from about 57 m to 59 m; † Incomplete breakthrough, center of mass (COM) travel time is a minimum or could not be calculated.

At relatively shallow wells (<65 m to the screen top) near the basin (KB-1, KBS-1, KBS-3), tracer was first detected <1 to 3 weeks after the beginning of the injection period. This was not the case at another nearby well, AMD-10, where the tracer was never detected in the five zones sampled, which have screen tops between 90 and 285 m below ground surface. This is in agreement with the October 1998 experiment [1,33], which showed that these zones are hydraulically connected with Anaheim Lake (Figures 1 and 2B), a recharge basin up gradient (to the east) of Kraemer Basin, once again indicating the importance of depth to the flow field.

The influence of depth is best explained by the complicated local hydrostratigraphy. The water recharged from the basins is not flowing through a sand box, rather it flows through conductive layer between confining and semi-confining layers [1,33,36]. This complicated hydrostratigraphy also helps to explain the reversal of arrival time (first vs. COM) at AM-10 and AMD-11/1 and points to a potential problem of using a tracer with a large signal to noise ratio. There is no doubt that the first arrival time is poorly defined: the arrival is defined by the first detection of the tracer and therefore is defined by which tracer is being used and how good the analytical system is. Therefore the signal to noise ratio of the tracer is vital for determining this arrival time and not the local hydrology. It may make sense in the future to define this arrival time with C/C_0 , where C_0 could be either the initial concentration in the recharge water or the local peak concentration. We believe that the former makes more sense because in is likely that the local peak concentration cannot capture unless continuous sampling is employed.

The travel times of the leading edge of the tracer arrival, which is defined by the first detection, were very similar (within three weeks) at six of the nine wells monitored during both the 2008 and 1998 experiments (Figure 5A; Table 1). The mean observed velocity of the leading edge was about 60 m/week (3 km/year). The exceptions were the three most distant wells along the southern flow path, AM-9, AM-10, and AM-14. During the 2008 experiment, the tracer arrival was traveling about 50% faster and, therefore, earlier detections were observed at these wells.

Figure 5. Comparison between the arrival times and distance from the January 2008 and October 1998 tracer experiments. (**A**) First arrival and (**B**) COM. Note the change of scale on the x-axis. During the January 2008 experiment, COM arrivals could not be calculated at all wells because of incomplete breakthroughs (see Table 1).



Monitoring wells AM-8, AM-48, AM-48A, and AM-49 are located along the northern flow path approximately the same distance down gradient. However, the screen depth of these wells differs. Tracer was not detected at the relatively shallow wells, AM-48A and AM-49 (screen depths between 20 and 30 m msl) while it was detected at the deep wells, AM-8 and AM-48 (screen depths between -20 and -31 m msl). This implies that the tracer had migrated vertically downward beneath the water table and was traveling through deep layers, with the shallow layers recharged by other sources, such as the nearby La Jolla Basin (Figure 2B). The movement of the COM was also very similar during two experiments (Figure 5B) and traveled with a velocity of about 40 m/week (2 km/year).

4. Conclusions

The results of the two deliberate tracer experiments conducted a decade apart were very similar despite differences in the local piezometric surface. The arrival times at nine of the twelve wells were nearly identical once the experimental uncertainty (*i.e.*, frequency of sampling) is considered. Apparently the hydraulic gradients near Kraemer Basin had not changed significantly between

experiments. This may be partly due to the fact that both experiments were conducted during similar times of high recharge at Kraemer Basin and reduced seasonal pumping in the winter-spring study period. As seen near other MAR facilities, vertical flow is important and must be considered when evaluating travel time information.

Acknowledgments

We would like to thank Alex Cervantes, David Field, Brian Okey, Nira Yamachika, Gary Yoshiba, and many others at Orange County Water District (OCWD) for hosting this research and providing assistance over many years. Daniel Petersen assisted with the field and laboratory work at Universoty of California Santa Barbara. Tim Becker helped draft some of the figures. The comments and suggestions of the two anonymous reviewers improved the manuscript considerably. Funding was provided by OCWD.

Author Contributions

All authors contributed equally to this paper.

Conflicts of Interest

The authors declare no conflict of interest.

References

- 1. Clark, J.F.; Hudson, G.B.; Davisson, M.L.; Woodside, G.; Herndon, R. Geochemical imaging of flow near an artificial recharge facility, Orange County, CA. *Ground Water* **2004**, *42*, 167–174.
- 2. Bouwer, H. Artificial recharge of groundwater: Hydrogeology and engineering. *Hydrogeol. J.* **2002**, *10*, 121–142.
- 3. Dillon, P. Future management of aquifer recharge. *Hydrogeol. J.* **2005**, *13*, 313–316.
- 4. National Research Council. *Prospects for Managed Underground Storage of Recoverable Water*; National Academy Press: Washington, DC, USA, 2007.
- 5. Yates, M.V.; Yates, S.R. A comparison of geostatistical methods for estimating virus inactivation rates in ground water. *Water Res.* **1987**, *21*, 1119–1125.
- 6. Fox, P.K.; Narayanaswamy, A.; Genz, A.; Drewes, J.E. Water quality transformations during soil aquifer treatment at the Mesa Northwest Water Reclamation Plant, USA. *Water Sci. Technol.* **2001**, *43*, 343–350.
- 7. Drewes, J.E.; Heberer, T.; Rauch, T.; Reddersen, K. Fate of pharmaceuticals during groundwater recharge. *Groundw. Monit. Remediat.* **2002**, *23*, 64–72.
- 8. Hiscock, K.M.; Grischeck, T. Attenuation of groundwater pollution by bank filtration. *J. Hydrol.* **2002**, *266*, 139–144.
- 9. Fox, P.K.; Makam, R. Surface area and travel time relationships in aquifer treatment systems. *Water Environ. Res.* **2009**, *81*, 2337–2343.

10. Johnson, T.A. Groundwater recharge using recycled municipal wastewater in Los Angeles County and the California Department of Public Health's draft regulations on aquifer retention time. *Ground Water* **2009**, *47*, 496–499.

- 11. Schlosser, P.; Stute, M.; Dörr, H.; Sonntag, C.; Münnich, K.O. Tritium/³He dating of shallow groundwater. *Earth Planet. Sci. Lett.* **1988**, *89*, 353–362.
- 12. Solomon, D.K.; Cook, P.G. ³H and ³He. In *Environmental Tracers in Subsurface Hydrology*; Cook, P.G., Herczeg, A.L., Eds.; Kluwer Academic Publishers: Boston, MA, USA, 2000; pp. 397–424.
- 13. Massman, G.; Sültenfuss, J.; Knappe, A.; Dünnbier, U.; Taute, T.; Pekdeger, A. Investigation of groundwater residence times during bank filtration in Berlin: A multi-tracer approach. *Hydrol. Process.* **2008**, *22*, 788–801.
- 14. McDermott, J.A.; Avisar, D.; Johnson, T.A.; Clark, J.F. Groundwater travel times near spreading ponds: Inferences from geochemical and physical approaches. *J. Hydrol. Eng. ASCE* **2008**, *13*, 1021–1028.
- 15. Schultz, T.R.; Randall, J.H.; Wilson, L.G.; Davis, S.N. Tracing sewage effluent recharge—Tucson, Arizona. *Ground Water* **1976**, *14*, 463–470.
- 16. Clark, J.F.; Smethie, W.M., Jr.; Simpson, H.J. Chlorofluorocarbons in the Hudson estuary during summer months. *Water Resour. Res.* **1995**, *31*, 2553–2560.
- 17. Busenberg, E.; Plummer, L.N. Dating young groundwater with sulfur hexafluoride: Natural and anthropogenic sources of sulfur hexafluoride. *Water Resour. Res.* **2000**, *36*, 3011–3030.
- 18. Gamlin, J.D.; Clark, J.F.; Woodside, G.; Herndon, R. Large-scale tracing of ground water with sulfur hexafluoride. *J. Environ. Eng. ASCE* **2001**, *127*, 171–174.
- 19. Clark, J.F.; Hudson, G.B.; Avisar, D. Gas transport below artificial recharge ponds: Insights from dissolved noble gases and a dual gas (SF₆ and ³He) tracer experiment. *Environ. Sci. Technol.* **2005**, *39*, 3939–3945.
- 20. Ledwell, J.R.; Watson, A.J.; Law, C.S. Evidence for slow mixing across the pycnocline from an open-ocean tracer-release experiment. *Nature* **1993**, *364*, 701–703.
- 21. Wanninkhof, R.; Ledwell, J.R.; Broecker, W.S. Gas exchange-wind speed relationship measured with sulfur hexafluoride on a lake. *Science* **1985**, *227*, 1224–1226.
- 22. Clark, J.F.; Wanninkhof, R.; Schlosser, P.; Simpson, H.J. Gas-exchange rates in the tidal Hudson River using a dual tracer technique. *Tellus Ser. B Chem. Phys. Meteorol.* **1994**, *46*, 274–285.
- 23. Hibbs, D.E.; Parkhill, K.L.; Gulliver, J.S. Sulfur hexafluoride gas tracer studies in streams. *J. Environ. Eng. ASCE* **1998**, *124*, 752–760.
- 24. Lester, D.; Greenberg, L.A. The toxicity of sulfur hexafluoride. *Arch. Ind. Hyd. Occup. Med.* **1950**, *2*, 348–349.
- 25. Wanninkhof, R.; Ledwell, J.R.; Watson, A.J. Analysis of sulfur hexafluoride in seawater. *J. Geophys. Res.* **1991**, *96*, 8733–8740.
- 26. Wilson, R.D.; Mackay, D.M. The use of sulfur hexafluoride as a conservative tracer in saturated sandy media. *Ground Water* **1993**, *31*, 719–724.
- 27. Wilson, R.D.; Mackay, D.M. SF₆ as a conservative tracer in saturated media with high intragranular porosity or high organic carbon content. *Ground Water* **1996**, *34*, 241–249.

28. Wanninkhof, R.; Mulholland, P.J.; Elwood, J.W. Gas exchange rates for a first-order stream determined with deliberate and natural tracers. *Water Resour. Res.* **1990**, *26*, 1621–1630.

- 29. Fry, V.A.; Istok, J.D.; Semprini, L.; O'Reilly, K.T.; Buscheck, T.E. Retardation of dissolved oxygen due to a trapped gas phase in porous media. *Ground Water* **1995**, *33*, 391–398.
- 30. Vulava, V.M.; Perry, E.B.; Romanek, C.S.; Seaman, J.C. Dissolved gases as partitioning tracers for determination of hydrogeological parameters. *Environ. Sci. Technol.* **2002**, *36*, 254–262.
- 31. 2009-10 Report on Groundwater Recharge in Orange County Groundwater Basin. Available online: http://www.ocwd.com/Portals/0/Pdf/09-10AnnualRechargeReport_All.pdf (accessed on 9 June 2014).
- 32. Gesch, D.B. The National Elevation Dataset. In *Digital Elevation Model Technologies and Applications: The DEM Users Manual*, 2nd ed.; Maune, D., Ed.; American Society for Photogrammetry and Remote Sensing: Bethesda, MD, USA, 2007; pp. 99–118.
- 33. Groundwater Replenishment System. Available online: www.gwrsystem.com (accessed on 5 April 2011).
- 34. Davisson, M.L.; Hudson, G.B.; Clark, J.F.; Woodside, G.; Herndon, R. Final report on isotope tracer investigations in the forebay of the Orange County groundwater basin. Lawrence Livermore Nation Laboratory UCRL-TR-201735, 2004. Available online: https://e-reports-ext.llnl.gov/pdf/303507.pdf (accessed on 14 April 2008).
- 35. Becker, T.E.; Clark, J.F.; Johnson, T.A. Heat ¹⁰B-enriched boric acid, and bromide as recycled groundwater tracers for Managed Aquifer Recharge: A case study. *J. Hydrol. Eng. ASCE*, submitted.
- 36. Thomson, A.F.B.; Carle, S.F.; Rosenberg, N.D.; Maxwell, R.M. Analysis of groundwater migration from artificial recharge in a large urban aquifer: A simulation perspective. *Water Resour. Res.* **1999**, *35*, 2981–2998.
- © 2014 by the authors; licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution license (http://creativecommons.org/licenses/by/3.0/).