



Critical Review

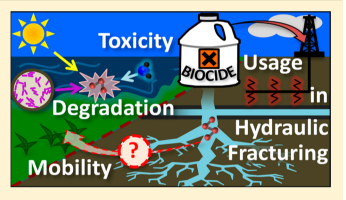
pubs.acs.org/est

Biocides in Hydraulic Fracturing Fluids: A Critical Review of Their Usage, Mobility, Degradation, and Toxicity

Genevieve A. Kahrilas, †, Jens Blotevogel, ‡, Philip S. Stewart, and Thomas Borch*,†,||

Supporting Information

ABSTRACT: Biocides are critical components of hydraulic fracturing ("fracking") fluids used for unconventional shale gas development. Bacteria may cause bioclogging and inhibit gas extraction, produce toxic hydrogen sulfide, and induce corrosion leading to downhole equipment failure. The use of biocides such as glutaraldehyde and quaternary ammonium compounds has spurred a public concern and debate among regulators regarding the impact of inadvertent releases into the environment on ecosystem and human health. This work provides a critical review of the potential fate and toxicity of biocides used in hydraulic fracturing operations. We identified the following physicochemical and toxicological aspects as well as knowledge gaps that should be considered when selecting



biocides: (1) uncharged species will dominate in the aqueous phase and be subject to degradation and transport whereas charged species will sorb to soils and be less bioavailable; (2) many biocides are short-lived or degradable through abiotic and biotic processes, but some may transform into more toxic or persistent compounds; (3) understanding of biocides' fate under downhole conditions (high pressure, temperature, and salt and organic matter concentrations) is limited; (4) several biocidal alternatives exist, but high cost, high energy demands, and/or formation of disinfection byproducts limits their use. This review may serve as a guide for environmental risk assessment and identification of microbial control strategies to help develop a sustainable path for managing hydraulic fracturing fluids.

■ INTRODUCTION

Biocides are widely used in food preservation, water treatment, healthcare sanitation, textile, and other industries. 1-4 During past decades, a wide variety of bioactive organic chemicals have been developed for disinfection, sterilization, and preservation purposes, including quaternary ammonium compounds, alcoholic and phenolic compounds, aldehydes, halogen-containing compounds, quinoline and isoquinoline derivatives, heterocyclic compounds, and peroxygens.^{5,6} Biocides have also been applied in oil reservoirs for many decades, particularly in water flooding operations during secondary oil recovery.⁷ Likewise, biocides are among the most common chemical additives used for hydraulic fracturing ("fracking" or "fraccing"), a process in which a water-based fluid is used to help induce cracks in oiland/or natural gas-containing unconventional formations such as shale rock. At total concentrations of up to >500 mg/L⁸ and total fluid volumes surpassing 10 million L per horizontal well,9 total amounts of biocide(s) used per hydraulic fracturing event can exceed 1,000 gallons.10

Bacterial control is necessary in hydraulic fracturing operations to prevent excessive biofilm formation downhole that may lead to clogging, consequently inhibiting gas extraction.¹¹ Biocides inhibit growth of sulfate-reducing bacteria (SRB), 12-14 which anaerobically generate sulfide during the organisms' respiration process. Sulfide species created in the subsurface may pose a risk regarding occupational safety and health when the fluid returns along with produced H2S gas. Furthermore, SRB and acid-producing bacteria (APB) may induce corrosion of the production casing/tubing underground, potentially leading to casing failure and environmental contamination by petroleum products.^{7,12,14–18}

Hydraulic fracturing operations provide bacterial species with many habitats favorable to their (unwanted) growth and

Received: July 30, 2014 November 20, 2014 Revised: Accepted: November 26, 2014 Published: November 26, 2014

16

Department of Chemistry, Colorado State University, 1872 Campus Delivery, Fort Collins, Colorado 80523, United States

[‡]Department of Civil and Environmental Engineering, Colorado State University, 1320 Campus Delivery, Fort Collins, Colorado 80523, United States

[§]Department of Chemical and Biological Engineering & Center for Biofilm Engineering, Montana State University, 366 EPS Building, Bozeman, Montana 59717, United States

Department of Soil and Crop Sciences, Colorado State University, 1170 Campus Delivery, Fort Collins, Colorado 80523, United States

proliferation. The major sources of bacterial contamination are (1) drilling mud, (2) water, (3) proppants, and (4) storage tanks. Prolonged storage of water prior to use, typically in lined or unlined earthen pits, 12,14 can lead to mass proliferation of microorganisms. Likewise, bacteria can thrive in stored produced water that was recycled for use in future fracturing operations. 19,20 The increased temperatures fracturing fluids are exposed to underground may also favor microbial growth, 21 and therefore many bacterial species (including anaerobic species that are native to shale formations) 16,22-25 may proliferate underground during hydraulic fracturing. A diverse array of bacteria including those within the taxa γ -proteobacteria, α -proteobacteria, δ-proteobacteria, Clostridia, Synergistetes, Thermotogae, Spirochetes, Bacteroidetes, and Archaea have all been found in untreated flowback water samples.²⁵ Sulfate reduction by piezophilic bacteria coupled to oxidation of methane or organic matter is stimulated at high pressure; 26-30 pressure increases naturally underground but is also artificially increased during the process of hydraulic fracturing, matching values relevant to the referenced studies. Similar pressure-stimulation effects have also been observed in iron-reducing bacteria. 26,31,32 In fact, reduction in viability of bacteria due to high pressure does not occur until 100-150 MPa (~15,000-22,000 psi),²⁶ which exceed most formation pressures.

Biocides are often (but not always) used in hydraulic fracturing fluid formulations. While their application is often mandatory in aboveground oil-water separation units, water storage tanks, and pipelines used to transport these fluids, it is believed that the extremely high temperatures inherent to some shale formations may naturally impede microbial growth. The temperature of shale plays in the continental U.S. varies greatly, even within a single formation. The gas-bearing shales of the Marcellus formation, which exist from 1,200 to 2,600 m underground, ^{33,34} typically fall within a temperature range of 40–100 °C, but can reach temperatures of 100-125 °C in the southwestern region.²⁰ In deeper shales such as the Haynesville in Texas/ Louisiana (3,200 to 4,100 m underground³⁴), the downhole temperature can reach almost 200 °C.^{34,35} However, a few studies have suggested that some bacteria are very persistent and may not be completely killed by the extreme underground conditions^{23,35,36} or that higher pressures may prevent bacterial death at higher temperatures.^{29,37} Furthermore, injection of colder fracturing fluids may lead to considerable cooling of the casing and target formation. Thus, biocides are sometimes added to fracturing fluids even in formations with temperatures exceeding 122 °C, the highest recorded temperature at which aerobic bacteria reproduction has been observed.³⁸

To achieve downhole bacterial control, a variety of biocides are currently being added to hydraulic fracturing fluids (Table 1, Supporting Information (SI) Figure S1). Selection of the biocide(s) used is highly dependent on the geology and biogeochemistry of the respective shale formation, and packages are individually tailored depending on their efficacy in bacterial control, compatibility with the respective environment, and cost efficiency. Compatibility with the respective environment, and cost efficiency. Compatibility with the respective environment, and cost efficiency. Furthermore, the doses needed for sufficient treatment. Furthermore, the reactivity toward other fluid additives must be considered, as many biocides are inherently reactive molecules and side reactions are undesirable.

Unlike water flooding, in which there are continuous inflows and, thus, there is a need for long-term suppression of microbial activity, the goal of biocide application in hydraulic fracturing is

Table 1. Chemical Identification Information on Hydraulic Fracturing ${\operatorname{Biocides}}^a$

Trade Name & CAS No.	Chemical Structure	Chemical Formula	МОА	Freq. of Use
Glutaraldehyde 111-30-8	O O II II HC CH	C ₅ H ₈ O ₂	Е	27%
Dibromo- nitrilopropionamide 10222-01-2	N Br NH ₂	C ₃ H ₂ Br ₂ N ₂ O	Е	24%
Tetrakis hydroxymethyl phosphonium sulfate 55566-30-8	HO OH O O	[(HOCH ₂) ₄ P] ₂ SO ₄	E	9%
Didecyl dimethyl ammonium chloride 7173-51-5	H ₃ C + CI -	C ₂₂ H ₄₈ NCI	L	8%
Chlorine dioxide 10049-04-4	0 C >0	CIO ₂	0	8%
Tributyl tetradecyl phosphonium chloride 81741-28-8	P CI-	C ₂₆ H ₅₆ PCI	L	4%
Alkyl dimethyl benzyl ammonium chloride 68424-85-1	CI CH ₃	C ₁₉ H ₃₄ NCI	L	3%
Methylisothiazolinone 2682-20-4	H ₃ C N - S	C₄H₅NOS	Е	3%
Chloro- methylisothiazolinone 26172-55-4	H₃C N−S O CI	C₄H₄NOSCI	Е	3%
Sodium Hypochlorite 7681-52-9	Na ⁺ CI—O ⁻	NaClO	0	3%
Dazomet 533-74-4	s N	C ₅ H ₁₀ N ₂ S ₂	Е	2%
Dimethyloxazolidine 51200-87-4	N, H	C ₅ H ₁₁ NO	Е	2%
Trimethyloxazolidine 75673-43-7	∑ _N ,CH₃	C ₆ H ₁₄ NO	Е	2%
N-Bromosuccinimide 128-08-5	o N O	C ₄ H ₄ BrNO ₂	E	1%
Bronopol 52-51-7	HO OHOO -	C ₃ H ₆ BrNO ₄	Е	<1%
Peracetic acid 79-21-0) О ОН	C₂H₄O₃	0	<1%

^aMode of action (MOA) listed is electrophilic (E), lytic (L), or oxidizing (O). Frequency of use is normalized to only those wells reported on FracFocus⁴⁸ in which biocides were used.

to reduce the deleterious microbial populations up-front to the lowest levels possible, in order to mitigate the risk of their colonization of the reservoir and well system long after the fracturing operation has been completed. This is due to the fact that reservoir souring in the fractures downhole cannot be cured after hydraulic fracturing is complete. While sterilization is unlikely, targeting deleterious bacteria with biocides that have been efficacy-tested for performance against those specific species can lead to long-term protection for months or longer, mitigating the risk for souring and microbially induced corrosion (MIC). Thus, it is critical to understand microbial dynamics in this context, i.e., types and concentrations of

microorganisms, carbon sources, nitrogen sources, and electron acceptors present, as well as growth-limiting factors. Furthermore, potential growth rates of microorganisms in fracturing fluids under subsurface conditions need to be considered, which are not yet fully understood. 23-25 To determine a suitable site-specific biocide (combination), 6-log reductions or greater in SRB, APB, or other deleterious bacterial populations are typically aimed for in preliminary laboratory testing according to NACE Standard TM0194.47 When choosing suitable biocide(s), typical specific parameters of the fracturing operation that can affect biocide performance are taken into account. Some of these parameters include compatibility with the other fracturing chemicals being used, flow rates of the fracture fluids, and the chemistry of the water used. Reservoir conditions such as temperature and formation geology may also be considered. However, selection of biocides is often done based on historical precedence and currently is not optimized on a well-by-well basis.

The following sections of this review will focus on the most common hydraulic fracturing biocides by frequency of application (Table 1) according to the national hydraulic fracturing chemical registry FracFocus.⁴⁸ FracFocus is managed by the Ground Water Protection Council and the Interstate Oil and Gas Compact Commission and is currently used as the official state chemical disclosure system in 10 U.S. states (CO, OK, LA, TX, ND, MT, MS, UT, OH, PA). While the registry is not inclusive of wells outside the listed states and is dependent on company-volunteered information in states that do not mandate reporting of injected chemicals, it is the largest registry of its type and allows an approximate estimation of national chemical use trends. Additionally, not all compounds listed as biocides on FracFocus are active ingredients with biocidal activity (i.e., ethylene glycol); regardless, these compounds are registered as biocides because they are components in commercial biocide product mixtures. To enable assessment of the potential environmental and health impacts of the biocides used in hydraulic fracturing fluids, we review their relevant pathways of environmental contamination, environmental mobility, stability and chemical behavior in a variety of natural environments, and toxicity. Alternatives to traditional chemical biocides will be highlighted as methods of achieving the necessary bacterial control in hydraulic fracturing operations while lowering environmental risks. Finally, areas in need of research are revealed as current knowledge gaps impede full understanding of environmental fate and transport of biocides.

HYDRAULIC FRACTURING BIOCIDES AND THEIR MODES OF ACTION

Biocides function by various modes of action to control bacteria—they are generally divided into oxidizing and non-oxidizing compounds. Oxidizing biocides such as bromine-(e.g., N-bromosuccinimide, NBS) and chlorine-based (e.g., chlorine dioxide and sodium hypochlorite) species rely on the action of released free radical species that attack cellular components. Peroxides are often used in pretreatment of obtained natural water sources due to the fact that they are extremely nonspecific and work efficiently on a wide range of bacteria. However, oxidizing biocides pose the risk of equipment corrosion as well as unwanted reactions with other hydraulic fracturing chemicals. Furthermore, their reaction may produce halogenated hydrocarbons and other unwanted disinfection byproducts (DBPs). Sp. Oxidizing biocides are too

short-lived to control microbial growth for long periods of time during fluid storage and to control growth of bacteria that may already be present in the well bore area from the drilling operation, making this class of biocide poorly suited for injection. For these reasons, oxidizing biocides are more commonly used for on-site treatment of stored fresh and wastewater, and less often in hydraulic fracturing fluids (Table 1, SI Figure S2) (though their use is increasing in some areas based on recent trial-and-error successes). However, oxidizing biocides are well-reviewed in the existing literature and therefore will not be emphasized in this critical review.

The most common biocides used in hydraulic fracturing fluids used are nonoxidizing organic chemicals, which will therefore be focused on. These biocides are split into two groups according to their respective modes of action: electrophilic and lytic (often used in conjuncture in fracturing fluids). Many of these biocides are not exclusive to hydraulic fracturing and are used by many other industries and products (SI Table S1).

Lytic Biocides. Lytic (also known as membrane-active) biocides are amphiphilic surfactants, and their activity is generally based on dissolution into the bacterial cell wall and its subsequent disruption. Specifically, their known mode of action involves binding to anionic functional groups on the membrane surface and subsequent perturbation and dissolution of the lipid bilayer, resulting in loss of osmotic regulation capacity and eventual lysis of the cells. SS

The two main lytic biocides used for hydraulic fracturing are the cationic quaternary ammonium/amine compounds ("QACs" or "quats") didecyl dimethylammonium chloride (DDAC or decanaminium; Table 1), and alkyldimethylbenzylammonium chloride (ADBAC, benzalkonium chloride, BAC, BC, or benzenemethanaminium), although they are often not distinguished specifically on disclosure forms. These biocides are characterized by a central quaternary nitrogen atom that carries a permanent positive charge and is bonded to four carbon-containing "R"-substituents.

QACs are injected into 22% of all wells registered on FracFocus and used in nearly every shale formation in the U.S., 48 though not solely as biocides since QACs also serve as cationic surfactants, corrosion inhibitors, and clay stabilizers. 56 The QACs used for biocidal purposes, DDAC (comprising 8% of all biocide use, Table 1), and ADBAC (3% of all use, Table 1) are often combined with electrophilic biocides such as glutaraldehyde to increase their efficiency through synergistic effects, thus lowering the total amounts of biocide needed for appropriate bacterial control. 48,57 The only other commonly used lytic biocide is tributyltetradecylphosphonium chloride (TTPC), whose U.S. EPA registration review is scheduled to be completed in 2017. 58 Therefore, information on this substance is limited.

Electrophilic Biocides. Electrophilic biocides typically have reactive electron-accepting functional groups (i.e., aldehydes) that react with electron-rich chemical groups such as exposed -SH (thiol) and -NH (secondary amine) groups in membrane proteins on bacterial cell walls. ⁵² Glutaraldehyde ("glut") is the most commonly used electrophilic biocide in hydraulic fracturing operations (Table 1). Similar to formaldehyde and other aldehydes, it is a potent cross-linker for amino and nucleic acids (SI Scheme S1), leading to cell wall damage and cytoplasmic coagulation. ⁵⁹ Glutaraldehyde is one of the main biocides being used in off-shore hydraulic fracturing operations internationally. ⁶⁰

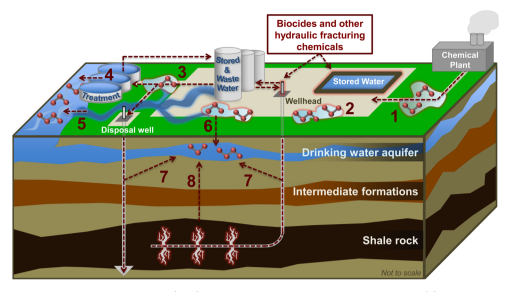


Figure 1. Modes of potential environmental exposure: (1–3) surface spills resulting in contamination of soil; (4) incomplete removal in treatment plants; (5) surface spill runoff into surface water; (6) surface spills leaching into shallow aquifer; (7) contamination of shallow groundwater via borehole leakage, fault lines, and abandoned wells; (8) contamination of shallow groundwater via induced fractures. Artwork is conceptual and not drawn to scale.

Both 2,2-dibromo-2-nitrilopropionamide (DBNPA, 2,2-dibromo-2-cyanoacetamide) and 2-bromo-2-nitropropane-1,3-diol (bronopol) are not considered oxidizing biocides despite small releases of bromine⁶¹ which are assumed to enhance their biocidal activity. ^{49,62} Both rapidly react with sulfur-containing nucleophiles such as glutathione or cysteine, thereby disrupting key cell components and biological functions. ⁶³

Another quaternary phosphonium compound currently registered for use as a biocide in hydraulic fracturing fluid is tetrakis(hydroxymethyl)phosphonium sulfate (THPS). Under alkaline conditions, it deformylates and releases tris(hydroxymethyl)phosphine (THP), which cleaves sulfur—sulfur bonds in the disulfide amino acids of the microbial cell wall (SI Scheme S2). Dazomet (3,5-dimethyl-1,3,5-thiadiazinane-2-thione or "mylon") is often combined with THPS in fracturing fluids. 12

Finally, although rare, use of sulfur-containing biocides is still reported (i.e., California and Texas, Table 1 and SI Figure S1). The most commonly used sulfur-containing biocide treatment is comprised of two chemicals, chloromethylisothiazolinone (CMIT, or MCI, 5-chloro-2-methyl-3(2H)-isothiazolinone) and methylisothiazolinone (MIT, or MI, 2-methyl-3(2H)-isothiazolinone). They react with a wide range of amino acids, inhibiting critical metabolic processes.

MODES OF ACCIDENTAL ENVIRONMENTAL CONTAMINATION

Environmental exposure to biocides after an inadvertent release may occur during (1) transportation of chemicals to well pads (Figure 1, pathway 1); (2) mixing of chemical additives with the bulk of the fracturing fluid (pathway 2); (3) injection of the mixed fluid into the borehole (often occurs simultaneously with in-line mixing; pathway 2); (4) handling, collection, and storage of chemical-containing produced water (pathways 5 and 6); and (5) reuse, treatment, recycling, and/or disposal of the produced water (pathways 3, 4, 5, and 7). The hydraulic fracturing event itself may have a considerable impact on the chemical stability of the organic additives and thus on the composition of the produced water. As the fluids are exposed

to the high pressures, temperatures, salt concentrations, and organic matter contents of the oil- and gas-bearing formations, abiotic transformation and sorption reactions may greatly accelerate the chemicals' depletion (Figure 2). To our knowledge, there is currently only one study that directly addresses this critical effect. For the biocide glutaraldehyde, it was shown that transformation increases with increasing temperature and pH and may be substantial under conditions encountered downhole. Consequently, it can be expected that a portion of the injected biocides will not resurface or may resurface as transformation product(s). Thus, the potential risks associated with biocide exposure may differ substantially before and after hydraulic fracturing.

Few studies have been conducted on the presence of organics in produced water from shales which had undergone hydraulic fracturing. 42,67,68 The compounds detected included polyaromatic hydrocarbons (PAHs), aliphatic hydrocarbons, and long-chain fatty acids. Certain chemicals unique to hydraulic fracturing, including ethoxylated surfactants and the biocide 1,3,5-trimethylhexahydro-1,3,5-triazin-2-thione (a dazomet derivative), were also detected in produced water. Levels of dissolved organics in the produced water as measured directly from active wellheads decreased sharply after the first 20 days; concentration of the biocide dropped from 1.5 mg/L on day 1 to 0.01 mg/L on day 8.

The most commonly reported accidents leading to environmental contamination are surface spills (pathways 1–3 in Figure 1), which can happen on-site (including well blowouts and casing failures)^{10,69,70} or during transportation to or off the site via pipelines, trains, or trucks.^{71–73} On-site spills, which may result in surface water, soil, or shallow groundwater contamination (pathways 5 and 6 in Figure 1),⁵³ are often a result of use of lined pits to temporarily store and evaporate flowback brine in order to reduce the volume of waste.^{10,69} More recently, storage tanks have replaced these pits in an attempt to reduce the potential for unintentional spills.⁷⁴

While comprehensive and complete data on surface spills for most shale plays are still lacking or are otherwise difficult to access, the Colorado Oil and Gas Conservation Commission

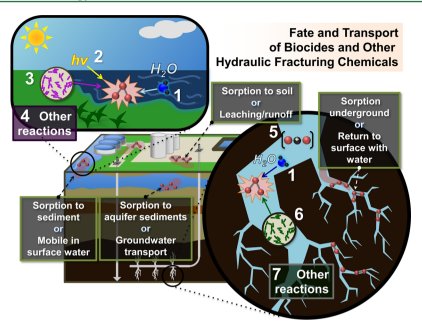


Figure 2. Fate and transport of hydraulic fracturing chemicals (including biocides) in different environments. Possible degradation pathways are numbered as follows: (1) hydrolysis; (2) direct or indirect photolysis; (3) aerobic biodegradation in the water or soil; (4) other chemical reaction with oxygen present; (5) complexation underground with dissolved inorganic species; (6) anaerobic biodegradation; (7) other chemical reaction (e.g., nucleophilic substitution or polymerization) under anoxic conditions, high pressure, and elevated temperature. Artwork is conceptual and not drawn to scale.

(COGCC) offers a robust database that includes any spill or release of exploration and production waste within the State of Colorado of one barrel or more outside and of five barrels or more inside of berms or other secondary containment.⁷⁵ In 2013, there were 591 reported spills, which released a total of 14,067 barrels (i.e., ~2,200,000 L), or 0.004% of all produced water. This relates to a total of 50,067 active oil and gas wells in Colorado, including 4,025 new wells that had been drilled in 2013.

To minimize wastewater volumes and thus potential environmental impacts, wastewater can be reused in subsequent hydraulic fracturing events, which is the dominant management practice in the Marcellus shale region. Of the 12,604 instances of unconventional waste disposal (which includes all fluid and solid waste generated from unconventional resource extraction) reported in Pennsylvania from January to June in 2013, 33.4% indicated treatment by a centralized plant and then recycling in another hydraulic fracturing operation, 30.3% indicated direct reuse of the wastewater in other drilling operations, 26.0% indicated disposal via injection in disposal wells, 9.5% indicated disposal via landfills, and only 0.3% indicated treatment for eventual discharge into surface waters. It is important to note that wastewater treatment strategies vary between states and these statistics may not necessarily be representative of those

While injection into deep underground reservoirs is currently the most common method of unconventional waste disposal, ^{78–80} little is known about the long-term impact and risk of this technique. ^{78,81} However, depending on exposure times of the organic chemical additives to the increased pressures and temperatures of these deep formations, (further) chemical transformation is conceivable. ⁶⁵ Environmental exposure to the injected solution may occur if the cement or casing is faulty (pathway 7, Figure 1); however, this has yet to be documented.

In the minority of instances where wastewater is treated for discharge, publicly owned treatment works (POTWs), municipal wastewater treatment plants (WWTPs), or commercially

operated industrial wastewater treatment plants are used to treat hydraulic fracturing wastewater. 77,82 However, most of the municipal treatment plants are designed for common compounds (nutrients and organic matter) and were not intended to treat the multitude and amounts of chemical species that resurface with flowback brine. Furthermore, assuming the presence of biocides in wastewater at sublethal concentrations, adaptation of surviving microorganisms to biocides becomes possible—a process that is well-documented for ADBAC^{83,84} and other biocides. 45,85 This biocidal resistance is sometimes accompanied by low-level increases in antibiotic resistance, 86 which is already an existing problem in WWTPs. 87 Fortunately, dilution below the minimum inhibitory concentration also enables the majority of these chemicals to undergo biodegradation in WWTPs and in natural waters, which will be discussed in more detail later. In hypothetical cases where biocides persist through WWTP treatment, however, contamination of water and soil may occur (pathway 4 in Figure 1).

Although WWTP effluent is typically released into natural streams and waterways, there are rare documented cases of its reuse for other purposes such as to irrigate crops ^{73,88} or to spread on roads to reduce dust and/or deice. ^{53,77,89,90} Finally, wastewater from hydraulic fracturing operations off the coast of southern California is sometimes discharged directly into the ocean; as of recently, the EPA is requiring full disclosure of all chemicals meant for direct discharge into the Pacific Ocean. ⁹¹

In contrast to accidents occurring at the surface, underground contamination phenomena may go unreported and undetected for years before any negative consequences are observed. As such, though not considered a dominant exposure pathway in hydraulic fracturing operations, underground exposure pathways are the focus of much peer-reviewed literature. The majority of research available suggests that natural upward migration of chemicals such as biocides from hydraulically induced fractures to aquifers (Figure 1, pathway 8) is unlikely. 92,93 Usually, hydraulic fracturing is performed >1,000 m

Table 2. Physicochemical Properties, Reaction Half-Lives, and Biodegradability of Biocides Used in Hydraulic Fracturing Operations^a

						hydrolysis half-life (25 °C)			
biocide	log K _{ow}	% water sol. at 25 $^{\circ}C$ log $K_{\rm ow}$ $$({\rm mg/L})$$	$\log K_{ m H}^{\ b}$	$\log K_{ m oc}$	S Hq	2 Hq	6 Hd	photolysis half-life (12 h day)	readily biodegradable (half-life < 30 days)
ADBAC	no data	no data >350,000 ¹⁰³	-10.87	5.810^{103}	stable 133	stable ¹³³	stable 133	stable (aq) ¹³³	no ¹³³
bronopol	-0.64	$200,000^{219}$	-10.88^{218}	-0.033	1.5 (pH 6, 20 °C) to 2 years ¹³	1.5 (pH 6, 20 °C) to 2 years ¹³⁷ 1.5 (pH 6, 20 °C) to 2 years ¹³⁷ 1.5 (pH 6, 20 °C) to 2 years ¹³⁷ 2 days (aq. pH 4) ¹³⁷	7 1.5 (pH 6, 20 $^{\circ}$ C) to 2 years 137	⁷ 2 days (aq, pH 4) ¹³⁷	no data
chlorine dioxide -3.22^{221}	s -3.22 ²²¹	3.01×10^{03220}	-1.40^{220}	no data	<25 min ²²¹	<25 min ²²¹	<25 min ²²¹	no data	no data
CMIT	-0.34		-7.45	0.711	stable	stable	22 days ²²³	no data	aq (aer) ²²³
dazomet	0.63^{218}	$3,000^{218}$	-9.30^{218}	1.000^{218}	23.8 h	180 h	45.6 h ¹⁴⁷	1.6 h (aq, pH 7) ¹⁴⁷	no data
DBNPA	0.82^{218}	15,000 ²¹⁸	-7.72^{218}	1.767	67 days	63 h	73 min ⁶²	14.8 h (aq, pH 5); 6.9 h (aq, pH 7) ⁶²	aq (aer, anaer) ⁶²
DDAC	$\approx 0^{134}$	700 ²²⁵	-9.16	5.64-6.20 ¹⁰⁴ 368 days	4 368 days	184.5 days	506 days ¹³²		no ¹³²
DMO	0.73^{227}	miscible ²²⁶	-5.52	0.747	<5 min ²²⁷	<5 min ²²⁷	<5 min ²²⁷	no data	no data
glutaraldehyde -0.33 ¹⁰²	-0.33^{102}	miscible ²²⁸	-4.48^{102}	$2.07 - 2.70^{102}$ 628 days	² 628 days	394 days	63.8 days ¹⁰²	196 days (aq, pH 5) ¹⁰² aq (aer, anaer) ¹⁰²	aq (aer, anaer) ¹⁰²
hypochlorite	-3.42^{221}	miscible ²²⁹	nonvolatile ²²¹	no data	<25 min ²²¹	<25 min ²²¹	<25 min ²²¹	no data	no data
MIT	-0.83	536,700	-7.30	0.440	stable	stable	22 days ²²³	no data	aq (aer) ²²³
N-bromo	-1.19	$14,700^{218}$	-6.41	0.259	unstable, no data	unstable, no data	unstable, no data	no data	no data
peracetic acid	-1.07	miscible ²³¹	-5.67^{218}	0.104		8.295 h ²³²		8.3 days ²³³	aq (aer, anaer) ²³³
THPS	-9.77	400,000 ²¹⁸	-12.64	-5.302	131 days	72 days	7 days ¹⁴²	<2 h (aq, pH 7) ¹⁶⁹	aq (aer, anaer), soil (aer) ¹⁶⁹
TMO	(≈DMO)	(≈DMO) miscible ²²⁶	-5.18	0.841	<5 min ²²⁷	<5 min ²²⁷	<5 min ²²⁷	no data	no data
TTPC	no data	no data miscible ²³⁴	3.75×10^{-8} mmHg, 39 °C ¹⁴⁵ 7.66 ^c	145 7.66°	unknown	unknown	unknown	no data	no data

 a Values in italics are calculated using EPI Suite v4.11. 101 Biodegradation reported in aqueous phase (aq) or soil under aerobic (aer) or anaerobic (anaer) conditions. b log $K_{H\nu}$ Henry's law coefficient. c Estimation uses MCI method (otherwise K_{OW} is used in K_{OC} predictions).

below a shallow aquifer used for drinking or irrigation water extraction. Even though some fractures have been documented to grow upward by as much as 450 m, their tops were still almost 1,500 m below the aquifer.⁹⁴

Underground contamination via borehole leakage, preexisting vertical cracks, fault lines, or uncharted boreholes (Figure 1, pathway 7) remains a possibility. Several studies using isotopic analysis have traced methane found in groundwater back to natural gas wells nearby. Darrah and co-workers (2014)⁹³ demonstrated that fugitive gas contamination was due to (1) release of intermediate-depth gas along the well annulus, likely due to cement failure; (2) release of target formation gas implicating improper, faulty, or failing production casings; and (3) underground well failure. While upward flow of leaking hydraulic fracturing fluids would be substantially slower than that of buoyant natural gas,⁵³ production well failure in the proximity or above an aquifer is a more likely potential pathway for groundwater contamination by fracturing fluid components. However, due to the lack of solid (baseline) data, further research is needed to address this question.

In conclusion, while a realistic potential for groundwater contamination via natural upward migration of deep-injected biocides may exist if the fractured formation is very shallow, it appears unlikely. However, several other scenarios exist which may result in unintentional environmental contamination with biocides, including surface spills and well integrity issues associated with casing or cement failure. 53,64,76,93,98–100

MOBILITY

Table 2 summarizes the key physicochemical properties that can be used to predict the mobility of the hydraulic fracturing biocides. The property estimation software EPI Suite v4.11¹⁰¹ was chosen after statistical comparison of the performance of different estimation methods (SI Tables S2 and S3) to close the large existing experimental data gaps regarding their water solubilities, Henry's law constants ($K_{\rm H}$), soil organic carbonwater partition constants ($K_{\rm OW}$).

In general, it can be seen that the organic-based hydraulic fracturing biocides considered here possess low $K_{\rm H}$ values and are thus not very volatile. Consequently, partitioning into the air phase is not considered a major pathway. However, biocides tend to be water-soluble or miscible, enabling their transport via surface water or groundwater depending on their affinity for sorption to natural solid phases (Table 2, Figure 2).

With the exception of glutaraldehyde ($\log K_{\rm OC} = 2.07 - 2.70$), 102 the $K_{\rm OC}$ values of the uncharged biocides lie predominantly below 100 (i.e., $\log K_{\rm OC} < 2$), indicating comparably low retardation during aqueous transport through the soil, especially soils which are low in organic matter content. Glutaraldehyde's mobility is reported to be moderate in soils and high in sediments, 102 and due to its rapid aerobic degradation in soil and water (see below), its potential for soil contamination may only be relevant under anaerobic conditions.

In contrast, ADBAC, DDAC, and TTPC possess substantially higher log $K_{\rm OC}$ values between 5.6 and 7.7, 101,103,104 which is due in part to their large hydrophobic moieties. In addition to hydrophobic interactions with soil organic matter, these cationic biocides can adsorb to permanent negatively charged clay surfaces, an effect which enables the use of these compounds as shale inhibitors/clay stabilizers (only quaternary ammonium/amine polyelectrolytes called "polyquats" with

multiple charges are used for this purpose). 56 Despite being sorbed, these biocides potentially remain bioactive. 106 Several studies reported strong sorption of cationic organic amines to the surfaces and interlayers of clay beyond its cation exchange capacity, causing extensive clay aggregation. $^{107-109}$

Experiments simulating river water contamination revealed that the quaternary ammonium compound paraquat (N,N'-dimethyl-4,4'-bipyridinium dichloride) becomes almost completely sorbed onto sediments $(\geq 97\%)$. Similar findings were reported from leaching studies with CTAB (cetyltrimethylammonium bromide), which showed strong sorption to silica and soil. Moreover, it was observed that adsorption of CTAB to silica surfaces occurs within minutes, indicating that QACs will quickly sorb to soils and sediments upon accidental release. This may result in the accumulation of chemical in soil or sediments affected by a spill.

Studies investigating QACs in municipal sludge observed a substantial extent of sorption. Since about 50% of the municipal biosolids in the U.S. are land-applied, an accumulation of QACs in agricultural soils over time may result, potentially leading to plant uptake as is seen with other pollutants such as organophosphates or to contamination of water resources through leaching and runoff.

Sorption of hydraulic fracturing biocides (and other organic additives) may also occur downhole in the resource-bearing fractured formations. Though unconventional formations such as shale are typically nonpermeable, the process of hydraulic fracturing induces many microfractures, which drastically increases the number of available sorption sites. Negatively charged sorption sites, to which cationic species such as DDAC and ADBAC are strongly attracted, are abundant in shale rock due to the presence of clays. 118 Furthermore, the total organic carbon content in unconventional shale formations can exceed 10%. 119 Thus, partitioning into organic carbon-dominated phases (e.g., kerogen) even for biocides with a rather low log $K_{\rm oc}$ < 2 may become a major pathway for their removal under downhole conditions. ¹⁰⁵ It has also been shown that elevated temperature increases the rate of chemical sorption of organics to soils, sediments, and clays 118 within time scales relevant for hydraulic fracturing operations. When further considering the high pressures (known to enhance adsorption processes 105,120) under downhole conditions, substantial sorption of hydrophobic biocides underground can be expected.

The presence of high salt concentrations may have a considerable impact on biocide sorption to organic and inorganic surfaces. The presence of salt is partly a result of injection of acid into the well prior to injection of the organic fracturing additives. This acid, usually hydrochloric (HCl), is used in concentrations reaching 20% in the injected fluid⁴⁸ and induces dissolution from the shale, including iron, calcium, and radium. For the uncharged biocides, increasing sorption with increasing salt concentration can be expected due to decreasing solubility as a result of salting out. For cationic biocides, which may sorb via electrostatic interactions in addition to hydrophobic forces, the effect of salt concentration is more complex as the salts increasingly compete for ion exchange sites, thus counteracting the salting-out effect. Several studies observed that the sorption affinities of QACs to both clays and organic matter generally decreased with increasing salt concentration.

The potential for bioaccumulation of the neutral organic biocides is predominantly low as their $K_{\rm OW}$ values consistently lie below 10 (i.e., log $K_{\rm OW}$ < 1; Table 2), for many even below 1

(i.e., log $K_{\rm OW}$ < 0), indicating a preference for the aqueous phase over the organic. However, as cationic surfactants typically accumulate at the interface between two phases and may be attracted to the negatively charged surface of cell membranes, the $K_{\rm OW}$ is not sufficient to predict bioaccumulation of ADBAC, DDAC, and TTCP. On the other hand, their potential to undergo strong ionic bonding may limit their mobility, and thus their potential for uptake by organisms: QACs have been found to be far less bioavailable (though potentially still bioactive) to organisms than other pollutants having similar $K_{\rm OW}$ values, 128,129 such as bisphenol A and organophosphates. 117

DEGRADATION

Lytic biocides have been reported to be relatively resistant to degradation in a wide range of environments. They are stable in the presence of strong oxidants, ¹³¹ under acidic and basic conditions, ¹³¹ and when exposed to heat, ⁵⁴ and are not readily degraded by hydrolysis, photolysis, or bacteria. ^{132–134} On the other hand, electrophilic biocides are reactive chemicals and are therefore relatively short-lived in natural environments where reduced functional groups of proteins and organic matter are prevalent. None of the chemicals react with themselves with the exception of glutaraldehyde (SI Scheme S3): unique among biocides, glutaraldehyde undergoes autopolymerization via aldol condensation, especially at high pH values. ¹³⁵ The resulting α,β -unsaturated polymer is not considered toxic, nor are the products of most known reactions between these biocides. Bronopol is an exception as it reacts with oxygen and thiols to produce superoxide, a reactive oxygen species. ⁶¹

Hydrolysis. Hydrolysis is a major degradation pathway common to electrophilic biocides. Characterized by the addition of a water molecule resulting in two smaller fragment molecules, this process is strongly affected by the pH of the surrounding environment. In some cases, the hydrolysis products can be more toxic and/or persistent than their parent compounds, which is the case with the biocides DBNPA, bronopol, THPS, 12,142–144 and dazomet. 12,145–147 DBNPA's major products via hydrolysis (which occurs readily above pH 8.5⁵²) are dibromoacetic acid and dibromoacetonitrile (DBA; SI Scheme S4). 62,136

Bronopol hydrolyzes within 3 h at 60 °C and pH 8, producing formaldehyde, nitrosamines, and other molecules. 137–141 Although the parent compound (bronopol) is rather short-lived in the environment, its degradation products are toxic and more persistent. The hydrolysis degradation products of THPS are THP molecules that subsequently degrade into tris(hydroxymethyl)phosphine oxide (THPO), accompanied by the release of two formaldehyde molecules. 12,142–144 Dazomet's hydrolysis half-life in water of pH 7 is 7.5 days (Table 2). 145 Its major products are methyl isothiocyanate, formaldehyde, monomethylamine, hydrogen sulfide, carbon disulfide (in acid soils), and nitrogen oxides. 12,145–147

Another influence on chemical stability underground is the pH of the surrounding environment. Analysis of produced water revealed a pH range of 5–7, controlled mainly by bicarbonate species intrinsic to the shale formations. ¹²² Although the biocides used in hydraulic fracturing are relatively stable at near-neutral pH, small changes in pH can nonetheless have a large impact on any acid- or base-catalyzed reactions. ^{65,66}

Photodegradation. Some of the hydraulic fracturing biocides contain chromophores, such as MIT, CMIT, dazomet and *N*-bromo, and may thus undergo direct photolysis

reactions when exposed to ultraviolet (UV) light. Experimentally, this has thus far only been tested for DBNPA and bronopol. Via direct photolysis, DBNPA yields dibromoacetic acid as the major degradation product (SI Scheme SS),⁶² Bronopol was shown to produce tris(2-hydroxymethyl-2-nitropropane-1,3-diol).¹³⁷ In addition, all biocides can be degraded by the action of reactive species from indirect photodegradation. However, no experimental data are available for this process.

Biodegradation. Microbial degradation and mineralization of biocides become possible upon dilution or degradation to sublethal levels, and/or due to proliferation of biocide-adapted bacteria. Minimum inhibitory concentrations not only depend on the type and mixture of biocide(s) but also on the type of organism and are typically lower for planktonic cells compared to biofilms. For many hydraulic fracturing biocides, typical MICs are on the order of milligrams to hundreds of milligrams per liter. For many hydraulic fracturing biocides,

Below MICs, bacterially induced degradation of biocides can occur in soil or surface water in the event of a spill, underground as a result of ineffective bacterial control, or at WWTPs. 149,150 In soils, it is generally concluded that cationic surfactants such as DDAC, ADBAC, and TTPC exhibit such strong sorption to soil that their bioavailability and thus biodegradation rates are substantially reduced. 132,133,151,152 One strain of Pseudomonas fluorescens TN4, which had been isolated from activated sludge from a WWTP, was shown to be not only highly resistant to DDAC, ADBAC, and other QACs but also capable of aerobically degrading them at concentrations of 50 mg/L DDAC (MIC 250 mg/L) and 100 mg/L for ADBAC and all other QACs (MICs > 1,000 mg/L). However, the study did not enable conclusions on whether complete mineralization occurred. In general, research pertaining to degradation of QACs in WWTPs is contradictory; most reports indicate that removal from wastewater is primarily based on sorption effects (not biodegradation) and that degradation of QACs by bacteria found in activated sludge is extremely limited.15

Aerobic bacteria rapidly mineralize glutaraldehyde with glutaric acid as an intermediate. In both wastewater and seawater, aerobic biodegradation has been reported at concentrations of up to 50 mg/L. Under anaerobic conditions over a time period of 123 days, it was reported that glutaraldehyde biodegradation led to the accumulation of 1,5-pentanediol and 3-formyl-6-hydroxy-2-cyclohexene-1-propanal, but no mineralization to $\rm CO_2$ was observed. 102

For most of the other biocides, information on their biodegradation behavior is sparse and often only available through their reregistration eligibility decision reports without citation of peer-reviewed studies. DBNPA was reported to undergo aerobic and anaerobic biodegradation on the order of hours, yielding six different products: oxalic acid, 2-cyanoacetamide, bromoacetamide, dibromoacetic acid, bromoacetic acid, and dibromoacetonitrile. ⁶² No information on complete mineralization was given. Biodegradation studies also showed that THPS is completely converted into CO_2 by bacteria. ¹⁴²

Downhole Conditions. As stated previously, very little is known about the reactivity of chemicals under the high temperatures and pressures of downhole conditions. Previous research on produced water revealed that it did not contain all of the same organic compounds which had initially been injected into the well, suggesting that sorption and/or degradation had occurred underground.⁴² As temperatures

Table 3. Reported Toxicological Data of Biocides Used in Hydraulic Fracturing Operations^a

	mamn	nalian (rat)	
biocide	LD ₅₀ (oral) (mg/kg)	LC ₅₀ (inhaled) (4 h, mg/L)	chronic toxicity
ADBAC	305	$0.054 - 0.51^{133}$	no evidence found ¹³³
bronopol	325	0.588 ¹³⁷	reproductive toxicity; degradation products formaldehyde is known human carcinogen (IARC 1) and nitrosamines are likely human carcinogens (IARC $2A$) ¹³⁸
chlorine dioxide	316	0.290^{221}	developmental toxicity in lab animals ²²¹
CMIT	105	$0.330^{222,223}$	assumed to be same or similar to methylisothiazolinone (MIT) ²²⁴
dazomet	519	8.4 ¹⁴⁵	developmental toxicity in lab animals (skeletal variations) ¹⁴⁷
DBNPA	207	0.320^{172}	degradation product DBA is possible human carcinogen (IARC 2B) and developmentally toxic in lab animals 169,170
DDAC	238	0.07^{132}	developmental toxicity (skeletal abnormalities); no evidence of carcinogenicity 132
DMO	1,173	1.10^{227}	degradation product formaldehyde is known human carcinogen (IARC 1) ¹³⁸
glutaraldehyde	460 ^b	>4.16 ^{b,162}	no evidence found ¹⁶²
hypochlorite	5,800	no data ²²¹	mutagenic in vitro; may react to form trihalomethanes, possible human carcinogens (IARC 2B) ²²¹
MIT	105	0.330^{223}	neurotoxic; no evidence of carcinogenicity ²²⁴
N-bromo	1,170	no data ²³⁰	no evidence found ¹⁴⁷
peracetic acid	1,540	0.450^{232}	no evidence found ²³¹
THPS	290	0.591^{142}	mutagenic in vitro; degradation product formaldehyde is known human carcinogen (IARC 1) ¹⁴²
TMO	1,173	1.10^{227}	degradation product formaldehyde is known human carcinogen (IARC 1) ¹³⁸
TTPC	1,002	< 0.9168	developmental toxicity in lab animals 168

[&]quot;Lowest reported value is displayed. Active ingredient (AI) concentration is 100% unless otherwise stated. ^bValues normalized to 100% AI from reported values which use 50% AI.

and pressures increase underground, chemical equilibrium will shift in accordance with Le Chatelier's principle, and reaction equilibrium will shift to favor endothermic products (negative $\Delta_{\rm r}H)$ and products with smaller volume (fewer molecules and liquid/solid phase) as compared to standard state conditions (25 °C at atmospheric pressure). The high downhole pressures and temperatures may not only lead to unexpected chemical reactions or degradation but may also alter the potential for biodegradation of organics (including biocides) underground.

The stability of aqueous organic chemicals at elevated temperatures is also highly influenced by the inorganic minerals present. ^{154–156} Therefore, it is important to attempt to understand the subsurface biogeochemistry that influences the fate of biocides ¹⁵⁶ and other organic additives during hydraulic fracturing. Complete fracturing of a production well typically requires 3–5 days, ¹⁵⁷ and during this time period the inorganics in solution may either complex with or catalyze degradation of the organic additives in fracturing fluids. Furthermore, the extreme pressures utilized in hydraulic fracturing may work in conjuncture with elevated temperatures underground to produce chemical reactions unexpected under normal surface conditions. ^{154,158–160} Although research is currently ongoing in this area, ^{65,66} the fate of biocides under these unique conditions is still unclear.

TOXICITY

Most of the biocides used in fracturing fluids are severe eye and skin irritants but have relatively low acute toxicity to mammals. Apart from a few exceptions, their oral median lethal dose (LD $_{50}$) values for rats range between 200 and >1,000 mg/kg (Table 3). However, the same biocides tend to be acutely toxic to aquatic life at low concentrations, especially to Mollusca such as oysters, which are highly sensitive to pollutants. 161 For example, glutaraldehyde displays only low to moderate toxicity toward aqueous organisms with the exception of oysters (LC $_{50}=0.78$ mg/L in the Eastern Oyster, Crassostrea virginica). 162 DBNPA is quite toxic to all aquatic life; the lowest

reported LC₅₀ for Daphnia magna (D. magna) is 0.66 mg/L (48 h) and 1.0 mg/L for rainbow trout (96 h). 62,163 Shellfish such as the estuarine oyster are especially sensitive to the presence of DBNPA, showing acute toxicity at levels below 0.070 mg/L.⁶² Additionally, DBNPA has been shown to affect the reproduction of D. magna at the subtoxic levels of 0.053 mg/L, and to negatively affect the growth of juvenile rainbow trout at concentrations as low as 0.019 mg/L¹⁶⁴ and fathead minnow at 1.8 mg/L. 165 DBA (breakdown product of DBNPA) also exhibits toxicity to aquatic life at very low concentrations ¹⁶⁵ (fathead minnow LC₅₀ is 0.55 mg/L within 96 h). ¹⁶⁶ Bronopol is very toxic to marine invertebrates such as oysters observed LC₅₀ is 1.6 mg/L for *D. magna*¹⁶⁷ and 0.77 mg/L for the Eastern Oyster. 137 It can therefore be concluded that substantial spills into surface waters or streams may have noticeable ecotoxicological effects on aquatic species.

Despite not being highly acutely toxic, certain biocides are suspected to possess developmental toxicity, carcinogenicity, mutagenicity, genotoxicity, and/or chronic toxicity (Table 3). Only a few of the hydraulic fracturing biocides have thus far been evaluated by the International Agency for Research on Cancer (IARC) or the U.S. EPA. For the remaining biocides, the evidence that does exist is insufficient to draw any firm conclusions.

Though research is scarce, developmental toxicity (i.e., teratogenicity) has been observed from several of the lytic biocides used in fracturing fluids, such as DDAC and TTPC, as well as the conventional chlorine dioxide (Table 3). Observations of developmental toxicity associated with DDAC have led to a recommended daily intake limit of 0.1 (mg/kg)/day for females of reproductive age. While preliminary EPA studies suggest TTPC exhibits some developmental toxicity, the inclusive final report has yet to be published. Other biocides have shown developmental toxicity associated only with very high concentrations, such as bronopol 137 or THPS, 169 or *in vitro*, such as MIT. 170

Carcinogenicity is rare among the biocides used for fracturing fluids (Table 3) and often is confirmed only *in vitro* at very high concentrations. Typically, any carcinogenicity is due to the

breakdown products of the biocides instead of the parent compound themselves, such as the possible human carcinogen DBA (IARC Group 2B), a breakdown product of DBNPA that has been shown to induce tumors in rat liver cells *in vitro*¹⁷¹ and to have carcinogenic effects in laboratory animals when administered in concentrations over 100 mg/L.¹⁷² Other examples include nitrosamines produced from hydrolysis of bronopol, ^{139–141} many of which are probably carcinogenic to humans (IARC Group 2A).^{148,173} Also, several of the biocides used in hydraulic fracturing operations are known or suspected formaldehyde generators. Formaldehyde is a known human carcinogen (IARC Group 1) if inhaled.^{137,138} However, when dissolved in aqueous media, formaldehyde exists predominantly in its less toxic hydrated form, methanediol (CH₂(OH)₂, methylene glycol), and its tendency to evaporate from aqueous solution is low.^{174,175} Examples of formaldehyde generators include bronopol, ¹³⁷ THPS, ^{12,144,176,177} and dazomet; ^{12,145–147} however, none of these have been found to be carcinogenic in laboratory tests.¹⁷⁸

A few compounds (though not carcinogenic) have been found to mutate DNA *in vitro*. Examples of mutagenic biocides include THPS. Other biocides, though not producing mutations, have been found to exert genotoxic effects *in vitro*, such as ADBAC in plant root tip cells and human lymphocytes at concentrations of 1.0 mg/L. However, *in vitro* effects do not necessarily imply human effects. Genotoxic effects from ADBAC exposure have also been observed in human respiratory cells (study used 1,000 mg/L¹⁸³) and fish cells at 0.12 mg/L. This study also revealed that DDAC (or its bromide salt, respectively) exerted genotoxicity toward mammalian cells at 0.3 mg/L. Finally, the electrophilic biocide THPS (or its chloride salt, respectively) was reported to be genotoxic to fish cells at concentrations of 0.2 μ g/L. 184

For a holistic risk assessment, the mobility and degradability of hydraulic fracturing biocides have to be included, which may limit their ability to exert toxicity on humans or other organisms. Attention must be paid, however, to transformation products that are known to be more toxic and/or more persistent than the parent biocide, such as DBNPA and its product DBA, further emphasizing the need for more detailed degradation studies. The charged, cationic biocides may be more resistant to breakdown, but their strong sorption to clay and organic matter is known to reduce their toxicity. ¹⁸⁵

■ ALTERNATIVE APPROACHES

Alternative products and technologies with biocidal activity exist or are currently being explored to enable a pathway to more sustainable bacterial control. However, none of them are without any environmental and/or human health impacts. One biocide that has been recognized for its relatively low environmental impact is THPS, which received the Presidential Green Chemistry Award in 1997 due to its low toxicity, low treatment levels, and rapid breakdown when used properly. However, a surface spill may result in formaldehyde generation due to the evaporation-driven conversion from its hydrated form, methanediol, a breakdown product of THPS.

Peracetic acid is a biocide already used in hydraulic fracturing operations, although thus far only in 0.01% of all wells registered on FracFocus (SI Figure S1). It is a stronger oxidizing agent than $\rm H_2O_2$ but has not been found to produce harmful DBPs¹⁸⁷ and is not expected to have adverse health effects including carcinogenesis.

Ozone has been widely used for drinking water and food disinfection. 189-191 It readily reacts with double bonds in

critical cellular components but is known to generate some DBPs such as bromate. Also, health effects such as various respiratory diseases, heart attack, and premature death are linked with ozone and its associated pollutants. Furthermore, ozone's rapid reaction kinetics are disadvantageous for downhole applications, which usually take days to weeks to completion. As with other oxidizing agents including peracetic acid, it may induce metal corrosion and thus steel casing failure.

Chlorine dioxide has a longer lifetime than ozone and is thus already used in hydraulic fracturing fluids (Table 1). Its corrosion potential is comparably low, but it may produce carcinogenic DBPs. ^{195,196} Because total dissolved organic carbon in produced water has been measured as high as 5,500 mg/L, ^{53,67} formation of DBPs resulting from use of chlorine dioxide, chlorine (generated by sodium hypochlorite), and bromine (generated by *N*-bromo) must be considered. ¹⁹⁷ A previous study correlated increased levels of THMs in WWTP effluent to Marcellus shale wastewater, especially brominated species, were observed. ¹⁹⁸ Furthermore, chlorine dioxide gas is explosive when exceeding 10% (v/v) in air and must thus be produced on-site. Historical illnesses outside hydraulic fracturing activity including one fatality resulting from exposure to less than 19 ppm have been reported. ¹⁹⁹

The injection of nitrate or nitrite has been broadly used as an alternative electron acceptor for microbial respiration to avert unwanted sulfate reduction. Especially in offshore water flooding operations, where seawater is continuously injected downhole, nitrate was proven to be very effective in reducing the amount and activity of SRB.

There are other alternatives which circumvent the need for chemical addition altogether, such as use of ultrasonic radiation 206 or UV light. 207 These technologies have proven to be extremely effective for microbial control, but their high energy demands and lack of residual effect currently prevent them from widespread use. Furthermore, electrochemical approaches, such as generation of electrochemically activated solutions, ²⁰⁸ can be used to generate reactive oxygen species such as hydroxyl radicals. However, in the presence of high chloride concentrations, electrochemical techniques can also generate hypochlorous acid (HOCl), which, as stated previously, is known to produce DBPs. Due to their typically much lower energy consumption, electrochemical technologies are widely applied in drinking water and wastewater treatment plants. Although their use for downhole applications is Although their use for downhole applications is limited, they may be viable alternatives for hydraulic fracturingrelated aboveground water operations.

DISCUSSION AND FUTURE WORK

Based on the currently available data regarding usage, mobility, degradation, and toxicity of the biocides used in hydraulic fracturing operations, the following conclusions can be drawn: (1) While the uncharged species can be mostly expected to be in the water phase where they are subject to biotic or abiotic degradation, the cationic quaternary ammonium and phosphonium compounds will strongly sorb to soil or sediments, where their (bio)availability to degrade may be limited and their toxicity attenuated. (2) In surface and shallow subsurface environments, many of the biocides are degradable through abiotic and biotic (especially aerobic) processes, but some can potentially transform into more toxic or persistent compounds, which may accumulate under certain conditions. (3) Although being a critical process, the understanding of the biocides' degradation and sorption under downhole conditions (high pressure, temperature, and salt

and organic matter concentrations) is extremely limited, currently not enabling reliable risk assessment. (4) Efforts in developing alternative biocides with lower human health risks are ongoing. Application of existing alternatives is currently limited by high cost, high energy demands, inapplicability under downhole conditions, or formation of DBPs.

Critical data regarding the fate and transport of hydraulic fracturing biocides are still missing despite the widespread and long-term use in other water-processing industries. This is especially true for biotic and abiotic (e.g., photolytic, hydrolytic) degradation processes, which are key to controlling organic contaminant removal from the environment. Future research addressing biocide degradation rates should thus focus on fundamental laboratory- and field-scale investigations under various environmental conditions as well as their runoff and leaching potential. Furthermore, as noted earlier, due to the unique exposure to deep geological formations either during hydraulic fracturing or after injection into deep disposal wells, their reactivity and sorption behavior downhole needs to be considered.

The biocides' susceptibility to various transformation²¹⁰ and degradation processes in combination with the fact that some degradation products have been reported to be more toxic and/ or persistent than the parent compound highlights the need for appropriate analytical detection methods to more accurately assess their potential environmental impact. Excellent screening methods based on high-resolution mass spectrometry exist for polar organic molecules^{211–213} and breakdown products,^{210,213–215} which could theoretically be modified and applied toward detecting hydraulic fracturing chemicals. Modification (typically in sample preparation) is necessary to accommodate the high salinity and TDS commonly associated with fracturing wastewater or measurement accuracy may suffer due to matrix suppression effects, as was recently found responsible for underestimation of radium levels in produced water from the Marcellus shale.²¹⁶

Furthermore, future environmental fate and transport studies need to address the complexity of hydraulic fracturing fluids. The presence of other organic additives, especially polymers that increase fluid viscosity such as gelling agents or the friction reducer polyacrylamide, ⁷⁶ may have substantial impacts on their transport through soil. Vice versa, the presence of biocides above their minimum inhibitory concentrations may impact natural biodegradation processes of other organic fracturing fluid additives.

As of today, no groundwater contamination by hydraulic fracturing additives in the U.S. has been irrefutably documented in the peer-reviewed literature. 53,76,100 However, due to the fact that hydraulic fracturing fluid spills have been reported previously (e.g., there were 591 documented spills in 2013 in Colorado alone),⁷⁵ this scenario does not appear unlikely. Consequently, it appears mandatory to address this concern at a minimum through (1) adequate baseline monitoring in combination with (2) comprehensive chemical analyses beyond currently analyzed and suggested chemical reporting parameters (i.e., bulk water characteristics such as pH and TDS, and concentration of inorganics including chloride, sodium, calcium, barium, strontium, magnesium, radium, uranium, and iron).²¹⁷ In the case of inadvertent releases, these data will enable the development of appropriate cleanup strategies, allow for ecosystem and human health risk assessment, and thus pave the way for a more sustainable approach to natural resource extraction.

ASSOCIATED CONTENT

S Supporting Information

Figure S1 showing use of biocides within U.S. shale plays, Figure S2 showing frequency of use and mode of action of biocides, Table S1 listing other uses of hydraulic fracturing biocides, Tables S2 and S3 providing data and statistical analyses of physicochemical property estimation, reaction schemes S1–S5 illustrating biocide transformation reactions, and text listing applicable references. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

*E-mail: thomas.borch@colostate.edu. Phone: +1-970-491-6235.

Author Contributions

¹G.A.K. and J.B. contributed equally to this work.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This research was supported by The Borch-Hoppess Fund for Environmental Contaminant Research. This literature review was conducted by using Google Scholar and ISI Web of Science as the primary search engines.

REFERENCES

- (1) Rutala, W. A.; Weber, D. J. Disinfection and sterilization in health care facilities: What clinicians need to know. *Clin. Infect. Dis.* **2004**, 39 (5), 702–709.
- (2) Holah, J. T.; Taylor, J. H.; Dawson, D. J.; Hall, K. E. Biocide use in the food industry and the disinfectant resistance of persistent strains of Listeria monocytogenes and Escherichia coli. *J. Appl. Microbiol.* **2002**, 92 (s1), 111S–120S.
- (3) Windler, L.; Height, M.; Nowack, B. Comparative evaluation of antimicrobials for textile applications. *Environ. Int.* **2013**, *53*, 62–73 DOI: 10.1016/j.envint.2012.12.010.
- (4) Stewart, P. S.; Rayner, J.; Roe, F.; Rees, W. M. Biofilm penetration and disinfection efficacy of alkaline hypochlorite and chlorosulfamates. *J. Appl. Microbiol.* **2001**, *91* (3), 525–532.
- (5) Russell, A. D. Principles and practice of disinfection, preservation and sterilization. *Principles and Practice of Disinfection, Preservation and Sterilization*; 3rd ed.; Blackwell Science: Oxford, U.K., 1983; pp 5–94.
- (6) Aqeel Ashraf, M.; Ullah, S.; Ahmad, I.; Kaleem Qureshi, A.; Balkhair, K. S.; Abdur Rehman, M. Green Biocides, a promising technology: Current and future applications to industry and industrial processes. *J. Sci. Food. Agric.* **2014**, *94* (3), 388–403 DOI: 10.1002/jsfa.6371.
- (7) Gieg, L. M.; Jack, T. R.; Foght, J. M. Biological souring and mitigation in oil reservoirs. *Appl. Microbiol. Biotechnol.* **2011**, 92 (2), 263–282 DOI: 10.1007/s00253-011-3542-6.
- (8) McCurdy, R. High rate hydraulic fracturing additives in non-Marcellus unconventional shales. *Proceedings of the Technical Workshops for the Hydraulic Fracturing Study: Chemical & Analytical Methods*, EPA 600/R-11/066; U.S. Environmental Protection Agency: Washington, DC, USA, 2011; pp 1721
- (9) Nicot, J. P.; Scanlon, B. R. Water use for shale-gas production in Texas, U.S. *Environ. Sci. Technol.* **2012**, 46 (6), 3580–3586 DOI: 10.1021/es204602t.
- (10) Aminto, A.; Olson, M. S. Four-compartment partition model of hazardous components in hydraulic fracturing fluid additives. *J. Nat. Gas Sci. Eng.* **2012**, *7*, 16–21 DOI: 10.1016/j.jngse.2012.03.006.
- (11) Bottero, S.; Picioreanu, C.; Enzien, M.; Van Loosdrecht, M. C. M.; Bruining, H.; Heimovaara, T. Formation damage and impact on gas flow caused by biofilms growing within proppant packing used in

- hydraulic fracturing. SPE International Symposium and Exhibition on Formation Damage Control, Lafayette, LA, USA; Society of Petroleum Engineers: Richardson, TX, USA, 2010.
- (12) Dawson, J.; Wood, M. A new approach to biocide application provides improved efficiency in fracturing fluids. SPE/EAGE European Unconventional Resources Conference and Exhibition, Vienna, Austria; Society of Petroleum Engineers: Richardson, TX, USA, 2012.
- (13) Lester, Y.; Yacob, T.; Morrissey, I.; Linden, K. G. Can we treat hydraulic fracturing flowback with a conventional biological process? The case of guar gum. *Environ. Sci. Technol. Lett.* **2013**, *1* (1), 133–136 DOI: 10.1021/ez4000115.
- (14) Fitcher, J. K.; Johnson, K.; French, K.; Oden, R.Use of microbiocides in Barnett shale gas well fracturing fluids to control bacteria related problems. *CORROSION 2008*, New Orleans, LA, USA; NACE International: Houston, TX, USA, 2008.
- (15) Alhajji, J.; Valliappan, M. Concerns over the selection of biocides for oil fields and power plants: A laboratory corrosion assessment. *15th World Petroleum Congress*; John Wiley & Sons: New York, 1998
- (16) Struchtemeyer, C. G.; Morrison, M. D.; Elshahed, M. S. A critical assessment of the efficacy of biocides used during the hydraulic fracturing process in shale natural gas wells. *Int. Biodeterior. Biodegrad.* **2012**, 71, 15–21 DOI: 10.1016/j.ibiod.2012.01.013.
- (17) Stark, M.; Allingham, R.; Calder, J.; Lennartz-Walker; Wai, K.; Thompson, P.; Zhao, S. Water and Shale Gas Development; Accenture Consulting: Dublin, Ireland, 2012.
- (18) Brondel, D.; Edwards, R.; Hayman, A.; Hill, D.; Mehta, S.; Semerad, T. Corrosion in the oil industry. *Oilfield Rev.* **1994**, *6* (2), 4–18.
- (19) Lysnes, K.; Bødtker, G.; Torsvik, T.; Bjørnestad, E. Ø.; Sunde, E. Microbial response to reinjection of produced water in an oil reservoir. *Appl. Microbiol. Biotechnol.* **2009**, 83 (6), 1143–1157.
- (20) Shaffer, D. L.; Arias Chavez, L. H.; Ben-Sasson, M.; Romero-Vargas Castrillon, S.; Yip, N. Y.; Elimelech, M. Desalination and reuse of high-salinity shale gas produced water: Drivers, technologies, and future directions. *Environ. Sci. Technol.* **2013**, 47 (17), 9569–9583 DOI: 10.1021/es401966e.
- (21) Gardner, L. R.; Stewart, P. S. Action of glutaraldehyde and nitrite against sulfate-reducing bacterial biofilms. *J. Ind. Microbiol. Biotechnol.* **2002**, 29 (6), 354–360 DOI: 10.1038/sj.jim.7000284.
- (22) Strong, L. C.; Gould, T.; Kasinkas, L.; Sadowsky, M.; Aksan, A.; Wackett, L. P. Biodegradation in waters from hydraulic fracturing: Chemistry, microbiology, and engineering. *J. Environ. Eng.* **2013**, *B4013001*, 1–8.
- (23) Struchtemeyer, C. G.; Elshahed, M. S. Bacterial communities associated with hydraulic fracturing fluids in thermogenic natural gas wells in North Central Texas, USA. *FEMS Microbiol. Ecol.* **2012**, *81* (1), 13–25 DOI: 10.1111/j.1574-6941.2011.01196.x.
- (24) Murali Mohan, A.; Hartsock, A.; Bibby, K. J.; Hammack, R. W.; Vidic, R. D.; Gregory, K. B. Microbial community changes in hydraulic fracturing fluids and produced water from shale gas extraction. *Environ. Sci. Technol.* **2013**, *47* (22), 13141–13150 DOI: 10.1021/es402928b.
- (25) Murali Mohan, A.; Hartsock, A.; Hammack, R. W.; Vidic, R. D.; Gregory, K. B. Microbial communities in flowback water impoundments from hydraulic fracturing for recovery of shale gas. *FEMS Microbiol. Ecol.* **2013**, *86* (3), 567–580 DOI: 10.1111/1574-6941.12183.
- (26) Picard, A.; Daniel, I. Pressure as an environmental parameter for microbial life—A review. *Biophys. Chem.* **2013**, *183*, 30–41 DOI: 10.1016/j.bpc.2013.06.019.
- (27) Fang, J.; Zhang, L.; Bazylinski, D. A. Deep-sea piezosphere and piezophiles: Geomicrobiology and biogeochemistry. *Trends Microbiol.* **2010**, *18* (9), 413–422.
- (28) Parkes, R. J.; Cragg, B. A.; Bale, S. J.; Goodman, K.; Fry, J. C. A combined ecological and physiological approach to studying sulphate reduction within deep marine sediment layers. *J. Microbiol. Methods.* **1995**, 23 (2), 235–249.
- (29) Kallmeyer, J.; Boetius, A. Effects of temperature and pressure on sulfate reduction and anaerobic oxidation of methane in hydrothermal sediments of Guaymas Basin. *Appl. Environ. Microbiol.* **2004**, *70* (2), 1231–1233.

- (30) Vossmeyer, A.; Deusner, C.; Kato, C.; Inagaki, F.; Ferdelman, T. G. Substrate-specific pressure-dependence of microbial sulfate reduction in deep-sea cold seep sediments of the Japan trench. *Front. Microbiol.* **2012**, 3 (253), 1–12 DOI: 10.3389/fmicb.2012.00253.
- (31) Picard, A.; Testemale, D.; Hazemann, J.-L.; Daniel, I. The influence of high hydrostatic pressure on bacterial dissimilatory iron reduction. *Geochim. Cosmochim. Acta* **2012**, *88*, 120–129 DOI: 10.1016/j.gca.2012.04.030.
- (32) Wu, W. F.; Wang, F. P.; Li, J. H.; Yang, X. W.; Xiao, X.; Pan, Y. X. Iron reduction and mineralization of deep-sea iron reducing bacterium Shewanella piezotolerans WP3 at elevated hydrostatic pressures. *Geobiology* **2013**, *11* (6), 593–601 DOI: 10.1111/gbi.12061.
- (33) King, H. Marcellus shale—Appalachian Basin natural gas play. *Geology.com*, 2008, http://geology.com/articles/marcellus-shale.shtml.
- (34) U.S. Shale Gas: An Unconventional Resource. Unconventional Challenges, White Paper; Halliburton: Houston, TX, USA, 2008.
- (35) Fichter, J.; Wunch, K.; Moore, R.; Summer, E.; Braman, S.; Holmes, P. How hot is too hot for bacteria? A technical study assessing bacterial establishment in downhole drilling, fracturing and stimulation operations. *CORROSION 2012*; NACE International: Houston, TX, USA2012.
- (36) Moore, S. L.; Cripps, C. M. Bacterial survival in fractured shale-gas wells of the Horn River Basin. *Northern Area Western Conference*, Calgary, Alberta, Canada, Vol. *S1* (4); NACE International: Houston, TX, USA, 2010.
- (37) Zobell, C. E.; Johnson, F. H. The Influence of Hydrostatic Pressure on the Growth and Viability of Terrestrial and Marine Bacteria. *J. Bacteriol.* **1949**, *57* (2), 179–189.
- (38) Takai, K.; Nakamura, K.; Toki, T.; Tsunogai, U.; Miyazaki, M.; Miyazaki, J.; Hirayama, H.; Nakagawa, S.; Nunoura, T.; Horikoshi, K. Cell proliferation at 122°C and isotopically heavy Ch₄ production by a hyperthermophilic methanogen under high-pressure cultivation. *Proc. Natl. Acad. Sci. U. S. A.* **2008**, *105* (31), 10949–10954.
- (39) Modern Shale Gas Development in the United States: A Primer; U.S. Department of Energy (DOE), U.S. Government Printing Office: Washington, DC, USA, 2009.
- (40) Rahm, B. G.; Riha, S. J. Toward strategic management of shale gas development: Regional, collective impacts on water resources. *Environ. Sci. Policy* **2012**, *17*, 12–23 DOI: 10.1016/j.envs-ci.2011.12.004
- (41) Rahim, Z.; Holditch, S. A. Effects of fracture fluid degradation on underground fracture dimensions and production increase. *J. Pet. Sci. Eng.* **2003**, *37* (1), 97–111.
- (42) Carter, K. E.; Hakala, J.; Hammack, R. W. Hydraulic fracturing and organic compounds—Uses, dispsal, and challenges. *SPE Eastern Regional Meeting*, Pittsburgh, PA, USA; Society of Petroleum Engineers: Richardson, TX, USA, 2013.
- (43) Sianawati, E.; Yin, B.; Williams, T.; Mcginley, H.; Nair, S. Microbial control management for oil and gas recovery operation. *SPE Kuwait Oil and Gas Show and Conference*; Society of Petroleum Engineers: Richardson, TX, USA, 2013.
- (44) Xu, D.; Li, Y.; Gu, T. A synergistic D-tyrosine and tetrakis hydroxymethyl phosphonium sulfate biocide combination for the mitigation of an SRB biofilm. *World J. Microbiol. Biotechnol.* **2012**, 28 (10), 3067–3074 DOI: 10.1007/s11274-012-1116-0.
- (45) Williams, T. M. Optimizing and Improving biocide performance in water treatment systems. *CORROSION 2008*; NACE International: Houston, TX, USA, 2008.
- (46) Corrin, E. R. Personal correspondence regarding general information on microbial control during hydraulic fracturing activities, 2014.
- (47) Field Monitoring of Bacterial Growth in Oil and Gas Systems, Standard Test Method Tm0194-2004; National Association of Corrosion Engineers: Houston, TX, USA, 2004; ISBN 1-57590-197-7.
- (48) Frac Focus Chemical Disclosure Registry, 2013, http://fracfocus.
- (49) Wiencek, M.; Chapman, J. S. Water treatment biocides: How do they work and why should you care?. *CORROSION* 99; NACE International: Houston, TX, USA, 1999.

- (50) Carpenter, J. F.; Nalepa, C. J. Bromine-based biocides for effective microbiological control in the oil field. *SPE International Symposium on Oilfield Chemistry*, The Woodlands, TX, USA; Society of Petroleum Engineers: Richardson, TX, USA, 2004.
- (51) Finnegan, M.; Linley, E.; Denyer, S. P.; Mcdonnell, G.; Simons, C.; Maillard, J. Y. Mode of action of hydrogen peroxide and other oxidizing agents: Differences between liquid and gas forms. *J. Antimicrob. Chemother.* **2010**, *65* (10), 2108–2115 DOI: 10.1093/jac/dkq308.
- (52) Bartlett, K.; Kramer, J. Comparative performance of industrial water treatment biocides. *CORROSION 2011*; NACE International: Houston, TX, USA2011.
- (53) Vengosh, A.; Jackson, R. B.; Warner, N.; Darrah, T. H.; Kondash, A. A critical review of the risks to water resources from unconventional shale gas development and hydraulic fracturing in the United States. *Environ. Sci. Technol.* **2014**, *48* (15), 8334–8348 DOI: 10.1021/es405118y.
- (54) Kramer, J. F.; Brien, F. O.; Strba, S. F. A new high performance quaternary phosphonium biocide for microbiological control in oilfield water systems. *CORROSION 2008*; NACE International: Houston, TX, USA, 2008.
- (55) Gilbert, P.; Moore, L. E. Cationic antiseptics: Diversity of action under a common epithet. *J. Appl. Microbiol.* **2005**, 99 (4), 703–715 DOI: 10.1111/j.1365-2672.2005.02664.x.
- (56) Patel, A. D. Design and development of quaternary amine compounds: Shale inhibition with improved environmental profile. SPE International Symposium on Oilfield Chemistry, The Woodlands, TX, USA; SPE International: Richardson, TX, USA, 2009.
- (57) Enzien, M.; Yin, B.; Love, D.; Harless, M.; Corrin, E. Improved microbial control programs for hydraulic fracturing fluids used during unconventional shale-gas exploration and production. *SPE International Symposium on Oilfield Chemistry*, The Woodlands, TX, USA; Society of Petroleum Engineers: Richardson, TX, USA, 2011.
- (58) Tri-N Butyl Tetradecyl Phosphonium Chloride (Ttpc) Amended Final Work Plan Registration Review; Docket No. EPA-HQ-OPP-2011-0952; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2012.
- (59) Maillard, J. Bacterial Target Sites for Biocide Action. J. Appl. Microbiol. 2002, 92 (s1), 16S-27S.
- (60) Mcginley, H. R.; Enzien, M. V.; Jennerman, G.; Harris, J. Studies on the chemical stability of glutaraldehyde in produced water. *SPE International Symposium on Oilfield Chemistry*, The Woodlands, TX, USA; Society of Petroleum Engineers: Richardson, TX, USA, 2011.
- (61) Williams, T. M.; Mcginley, H. R. Deactivation of industrial water treatment biocides. *NACE International Corrosion Conference & Expo*; NACE International: Houston, TX, USA, 2010.
- (62) Reregistration Eligibility Decision (RED) 2,2-Dibromo-3-nitrilopropionamide (DBNPA), EPA 738-R-94-026; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 1994.
- (63) Paulus, W. Directory of Microbicides for the Protection of Materials: A Handbook.; Springer: Dordrecht, The Netherlands, 2005.
- (64) Study of the Potential Impacts of Hydraulic Fracturing on Drinking Water Resources, EPA 601/R-12/011; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2012.
- (65) Kahrilas, G.; Blotevogel, J.; Corrin, E. R.; Borch, T. Fate of hydraulic fracturing chemicals under down-hole conditions. *Abstracts of Papers of the American Chemical Society*, Vol. 246; American Chemical Society: Washington, DC, USA, 2013.
- (66) Blotevogel, J.; Kahrilas, G.; Corrin, E. R.; Landry, C. F.; Borch, T. Degradation of the biocide glutaraldehyde under down-hole conditions. *Abstracts of Papers of the American Chemical Society*, Vol. 246, 86-ENVR; American Chemical Society: Washington, DC, USA, 2014.
- (67) Orem, W.; Tatu, C.; Varonka, M.; Lerch, H.; Bates, A.; Engle, M.; Crosby, L.; Mcintosh, J. Organic substances in produced and formation water from unconventional natural gas extraction in coal and shale. *Int. J. Coal Geol.* **2014**, *126*, 20–31.

- (68) Thurman, E. M.; Ferrer, I.; Blotevogel, J.; Borch, T. Analysis of hydraulic fracturing flowback and produced waters using accurate mass: identification of ethoxylated surfactants. *Anal. Chem.* **2014**, *86* (19), 9653–9661 DOI: 10.1021/ac502163k.
- (69) Soeder, D. J.; Kappel, W. M. Water Resources and Natural Gas Production from the Marcellus Shale, FS 2009-3032; United States Geological Survey: Baltimore, MD, USA, 2009.
- (70) Staaf, E. Risky Business: An Analysis of Marcellus Shale Gas Drilling Violations in Pennsylvania 2008–2011; PennEnvironment Research & Policy Center. [ONLINE] 2012, http://pennenvironmentcenter.org/sites/environment/files/reports/Risky%20Business%20Violations%20Report 0.pdf.
- (71) Larson, J.; Kusnetz, N. North Dakota Spills. ProPublica, 2012, http://www.propublica.org/special/north-dakota-spills.
- (72) Rassenfoss, S. From flowback to fracturing: Water recycling grows in the Marcellus Shale. *J. Pet. Technol.* **2011**, *63*, 48–51.
- (73) Veil, J. A.; Puder, M. G.; Elcock, D.; Redweik R. J., Jr. A White Paper Describing Produced Water from Production of Crude Oil, Natural Gas, and Coal Bed Methane; Argonne National Laboratory, U.S. Department of Energy: Argonne, IL, USA, 2004.
- (74) Farag, A. M.; Harper, D. D. A review of environmental impacts of salts from produced waters on aquatic resources. *Int. J. Coal Geol.* **2013**, *126*, 157–161.
- (75) Colorado Oil and Gas Conservation Commission. *Master Records Database, Spill/Release Information,* Mar. 05, 2014, http://cogcc.state.co.us/.
- (76) Vidic, R. D.; Brantley, S. L.; Vandenbossche, J. M.; Yoxtheimer, D.; Abad, J. D. Impact of shale gas development on regional water quality. *Science* **2013**, *340* (6134), 1235009–1235009 DOI: 10.1126/science.1235009.
- (77) Frac Tracker. 2013, http://www.fractracker.org/.
- (78) Hammer, R.; Vanbriesen, J. In Fracking's Wake: New Rules Are Needed to Protect Our Health and Environment From Contaminated Wastewater, NRDC Document D:12-05-A; Natural Resources Defense Council (NDRC): New York, 2012; pp 1–113.
- (79) Gregory, K. B.; Vidic, R. D.; Dzombak, D. A. Water management challenges associated with the production of shale gas by hydraulic fracturing. *Elements* **2011**, *7* (3), 181–186 DOI: 10.2113/gselements.7.3.181.
- (80) Lutz, B. D.; Lewis, A. N.; Doyle, M. W. Generation, transport, and disposal of wastewater associated with marcellus shale gas development. *Water Resour. Res.* **2013**, 49 (2), 647–656 DOI: 10.1002/wrcr.20096.
- (81) Myers, T. Potential contaminant pathways from hydraulically fractured shale to aquifers. *Ground Water* **2012**, *50* (6), 872–882 DOI: 10.1111/j.1745-6584.2012.00933.x.
- (82) Ferrar, K. J.; Michanowicz, D. R.; Christen, C. L.; Mulcahy, N.; Malone, S. L.; Sharma, R. K. Assessment of effluent contaminants from three facilities discharging marcellus shale wastewater to surface waters in Pennsylvania. *Environ. Sci. Technol.* **2013**, 47 (7), 3472–3481 DOI: 10.1021/es301411q.
- (83) Oh, S.; Tandukar, M.; Pavlostathis, S. G.; Chain, P. S.; Konstantinidis, K. T. Microbial community adaptation to quaternary ammonium biocides as revealed by metagenomics. *Environ. Microbiol.* **2013**, *15* (10), 2850–2864 DOI: 10.1111/1462-2920.12154.
- (84) Bore, E.; Hebraud, M.; Chafsey, I.; Chambon, C.; Skjaeret, C.; Moen, B.; Moretro, T.; Langsrud, O.; Rudi, K.; Langsrud, S. Adapted tolerance to benzalkonium chloride in Escherichia coli K-12 Studied by transcriptome and proteome analyses. *Microbiology* **2007**, *153* (Pt 4), 935–946 DOI: 10.1099/mic.0.29288-0.
- (85) Russell, A. D. Mechanisms of bacterial resistance to biocides. *Int. Biodeterior. Biodegrad.* **1995**, *36* (3), 247–265.
- (86) Russell, A. D. Biocide use and antibiotic resistance: The relevance of laboratory findings to clinical and environmental situations. *Lancet Infect. Dis..* **2003**, 3 (12), 794–803.
- (87) Pruden, A. Balancing water sustainability and public health goals in the face of growing concerns about antibiotic resistance. *Environ. Sci. Technol.* **2014**, 48 (1), 5–14 DOI: 10.1021/es403883p.

- (88) Shariq, L. Uncertainties associated with the reuse of treated hydraulic fracturing wastewater for crop irrigation. *Environ. Sci. Technol.* **2013**, 47 (6), 2435–2436 DOI: 10.1021/es4002983.
- (89) Warner, N. R.; Christie, C. A.; Jackson, R. B.; Vengosh, A. Impacts of shale gas wastewater disposal on water quality in Western Pennsylvania. *Environ. Sci. Technol.* **2013**, 47 (20), 11849–11857 DOI: 10.1021/es402165b.
- (90) Skalak, K. J.; Engle, M. A.; Rowan, E. L.; Jolly, G. D.; Conko, K. M.; Benthem, A. J.; Kraemer, T. F. Surface disposal of produced waters in Western and Southwestern Pennsylvania: Potential for accumulation of alkali-earth elements in sediments. *Int. J. Coal Geol.* **2013**, *126*, 162–170.
- (91) Federal Register: Reissuance of National Pollutant Discharge Elimination System (NPDES) General Permit for Offshore Oil and Gas Exploration, Development and Production Operations Off Southern California, FRL-99055-23-Region 9; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2014.
- (92) Flewelling, S. A.; Sharma, M. Constraints on upward migration of hydraulic fracturing fluid and brine. *Groundwater* **2014**, *52* (1), 9–19 DOI: 10.1111/gwat.12095.
- (93) Darrah, T. H.; Vengosh, A.; Jackson, R. B.; Warner, N. R.; Poreda, R. J. Noble gases identify the mechanisms of fugitive gas contamination in drinking-water wells overlying the Marcellus and Barnett shales. *Proc. Natl. Acad. Sci. U. S. A.* **2014**, *111* (39), 14076—14081 DOI: 10.1073/pnas.1322107111.
- (94) Fisher, K. M.; Warpinski, N. R. Hydraulic-fracture-height growth: Real data. SPE Prod. Oper. 2012, 27 (01), 8–19 http://dx.doi.org/10.2118/145949-PA.
- (95) Jackson, R. B.; Vengosh, A.; Darrah, T. H.; Warner, N. R.; Down, A.; Poreda, R. J.; Osborn, S. G.; Zhao, K.; Karr, J. D. Increased stray gas abundance in a subset of drinking water wells near Marcellus shale gas extraction. *Proc. Natl. Acad. Sci. U. S. A.* **2013**, *110* (28), 11250–11255 DOI: 10.1073/pnas.1221635110.
- (96) Osborn, S. G.; Vengosh, A.; Warner, N. R.; Jackson, R. B. Methane contamination of drinking water accompanying gas-well drilling and hydraulic fracturing. *Proc. Natl. Acad. Sci.* **2011**, *108* (20), 8172–8176.
- (97) Li, H.; Carlson, K. H. Distribution and origin of groundwater methane in the Wattenberg oil and gas field of Northern Colorado. *Environ. Sci. Technol.* **2014**, 48 (3), 1484–1491 DOI: 10.1021/es404668b.
- (98) Butler, K.; Engleberg, D.; Ross, J.; Soule, G. B. Response to Congressional Inquiry Regarding the EPA's Emergency Order to the Range Resources Gas Drilling Company, 14-P-0044; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2013.
- (99) Dussealt, M.; Nawrocki, G. M. Why Oil Wells Leak: Cement Behavior and Long-Term Consequences. *International Oil and Gas Conference and Exhibition in China*, SPE 64733; Society of Petroleum Engineers: Richardson, TX, USA, 2000.
- (100) Jackson, R. E.; Gorody, A. W.; Mayer, B.; Roy, J. W.; Ryan, M. C.; Van Stempvoort, D. R. Groundwater protection and unconventional gas extraction: The critical need for field-based hydrogeological research. *Groundwater* **2013**, *51* (4), 488–510 DOI: 10.1111/gwat.12074.
- (101) EPIWIN, version 4.11; U.S. EPA & Syracuse Research Corp.: Washington, DC, and Syracuse, NY, USA, 2012.
- (102) Leung, H. W. Ecotoxicology of glutaraldehyde: Review of environmental fate and effects studies. *Ecotoxicol. Environ. Saf.* **2001**, 49 (1), 26–39 DOI: 10.1006/eesa.2000.2031.
- (103) Index of Robust Summaries for ADBAC Joint Venture HPV Chemicals Challenge: Appendix A; Code No. 201-16856B; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2011.
- (104) Canadian Water Quality Guidelines for the Protection of Aquatic Life (DDAC); Canadian Council of Ministers of the Environment: Winnipeg, Canada, 1999.

- (105) Cornelissen, G.; Gustafsson, Ö.; Bucheli, T. D.; Jonker, M. T. O.; Koelmans, A. A.; Van, N.; Paul, C. M. Extensive sorption of organic compounds to black carbon, coal, and kerogen in sediments and soils: Mechanisms and consequences for distribution, bioaccumulation, and biodegradation. *Environ. Sci. Technol.* **2005**, 39 (18), 6881–6895 DOI: 10.1021/es050191b.
- (106) Harney, M. B.; Pant, R. R.; Fulmer, P. A.; Wynne, J. H. Surface self-concentrating amphiphilic quaternary ammonium biocides as coating additives. *ACS Appl. Mater. Interfaces* **2009**, *1* (1), 39–41 DOI: 10.1021/am800046r.
- (107) Cowan, C. T.; White, D. The mechanism of exchange reactions occurring between sodium montmorillonite and various n-primary aliphatic amine salts. *Trans. Faraday Soc.* **1958**, *54*, 691–697.
- (108) Kwolek, T.; Hodorowicz, M.; Stadnicka, K.; Czapkiewicz, J. Adsorption isotherms of homologous alkyldimethylbenzylammonium bromides on sodium montmorillonite. *J. Colloid Interface Sci.* **2003**, 264 (1), 14–19 DOI: 10.1016/S0021-9797(03)00414-4.
- (109) Xu, S.; Boyd, S. A. Cationic surfactant sorption to a vermiculitic subsoil via hydrophobic bonding. *Environ. Sci. Technol.* **1995**, 29 (2), 312–320.
- (110) Wang, Y.-S.; Yen, J.-H.; Hsieh, Y.-N.; Chen, Y.-L. Dissipation of 2, 4-D glyphosate and paraquat in river water. *Water, Air, Soil Pollut.* **1994**, 72 (1–4), 1–7.
- (111) Atkin, R.; Craig, V. S. J.; Biggs, S. Adsorption kinetics and structural arrangements of cationic surfactants on silica surfaces. *Langmuir* **2000**, *16* (24), 9374–9380 DOI: 10.1021/la0001272.
- (112) Lu, L.; Zhu, L. Reducing plant uptake of pahs by cationic surfactant-enhanced soil retention. *Environ. Pollut.* **2009**, *157* (6), 1794–1799 DOI: 10.1016/j.envpol.2009.01.028.
- (113) Pan, G.; Jia, C.; Zhao, D.; You, C.; Chen, H.; Jiang, G. Effect of cationic and anionic surfactants on the sorption and desorption of perfluorooctane sulfonate (PFOS) on natural sediments. *Environ. Pollut.* **2009**, *157* (1), 325–330 DOI: 10.1016/j.envpol.2008.06.035.
- (114) Wagner, J.; Chen, H.; Brownawell, B. J.; Westall, J. C. Use of cationic surfactants to modify soil surfaces to promote sorption and retard migration of hydrophobic organic compounds. *Environ. Sci. Technol.* 1994, 28 (2), 231–237 DOI: 10.1021/es00051a008.
- (115) Ismail, Z. Z.; Tezel, U.; Pavlostathis, S. G. Sorption of quaternary ammonium compounds to municipal sludge. *Water Res.* **2010**, 44 (7), 2303–2313 DOI: 10.1016/j.watres.2009.12.029.
- (116) García, M. T.; Campos, E.; Sánchez-Leal, J.; Comelles, F. Sorption of alkyl benzyl dimethyl ammonium compounds by activated sludge. *J. Dispersion Sci. Technol.* **2006**, 27 (5), 739–744 DOI: 10.1080/01932690600662554.
- (117) Eggen, T.; Heimstad, E. S.; Stuanes, A. O.; Norli, H. R. Uptake and translocation of organophosphates and other emerging contaminants in food and forage crops. *Environ. Sci. Pollut. Res. Int.* **2013**, *20* (7), 4520–4531 DOI: 10.1007/s11356-012-1363-5.
- (118) Ten Hulscher, T. E. M.; Cornelissen, G. Effect of temperature on sorption equilibrium and sorption kinetics of organic micropollutants—A review. *Chemosphere* **1996**, 32 (4), 609–626.
- (119) Montgomery, S. L.; Jarvie, D. M.; Bowker, K. A.; Pollastro, R. M. Mississippian Barnett shale, Fort Worth Basin, North-Central Texas: Gas-shale play with multi-trillion cubic foot potential. *AAPG Bull.* **2005**, 89 (2), 155–175.
- (120) Krooss, B. M.; Van Bergen, F.; Gensterblum, Y.; Siemons, N.; Pagnier, H. J. M.; David, P. High-pressure methane and carbon dioxide adsorption on dry and moisture-equilibrated Pennsylvanian coals. *Int. J. Coal Geol.* **2002**, *51* (2), 69–92.
- (121) Barbot, E.; Vidic, N.; Gregory, K. B.; Vidic, R. D. Spatial and temporal correlation of water quality parameters of produced waters from Devonian-age shale following hydraulic fracturing. *Environ. Sci. Technol.* **2013**, 47 (6), 2562–2569 DOI: 10.1021/es304638h.
- (122) Haluszczak, L. O.; Rose, A. W.; Kump, L. R. Geochemical evaluation of flowback brine from Marcellus gas wells in Pennsylvania, USA. *Appl. Geochem.* **2013**, 28, 55–61 DOI: 10.1016/j.apgeochem.2012.10.002.
- (123) Rowan, E. L.; Engle, M. A.; Kirby, C. S.; Kraemer, T. F. Radium Content of Oil- and Gas-Field Produced Waters in the Northern

- Appalachian Basin (USA): Summary and Discussion of Data, Scientific Investigations Report 2011-5135; U.S. Geological Survey, U.S. Government Printing Office: Washington, DC, USA, 2011.
- (124) Engle, M. A.; Rowan, E. L. Geochemical evolution of produced waters from hydraulic fracturing of the Marcellus shale, Northern Appalachian Basin: A multivariate compositional data analysis. *Int. J. Coal Geol.* **2013**, *126*, 45–56.
- (125) Delle Site, A. Factors affecting sorption of organic compounds in natural sorbent/water systems and sorption coefficients for selected pollutants. A review. *J. Phys. Chem. Ref. Data* **2001**, *30*, 187–439.
- (126) Ersoy, B.; Çelik, M. S. Effect of hydrocarbon chain length on adsorption of cationic surfactants onto clinoptilolite. *Clays Clay Miner.* **2003**, *51* (2), 172–180 DOI: 10.1346/CCMN.2003.0510207.
- (127) Droge, S.; Goss, K. U. Effect of sodium and calcium cations on the ion-exchange affinity of organic cations for soil organic matter. *Environ. Sci. Technol.* **2012**, *46* (11), 5894–5901 DOI: 10.1021/es204449r.
- (128) Knezovich, J. P.; Lawton, M. P.; Inouye, L. S. Bioaccumulation and tissue distribution of a quaternary ammonium surfactant in three aquatic species. *Bull. Environ. Contam. Toxicol.* **1989**, 42 (1), 87–93.
- (129) Pateiro-Moure, M.; Arias-Estévez, M.; Simal-Gándara, J. Critical review on the environmental fate of quaternary ammonium herbicides in soils devoted to vineyards. *Environ. Sci. Technol.* **2013**, 47 (10), 4984–4998 DOI: ds.doi.org/10.1021/es400755h.
- (130) Saiyood, S.; Vangnai, A. S.; Thiravetyan, P.; Inthorn, D. Bisphenol a removal by the Dracaena plant and the role of plant-associating bacteria. *J. Hazard. Mater.* **2010**, *178* (1–3), *777*–785 DOI: 10.1016/j.jhazmat.2010.02.008.
- (131) Jovović, M.; Kostić, N.; Jančić-Stojanović, B.; Malenović, A. Investigation of tropicamide and benzalkonium chloride stability using liquid chromatography. *J. Liq. Chromatogr. Relat. Technol.* **2012**, 35 (2), 231–239 DOI: 10.1080/10826076.2011.597072.
- (132) Reregistration Eligibility Decision for Aliphatic Alkyl Quaternaries (DDAC), EPA739-R-06-008; United States Environmental Agency, U.S. Government Printing Office: Washington, DC, USA, 2006.
- (133) Reregistration Eligibility Decision for Alkyl Dimethyl Benzyl Ammonium Chloride (ADBAC), EPA739-R-06-009; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2006.
- (134) Henderson, N. D. Environmental Impact and Toxic Effects of DDAC; Environmental Protection Division, BC Ministry of Environment, Lands and Parks: Victoria, British Columbia, Canada, 1992.
- (135) Migneault, I.; Dartiguenave, C.; Bertrand, M. J.; Waldron, K. C. Glutaraldehyde: Behavior in aqueous solution, reaction with proteins, and application to enzyme crosslinking. *BioTechniques* **2004**, *37* (5), 790–806.
- (136) Exner, J. H.; Burk, G. A.; Kyriacou, D. Rates and products of decomposition of 2,2-dibromo-3-nitrilopropionamide. *J. Agric. Food Chem.* **1973**, 21 (5), 838–842.
- (137) Reregistration Eligibility Decision Bronopol; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 1995.
- (138) Swenberg, J. A.; Kerns, W. D.; Mitchell, R. I.; Gralla, E. J.; Pavkov, K. L. Induction of squamous cell carcinomas of the rat nasal cavity by inhalation exposure to formaldehyde vapor. *Cancer Res.* **1980**, 40 (9), 3398–3402.
- (139) Loeppky, R. N. Nitrosamine and N-Nitroso Compound Chemistry and Biochemistry; ACS Symposium Series; American Chemical Society: Washington, DC, USA, 1994.
- (140) Challis, B. C.; Yousaf, T. I. Facile formation of N-nitrosamines from bromonitromethane and secondary amines. *J. Chem. Soc., Chem. Commun.* **1990**, 22, 1598–1599.
- (141) Dunnett, P. C.; Telling, G. M. Study of the fate of bronopol and the effects of antioxidants on N-nitrosamine formation in shampoos and skin creams. *Int. J. Cosmet. Sci.* **1984**, *6* (5), 241–247 DOI: 10.1111/j.1467-2494.1984.tb00381.x.
- (142) Scoping Document: Product Chemistry/Environmental Fate/ Ecotoxicity of: Tetrakis (Hydroxymethyl) Phosphonium Sulfate; EPA-HQ-OPP-2011-0067-0003; United States Environmental Protection

- Agency, U.S. Government Printing Office: Washington, DC, USA, 2011.
- (143) Zhao, K.; Wen, J.; Gu, T.; Kopliku, A.; Cruz, I. Mechanistic modeling of anaerobic Thps degradation in seawater under various conditions. *CORROSION 2008*; NACE International: Houston, TX, USA, 2008.
- (144) Ya, L.; Shuangxi, S.; Kaiqi, S.; Lan, J. Release of free formaldehyde in THP salt tannages. *J. Soc. Leather Technol. Chem.* **2008**, 92, 167–169.
- (145) Material Safety Data Sheet for Dazomet; AMVAC Chemical Corp.: Los Angeles, CA, USA, 2011.
- (146) Amended Reregistration Eligibility Decision for Dazomet, EPA 738-R-09-309; United States Environmental Protection Agency (USEPA), U.S. Government Printing Office: Washington, DC, USA, 2009
- (147) NRA Special Review of Metham Sodium, Dazomet and Methylisothiocyanate (Mitc); National Registration Authority: Canberra, Australia, 1997.
- (148) Douglass, M. L.; Kabacoff, B. L.; Anderson, G. A.; Chent, M. C. The chemistry of nitrosamine formation, inhibition and destruction. *J. Soc. Cosmet. Chem.* **1978**, 29, 581–606.
- (149) Lee, Y.; Von Gunten, U. Oxidative transformation of micropollutants during municipal wastewater treatment: Comparison of kinetic aspects of selective (chlorine, chlorine dioxide, ferrate vi, and ozone) and non-selective oxidants (hydroxyl radical). *Water Res.* **2010**, 44 (2), 555–566 DOI: 10.1016/j.watres.2009.11.045.
- (150) Luft, A.; Wagner, M.; Ternes, T. A. Transformation of biocides irgarol and terbutryn in the biological wastewater treatment. *Environ. Sci. Technol.* **2014**, *48* (1), 244–254 DOI: 10.1021/es403531d.
- (151) Garcia, M. T.; Campos, E.; Sanchez-Leal, J.; Ribosa, I. Effect of the alkyl chain length on the anaerobic biodegradability and toxicity of quaternary ammonium based surfactants. *Chemosphere* **1999**, 38 (15), 3473–3483.
- (152) Sutterlin, H.; Alexy, R.; Coker, A.; Kummerer, K. Mixtures of quaternary ammonium compounds and anionic organic compounds in the aquatic environment: Elimination and biodegradability in the closed bottle test monitored by LC-MS/MS. *Chemosphere* **2008**, 72 (3), 479–484 DOI: 10.1016/j.chemosphere.2008.03.008.
- (153) Nishihara, T.; Okamoto, T.; Nishiyama, N. Biodegradation of didecyldimethylammonium chloride by pseudomonas fluorescens TN4 isolated from activated sludge. *J. Appl. Microbiol.* **2000**, *88* (4), 641–647.
- (154) Seewald, J. S. Aqueous geochemistry of low molecular weight hydrocarbons at elevated temperatures and pressures: Constraints from mineral buffered laboratory experiments. *Geochim. Cosmochim. Acta* **2001**, *65* (10), 1641–1664.
- (155) Seewald, J. S. Organic-inorganic interactions in petroleum-producing sedimentary basins. *Nature.* **2003**, *426* (6964), 327–333.
- (156) Borch, T.; Kretzschmar, R.; Kappler, A.; Cappellen, P. V.; Ginder-Vogel, M.; Voegelin, A.; Campbell, K. Biogeochemical redox processes and their impact on contaminant dynamics. *Environ. Sci. Technol.* **2010**, 44 (1), 15–23 DOI: 10.1021/es9026248.
- (157) The Real Facts about Fracture Stimulation; American Exploration & Production Council: Washington, DC, USA, 2013.
- (158) Shipp, J.; Gould, I. R.; Herckes, P.; Shock, E. L.; Williams, L. B.; Hartnett, H. E. Organic functional group transformations in water at elevated temperature and pressure: Reversibility, reactivity, and mechanisms. *Geochim. Cosmochim. Acta* **2013**, *104*, 194–209 DOI: 10.1016/j.gca.2012.11.014.
- (159) Seewald, J. S.; Zolotov, M. Y.; Mccollom, T. Experimental investigation of single carbon compounds under hydrothermal conditions. *Geochim. Cosmochim. Acta* **2006**, 70 (2), 446–460 DOI: 10.1016/j.gca.2005.09.002.
- (160) Siskin, M.; Katritzky, A. R. Reactivity of organic compounds in superheated water: General background. *Chem. Rev.* **2001**, *101* (4), 825–836 DOI: 10.1021/cr000088z.
- (161) Comprehensive Everglades Restoration Plan 2009 System Status Report. Appendix—Indicator 1.1. Comprehensive Everglades

- Restoration Plan (CERP); USACE and South Florida Water Management District: West Palm Beach, FL, USA, 2010.
- (162) Reregistration Eligibility Decision for Glutaraldehyde, EPA 739-R-07-006; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2007.
- (163) Material Safety Data Sheet for DBNPA; Dow Chemical: Midland, MI, USA, 2012.
- (164) Chen, F. The chronic aquatic toxicity of a microbicide dibromonitrilopropionamide. *Toxicol Ind. Health.* **2012**, 28 (2), 181–185 DOI: 10.1177/0748233711410904.
- (165) Mayes, M. A.; Blanchard, F. A.; Hopkins, D. L.; Takahashi, I. T. Static acute toxicity of dibromonitrilopropionamide and selected degradation products to the fathead minnow (Pimephales promelas Rafinesque). *Environ. Toxicol. Chem.* 1985, 4 (6), 823–830.
- (166) Material Safety Data Sheet for Dibromoacetonitrile; Sigma-Aldrich: St. Louis, MO, USA, 2013.
- (167) Material Safety Data Sheet for 2-Bromo-2-nitro-1,3-propanediol; Sigma-Aldrich: St. Louis, MO, USA, 2014.
- (168) Agency's Response to the Registrant Response for Acute Neurotoxicity and Repeated Exposure Inhalation Study and Determination of Inhalation Moe and Developmental Toxicity (TTPC), Memorandum; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2006.
- (169) Bilthoven, G. J. Flame Retardants: Tris(2-Butoxyethyl) Phosphate, Tris(2-Ethylhexyl) Phosphate, and Tetrakis(hydroxymethyl) Phosphonium Salts, Environmental Health Criteria 218; World Health Organization: Geneva, Switzerland, 2000.
- (170) He, K.; Huang, J.; Lagenaur, C. F.; Aizenman, E. Methylisothiazolinone, a neurotoxic biocide, disrupts the association of SRC family tyrosine kinases with focal adhesion kinase in developing cortical neurons. *J. Pharmacol. Exp. Ther.* **2006**, 317 (3), 1320–1329 DOI: 10.1124/jpet.106.103044.
- (171) Assaf, N.; Salem, N. A.; Khalil, W. K.; Ahmed, H. H. Evaluation of the potential toxicity of dibromoacetonitrile-induced apoptosis and tumor-initiating activity in rat liver. *Food Chem. Toxicol.* **2011**, 49 (12), 3055–3062 DOI: 10.1016/j.fct.2011.09.030.
- (172) NTP Technical Report of the Toxicology and Carcinogenesis Studies of Dibromoacetonitrile in F344/n Rats and B6c3f1Mice, NTP TR 544, NIH Publication No. 10-5886; National Toxicology Program, National Institutes of Health Public Health Service, U.S. Department of Health and Human Services: Research Triangle Park, NC, USA, 2010.
- (173) Lijinsky, W.; Epstein, S. S. Nitrosamines as environmental carcinogens. *Nature* **1970**, 225, 21–23.
- (174) Matubayasi, N.; Morooka, S.; Nakahara, M.; Takahashi, H. Chemical equilibrium of formaldehyde and methanediol in hot water: Free-energy analysis of the solvent effect. *J. Mol. Liq.* **2007**, *134* (1–3), 58–63 DOI: 10.1016/j.molliq.2006.12.002.
- (175) Ottosson, N.; Aziz, E. F.; Bradeanu, I. L.; Legendre, S.; Öhrwall, G.; Svensson, S.; Björneholm, O.; Eberhardt, W. An electronic signature of hydrolysation in the X-ray absorption spectrum of aqueous formaldehyde. *Chem. Phys. Lett.* **2008**, *460* (4–6), 540–542 DOI: 10.1016/j.cplett.2008.06.043.
- (176) Tetrakis (Hydroxymethyl) Phosphonium Sulfate (THPS) Summary Document: Registration Review, Docket No. EPA-HQ-OPP-2011-0067; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2011.
- (177) Summary of Human Health Effects Data for the THPS Registration Review Decision Document, EPA-HQ-OPP-2011-0067-0011; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2011.
- (178) Van Duuren, B. L.; Loewengart, G.; Seidman, I.; Smith, A. C.; Melchionne, S. Mouse skin carcinogenicity tests of the flame retardants tris(2,3-dibromopropyl) phosphate, tetrakis-(hydroxymethyl) phosphonium chloride, and polyvinyl bromide. *Cancer Res.* 1978, 38 (10), 3236–3240.
- (179) Speit, G.; Neuss, S.; Schutz, P.; Frohler-Keller, M.; Schmid, O. The genotoxic potential of glutaraldehyde in mammalian cells in vitro

- in comparison with formaldehyde. *Mutat. Res.* **2008**, *649* (1–2), 146–154 DOI: 10.1016/j.mrgentox.2007.08.010.
- (180) Zeiger, E.; Gollapudi, B.; Spencer, P. Genetic toxicity and carcinogenicity studies of glutaraldehyde—A review. *Mutat. Res.* **2005**, 589 (2), 136–151 DOI: 10.1016/j.mrrev.2005.01.001.
- (181) Carcinogenesis Studies of Glutaraldehyde (CAS No. 111-30-8) in F344/n Rats and B6c3f1Mice (Inhalation Studies), NTP TR 490, NIH Publication No. 99-3980; National Toxicology Program (NTP), U.S. Department of Health and Human Services: Research Triangle Park, NC, USA, 1999.
- (182) Ferk, F.; Misik, M.; Hoelzl, C.; Uhl, M.; Fuerhacker, M.; Grillitsch, B.; Parzefall, W.; Nersesyan, A.; Micieta, K.; Grummt, T.; Ehrlich, V.; Knasmuller, S. Benzalkonium chloride (Bac) and dimethyldioctadecyl-ammonium bromide (Ddab), two common quaternary ammonium compounds, cause genotoxic effects in mammalian and plant cells at environmentally relevant concentrations. *Mutagenesis.* **2007**, *22* (6), 363–370 DOI: 10.1093/mutage/gem027.
- (183) Deutschle, T.; Porkert, U.; Reiter, R.; Keck, T.; Riechelmann, H. In vitro genotoxicity and cytotoxicity of benzalkonium chloride. *Toxicol. In Vitro* **2006**, 20 (8), 1472–1477 DOI: 10.1016/j.tiv.2006.07.006.
- (184) Sanchez-Fortun, S.; Llorente, M. T.; Castano, A. Genotoxic effects of selected biocides on Rtg-2 fish cells by means of a modified fast micromethod assay. *Aquat. Toxicol.* **2005**, 73 (1), 55–64 DOI: 10.1016/j.aquatox.2005.03.002.
- (185) Chen, Y.; Geurts, M.; Sjollema, S. B.; Kramer, N. I.; Hermens, J. L.; Droge, S. T. Acute toxicity of the cationic surfactant C12-benzalkonium in different bioassays: How test design affects bioavailability and effect concentrations. *Environ. Toxicol. Chem.* **2014**, 33 (3), 606–615 DOI: 10.1002/etc.2465.
- (186) THPS Biocides: A New Class of Antimicrobial Chemistry. 1997 Designing Greener Chemicals Award; United States Environmental Protection Agency: 1997. http://www2.epa.gov/greenchemistry/1997-designing-greener-chemicals-award.
- (187) Veschetti, E.; Cutilli, D.; Bonadonna, L.; Briancesco, R.; Martini, C.; Cecchini, G.; Anastasi, P.; Ottaviani, M. Pilot-plant comparative study of peracetic acid and sodium hypochlorite wastewater disinfection. *Water Res.* **2003**, *37* (1), 78–94.
- (188) Peroxyacetic Acid; Exemption from the Requirement of a Tolerance. *United States Environmental Protection Agency, Federal Register*; U.S. Government Printing Office; Washington, DC, USA, 2009.
- (189) Glaze, W. H.; Kang, J. W.; Chapin, D. H. The chemistry of water treatment processes involving ozone, hydrogen peroxide and W radiation. *Ozone: Sci. Eng.* 1987, 9 (4), 335–352.
- (190) Glaze, W. H. Drinking-water treatment with ozone. *Environ. Sci. Technol.* **1987**, 21 (3), 224–230.
- (191) Kirn, J. G.; Yousef, A. E.; Dave, S. Application of ozone for enhancing the microbiological safety and quality of foods. *J. Food Prot.* **1999**, *62* (9), 1071–1080.
- (192) Richardson, S. D.; Thruston, A. D.; Caughran, T. V.; Chen, P. H.; Collette, T. W.; Floyd, T. L.; Schenck, K. M.; Lykins, B. W.; Sun, G.; Majetich, G. Identification of new ozone disinfection byproducts in drinking water. *Environ. Sci. Technol.* **1999**, 33 (19), 3368–3377 DOI: 10.1021/es981218c.
- (193) Hua, G.; Reckhow, D. A. Comparison of disinfection byproduct formation from chlorine and alternative disinfectants. *Water Res.* **2007**, *41* (8), 1667–1678 DOI: 10.1016/j.watres.2007.01.032.
- (194) Weinhold, B. Ozone nation: EPA standard panned by the people. *Environ. Health Perspect.* **2008**, *116* (7), A 302–A 305.
- (195) Lu, J.; Zhang, T.; Ma, J.; Chen, Z. Evaluation of disinfection by-products formation during chlorination and chloramination of dissolved natural organic matter fractions isolated from a filtered river water. *J. Hazard. Mater.* **2009**, *162* (1), 140–145 DOI: 10.1016/j.jhazmat.2008.05.058.
- (196) Duong, H. A.; Berg, M.; Hoang, M. H.; Pham, H. V.; Gallard, H.; Giger, W.; Gunten, U. V. Trihalomethane formation by chlorination of ammonium- and bromide-containing groundwater in

- water supplies of Hanoi, Vietnam. Water Res. 2003, 37 (13), 3242–3252 DOI: 10.1016/S0043-1354(03)00138-6.
- (197) Hladik, M. L.; Focazio, M. J.; Engle, M. Discharges of produced waters from oil and gas extraction via wastewater treatment plants are sources of disinfection by-products to receiving streams. *Sci. Total Environ.* **2014**, 466–467, 1085–1093 DOI: 10.1016/j.scitotenv.2013.08.008.
- (198) States, S.; Cyprych, G.; Stoner, M.; Wydra, F.; Kuchta, J.; Monnell, J.; Casson, L. Marcellus shale drilling and brominated THMS in Pittsburgh, PA., drinking water (PDF). *J.*—*Am. Water Works Assoc.* **2013**, *105* (8), E432–E448.
- (199) Elkins, H. B. Chlorine Dioxide, ClO₂, The Chemistry of Industrial Toxicology; John Wiley & Sons: New York, NY, 1950.
- (200) Londry, K. L.; Suflita, J. M. Use of nitrate to control sulfide generation by sulfate-reducing bacteria associated with oily waste. *J. Ind. Microbiol. Biotechnol.* **1999**, 22 (6), 582–589.
- (201) Stott, J. F. D. Implementation of nitrate treatment for reservoir souring control: Complexities and pitfalls. *SPE International Conference and Exhibition on Oilfield Corrosion*, Aberdeen, U.K.; Society of Petroleum Engineers: Richardson, TX, USA, 2012.
- (202) Jurelevicius, D.; Von Der Weid, I.; Korenblum, E.; Valoni, E.; Penna, M.; Seldin, L. Effect of nitrate injection on the bacterial community in a water-oil tank system analyzed by PCR-DGGE. *J. Ind. Microbiol. Biotechnol.* **2008**, 35 (4), 251–255 DOI: 10.1007/s10295-007-0297-2.
- (203) Kjellerup, B. V.; Veeh, R. H.; Sumithraratne, P.; Thomsen, T. R.; Buckingham-Meyer, K.; Frolund, B.; Sturman, P. Monitoring of microbial souring in chemically treated, produced-water biofilm systems using molecular techniques. *J. Ind. Microbiol. Biotechnol.* **2005**, 32 (4), 163–170 DOI: 10.1007/s10295-005-0222-5.
- (204) Davidova, I.; Hicks, M. S.; Fedorak, P. M.; Suflita, J. M. The influence of nitrate on microbial processes in oil industry production waters. *J. Ind. Microbiol. Biotechnol.* **2001**, *27* (2), 80–86.
- (205) Hubert, C.; Voordouw, G. Oil field souring control by nitrate-reducing Sulfurospirillum Spp. that outcompete sulfate-reducing bacteria for organic electron donors. *Appl. Environ. Microbiol.* **2007**, 73 (8), 2644–2652 DOI: 10.1128/AEM.02332-06.
- (206) Hulsmans, A.; Joris, K.; Lambert, N.; Rediers, H.; Declerck, P.; Delaedt, Y.; Ollevier, F.; Liers, S. Evaluation of process parameters of ultrasonic treatment of bacterial suspensions in a pilot scale water disinfection system. *Ultrason. Sonochem.* **2010**, *17* (6), 1004–1009 DOI: 10.1016/j.ultsonch.2009.10.013.
- (207) Halliburton. Halliburton Cleansuite Technologies, 2014, http://www.halliburton.com/public/projects/pubsdata/Hydraulic_Fracturing/CleanSuite Technologies.html.
- (208) Robinson, G.; Thorn, R.; Reynolds, D. The effect of long-term storage on the physiochemical and bactericidal properties of electrochemically activated solutions. *Int. J. Mol. Sci.* **2012**, *14* (1), 457–469 DOI: 10.3390/ijms14010457.
- (209) Electrochemistry and the environment. J. Appl. Electrochem.. 1994, 24 (11), 1077–1091.
- (210) Jeon, J.; Kurth, D.; Hollender, J. Biotransformation pathways of biocides and pharmaceuticals in freshwater crustaceans based on structure elucidation of metabolites using high resolution mass spectrometry. *Chem. Res. Toxicol.* **2013**, *26* (3), 313–324 DOI: 10.1021/tx300457f.
- (211) Chiaia-Hernandez, A. C.; Krauss, M.; Hollender, J. Screening of lake sediments for emerging contaminants by liquid chromatography atmospheric pressure photoionization and electrospray ionization coupled to high resolution mass spectrometry. *Environ. Sci. Technol.* **2013**, 47 (2), 976–986 DOI: 10.1021/es303888v.
- (212) Moschet, C.; Gotz, C.; Longree, P.; Hollender, J.; Singer, H. Multi-level approach for the integrated assessment of polar organic micropollutants in an international lake catchment: The example of Lake Constance. *Environ. Sci. Technol.* **2013**, 47 (13), 7028–7036 DOI: 10.1021/es304484w.
- (213) Schymanski, E. L.; Singer, H. P.; Longree, P.; Loos, M.; Ruff, M.; Stravs, M. A.; Ripolles Vidal, C.; Hollender, J. Strategies to characterize polar organic contamination in wastewater: Exploring the

- capability of high resolution mass spectrometry. *Environ. Sci. Technol.* **2014**, 48 (3), 1811–1818 DOI: 10.1021/es4044374.
- (214) Huntscha, S.; Rodriguez Velosa, D. M.; Schroth, M. H.; Hollender, J. Degradation of polar organic micropollutants during riverbank filtration: complementary results from spatiotemporal sampling and push-pull tests. *Environ. Sci. Technol.* **2013**, 47 (20), 11512–11521 DOI: 10.1021/es401802z.
- (215) Moschet, C.; Piazzoli, A.; Singer, H.; Hollender, J. Alleviating the reference standard dilemma using a systematic exact mass suspect screening approach with liquid chromatography-high resolution mass spectrometry. *Anal. Chem.* **2013**, 85 (21), 10312–10320 DOI: 10.1021/ac4021598.
- (216) Nelson, A. W.; May, D.; Knight, A. W.; Eitrheim, E. S.; Mehrhoff, M.; Shannon, R.; Litman, R.; Schultz, M. K. Matrix complications in the determination of radium levels in hydraulic fracturing flowback water from Marcellus shale. *Environ. Sci. Technol. Lett.* **2014**, *1* (3), 204–208 DOI: 10.1021/ez5000379.
- (217) Bibby, K. J.; Brantley, S. L.; Reible, D. D.; Linden, K. G.; Mouser, P. J.; Gregory, K. B.; Ellis, B. R.; Vidic, R. D. Suggested reporting parameters for investigations of wastewater from unconventional shale gas extraction. *Environ. Sci. Technol.* **2013**, 47 (23), 13220–13221 DOI: 10.1021/es404960z.
- (218) Physical/Chemical Property Database (Physprop); SRC Environmental Science Center, Syracuse Research Corp.: Syracuse, NY, USA, 1994.
- (219) Yalkowsky, S. H.; He, Y. Handbook of Aqueous Solubility Data; CRC Press: Boca Raton, FL, USA, 2003.
- (220) Material Safety Data Sheet for Chlorine Dioxide; 2004.
- (221) Reregistration Eligibility Decision (RED) for Chlorine Dioxide and Sodium Chlorite, EPA 738-R-06-007; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 2006.
- (222) Material Safety Data Sheet for 5-Chloro-2-methyl-4-isothiazolin-3-one Solution; Sigma-Aldrich: St. Louis, MO, USA, 2013.
- (223) Reregistration Eligibility Decision (RED) Methylisothiazolinone, EPA738-R-98-012; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 1998.
- (224) Du, S.; Mclaughlin, B.; Pal, S.; Aizenman, E. In vitro neurotoxicity of methylisothiazolinone, a commonly used industrial and household biocide, proceeds via a zinc and extracellular signal-regulated kinase mitogen-activated protein kinase-dependent pathway. *J. Neurosci.* 2002, 22 (17), 7408–7416.
- (225) Fatty Nitrogen Derived Cationics Category High Production Volume (HPV) Chemicals Challenge Assessment of Data Availability, AR201–13407A; United States Environmental Protection Agency, Toxicology/Regulatory Services, Inc., U.S. Government Printing Office: Washington, DC, USA, 2001.
- (226) Material Safety Data Sheet for 4,4-Dimethyloxazolidine; 2012.
- (227) Reregistration Eligibility Decision (RED) 4,4-Dimethyloxazolidine, EPA-738-R-96-007; United States Environmental Protection Agency, U.S. Government Printing Office: Washington, DC, USA, 1996.
- (228) Material Safety Data Sheet: Glutaraldehyde Solution, 50%; 2012.
- (229) Material Safety Data Sheet-Extented: Sodium Hypochlorite, 5%; Oltchim: Valcea, Romania, 2011.
- (230) Material Safety Data Sheet for N-Bromosuccinimide; Sigma-Aldrich: St. Louis, MO, USA, 2013.
- (231) Material Safety Data Sheet Peracetic Acid; Sigma-Aldrich; St. Louis, MO, USA, 2013.
- (232) Reregistration Eligibility Decision Peroxy Compounds, EPA-738-F-93-026; United States Environmental Agency, U.S. Government Printing Office: Washington, DC, USA, 1993.
- (233) Summary Review of Available Literature for Hydrogen Peroxide and Peroxyacetic Acid for New Use to Treat Wastewater; United States Environmental Protection Agency, Office of Prevention, Pesticides, and Toxic Substances, U.S. Government Printing Office: Washington, DC, USA, 2007.
- (234) Material Safety Data Sheet: Bellacide 350 (TTPC); BWA Water Additives: Manchester, U.K., 2009.