Physicochemical Characterization of Horizontal Directional Drilling Residuals

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Abstract: Horizontal directional drilling (HDD) is a trenchless technology mainly used for the installation of gas, sewer lines, and fiber optic cables. Spent HDD fluid is a by-product of the boring process. However, little is known of its constituency, although the land application of spent HDD residuals would be an economical and less time-consuming method of disposal. Physicochemical analysis of spent HDD residuals from broad geographic regions was conducted to determine if the land application would be an environmentally safe option for disposal. Fifty-eight HDD samples were collected from 26 states throughout the United States. After separation of the liquid and solid portion, the materials were assessed for carbon (C), nitrogen (N), plant nutrients, soluble and total trace metals, total dissolved solids, sodium adsorption ratio (SAR), and potential “leachable metals”. All trace metal concentrations in the digested solid portion were well below the limits for biosolids set by the EPA 40 Part 503 rule for land application. Metal concentrations did not exceed the limits of drinking water standards. Besides the total amount of solids, there was nothing chemically limiting HDD residuals from land application. However, if boring through soil with suspected contamination, testing the residual before the application is recommended.

Keywords: residuals; horizontal directional drilling; industrial by-products; metals; synthetic precipitation leaching procedure; land application

1. Introduction

A wide variety of waste materials and by-products have been land-applied as soil amendments to improve soil quality for centuries [1]. Many by-products are applied as a means of waste disposal. Often the sources of these waste residuals are agricultural, municipal, and industrial wastes or by-products. These materials, in their broadest sense, vary widely in terms of their overall beneficial and environmental impacts. Information acquired over years of research suggests that sustainable land application systems can be established and maintained under a wide range of conditions [2]. Because of environmental concerns and economic restraints in regards to landfilling and incineration, public interest in land application of waste is continuing to grow [3]. As the interest in land application increases so does public speculation on potential contaminants added to the land. Perhaps the most important factor in regards to the land application of agricultural, municipal, and industrial by-products is earning the public’s trust concerning safety. Gaining the public’s trust will require waste management in such a way that protects human and animal health as well as safeguards soil and water resources, and maintains a quality ecosystem for the long term [4].

An industrial by-product with a growing need for disposal is the spent drilling fluid from the oil and natural gas industry. The chemical makeup and physical properties of drilling mud constitute
its potentially hazardous nature and ability to impact the environment [5]. Drilling muds generally contain bentonite clays, synthetic organic polymers, production water, barite, formation cuttings, and salts [6], but may have other constituents such as petroleum hydrocarbons, heavy metals, and high salt or sodium. The land application or disposal of spent oil and gas drilling fluid is regulated by state regulatory agencies.

The horizontal directional drilling (HDD) industry is on the rise worldwide. In North America alone, the number of drilling units grew from 12 units in 1984 to more than 2000 units in 1995 [7], and this number has only continued to grow over the last 25 years [8,9]. The directional drilling market in Europe occupies the third position in the industry, after the US and Africa, although it leads the market as far as drilling operations for telecommunication tools are concerned. This is probably a consequence of European government regulations regarding oil and gas field developments being highly strict. In the last decade, the European Union (EU) Parliament approved a new law on the safety of oil and gas operations in which oil and gas companies have to submit a special report on possible hazards before commencing any drilling operations in EU waters. Horizontal directional drilling is a trenchless technology and its popularity is likely a direct result of its noninvasive nature and reduced project costs [10,11]. It is usually used for the installation of underground utilities for municipalities ranging from 50 mm utility conduits to 900 mm sewer [10]. Spent HDD residuals are an ‘industrial’ by-product resulting from the drilling or boring process, which utilizes drilling fluid. Drilling fluids are used in the boring process as a means to stabilize the borehole, float cuttings to the surface, and keep the bit cool. Once the drilling fluid’s density reaches 1.078–1.198 mg mL\(^{-1}\) (9–10 lbs gallon\(^{-1}\)) it is no longer viable and needs to be disposed of [12]. At that point, the by-product is known as “mud” or “residual”. The HDD residual constituents are usually water, bentonite (montmorillonite), and/or polymers [12–14], as well as surfactants and soda ash when needed. These additives may or may not be added to aid in the drilling process, depending on the type of soil that is being drilled through. The regulations and bylaws for the proper disposal of the spent HDD residuals vary by municipality as well as by state and are not federally regulated. In Oklahoma, the Corporation Commission (OCC) is the regulating authority for disposal. While most states possess strict rules for the disposal of oil and gas drilling residuals, no regulations or guidelines are currently available for HDD residuals.

Horizontal directional drilling residual containment and disposal is a growing issue, as the number of directional drilling rigs is on the rise paired with the stigma of its association with spent oil and gas drilling mud [10]. Today there is growing speculation about HDD residuals and the safety of their constituents or chemical makeup. As stated above, it is often mistaken for oil and gas drilling mud even though the two by-products are very different. The HDD residuals are simply subsoils mixed with some additives originally used in the drilling fluid. There is a need to separate the public’s perception of the spent HDD drilling residuals from that of the oil and gas industry. To date, there is no published research on the chemical and physical makeup of spent HDD residuals or their environmental impact when applied on land. Thus, the objective of this research is to collect and fully characterize many HDD residuals from multiple locations in the United States in order to make an overall assessment of their suitability for land application.

2. Materials and Methods

2.1. HDD Residual Sample Collection and Preparation

For a broad overall characterization of HDD residuals, it was necessary to obtain samples from multiple locations where HDD operations took place. To do this, packages were sent to contractors throughout the country. These packages contained sample bottles, instructions on how to obtain a representative sample, by fulfilling the proper containers (1 L plastic bottles) with mud, and a written survey. A representative HDD residue sample was considered by either completely filling the sample bottle with mud over an extended period of time as it flows out of the borehole throughout the drilling job or taking samples directly from the ditch or “pit” shortly after the job is complete.
samples were received representing 26 US states across the US (Figure 1). Solid content (gravimetric) was determined by shaking samples for 20 min followed by subsampling and placement in an oven at 65 °C for 24 h. To conduct several analyses, the solid and liquid portion was separated by centrifugation at 5000 rpm for 10 min followed by filtration with a Whatman #41 paper. Since the as-is materials were normally occurring as slurries, presenting a wide range of moisture w/w and the solid contents ranging from 0.04 to 0.72 g solid g total slurry −1, it was more appropriate to perform the physicochemical analyses of samples in both conditions separated solid and water portions and discuss them separately.

![Figure 1. The US States where horizontal directional drilling (HDD) residues were collected (green).](image)

2.2. Analysis of Separated Solids Fraction

Two grams of solids were digested using EPA 3050 method (Environmental Protection Agency, EPA 1996) in trace metal grade nitric acid (HNO₃) and trace metal grade hydrochloric acid (HCl), and 30% hydrogen peroxide solution. The digests were filtered using Whatman #41 paper and brought to volume with deionized (DI) water in 50 mL volumetric flasks, and analyzed for various elements, including aluminum (Al), sodium (Na), sulfur (S), calcium (Ca), magnesium (Mg), phosphorus (P), boron (B), barium (Ba), potassium (K), chromium (Cr), manganese (Mn), iron (Fe), cobalt (Co), nickel (Ni), copper (Cu), zinc (Zn), cadmium (Cd), arsenic (As), and lead (Pb), using an Inductively Coupled Plasma Atomic Emission Spectroscopy (ICP-AES). The results were compared to the EPA Regional Residential Screening Levels (EPA, 2013) as well as EPA 40 CFR 503 Criteria for Exceptional Quality Biosolids (EPA 2007).

Plant-available nutrients P, K, Ca, and Mg were extracted from the separated solids fraction with Mehlich-3 (0.2 M CH₃COOH + 0.015 M NH₄F + 0.013 M HNO₃ + 0.001 M EDTA + 0.25 M H₂NO₃) at a ratio of 1:10 solids:solution for five minutes, followed by filtration with Whatman #42 filter paper and analysis by ICP-AES [15]. Total N and carbon (C) were quantified on the separated solids fraction using LECO TruSpec CN Analyzer (Leco Corporation, St. Joseph, MI, USA).

A modified version of the synthetic precipitation leaching procedure (SPLP) was also performed on the solid portion to simulate potential leachability to groundwater [16]. Extraction solution was made with DI water acidified to a pH of 4.2 using a 60:40 ratio solution of trace metal grade nitric acid (HNO₃) to reagent-grade sulfuric acid (H₂SO₄). Solids were extracted at a 1:20 solid to solution mass ratio for 18 h, followed by centrifugation at 5000 rpm for 10 min and filtration through 0.45 μm glass fiber filters using vacuum filtration. Extracts were analyzed for Mo, Al, B, Na, S, Ca, Mg, P, K, Cr, Mn, Fe, Co, Cu, Zn, Cd, As, and Pb by ICP-AES.
2.3. Analysis of Separated Liquid Fraction

The separated liquid portion was analyzed for Mo, Al, Na, S, Ca, Mg, P, K, Cr, Mn, Fe, Co, Cu, Zn, Cd, As, and Pb by ICP-AES. This is referred to as “water-soluble” contents. Results were compared to EPA Aquatic Life Criteria Table (EPA 2013). Sodium adsorption ratio was calculated for each sample as the following: \( \text{SAR} = \frac{\text{Na}}{\sqrt{((\text{Ca}+\text{Mg})/2)}} \), where Na, Ca, and Mg are in units of meq L\(^{-1}\). Electrical Conductivity (EC) and pH were tested on both HDD residual conditions, the as-is and the separated water portion, using standard EC and pH meters.

2.4. Quality Control

The analyses of the 58 samples included laboratory check samples for quality assurance and quality control (QA/QC). Reagent blanks and reference samples were analyzed every nine samples. Acceptable recoveries for the certified references were \( \pm 10\% \) of certified values. If a check sample failed control limits, then the nine samples analyzed before and after the failure were reanalyzed. Reference samples from the North American Proficiency Testing Program (NAPTP) were utilized for QA/QC of pH, Mehlich-3 P, K, Ca, and Mg, DC, total C, and total N. Certified biosolids-treated reference soil (CRM005–050, RTC Corp., Laramie, WY, USA) was included in the determination of the total element contents in the liquid and solid portions of HDD residuals. Acceptable method blank concentrations of all elements analyzed were below the established instrumental Limits of Detection (LOD), and the LOD of metals were in the range of 0.1 to 0.2 mg L\(^{-1}\).

2.5. Statistical Analysis

A descriptive statistical analysis was performed using Microsoft Excel (2013). Mean, median, minimum, maximum, and standard deviation were calculated for some elements of interest from the extraction methods listed above. The same parameters were also calculated for total C and N, pH, EC, solid content, and sodium adsorption ratio (SAR).

3. Results

3.1. Chemical Properties of HDD Residuals

Solid content had a broad range of 0.04–0.72 gr solid g slurry\(^{-1}\) with mean and median values of 0.35 and 0.34 g solid g total slurry\(^{-1}\), respectively (Table 1). This wide range is likely due to the variability of the original drilling fluid used and the various formations or sub-soils being bored through; for example, sometimes only water is used, without bentonite clay. The electrical conductivity of the residuals ranged from a low of 118 \(\mu\text{S cm}^{-1}\) to a high of 3950 \(\mu\text{S cm}^{-1}\) with mean and median values of 1198 and 1124 \(\mu\text{S cm}^{-1}\), respectively (Table 1). All samples were under the threshold of 4000 \(\mu\text{S cm}^{-1}\) for salt-sensitive plants [17]. Electrical conductivity and density were used to calculate loading rates of salt for all samples. Using an application rate of 112 metric tons of solids ha\(^{-1}\), the largest loading rate would apply 3723 kg ha\(^{-1}\) salt. This is well below the limit of 6724 kg total dissolved solids ha\(^{-1}\) allowed by the OCC for the land application of water-base drilling mud from oil and gas exploration (Oklahoma administrative code and register, Title 165:10-7-19). The majority of samples had pH values similar to that of a typical subsoil, ranging from 4.7 to 9.9 with a mean of 7.5 and a median of 7.6 (Table 1). The high pH samples were likely indicative of free carbonates, which could occur naturally in subsoils, or from bicarbonate additives in the original drilling fluid. Overall, the residuals were low in total carbon, ranging from 0.09% to 6.61% with a mean of 1.42 and a median of 0.51 (Table 1). Nitrogen content ranged from 0.01% to 0.26% with a mean and median of 0.04. Low C and N contents were expected, as the majority of the residual is smectitic clay and subsurface (deeper than 1 m) soil and cuttings. The lack of C and N in the residual makes it of little value as a potential C or N source if land applied.
Table 1. Descriptive statistics for electrical conductivity (EC), in µS cm\(^{-1}\), and pH for both the as-is and separated water portion, and percentage of total carbon (C), total nitrogen (N), and the carbon:nitrogen ratio (C:N) of the solid portion of the residuals. Solid content is expressed as g solid g total slurry\(^{-1}\). \(n = 58\).

| HDD Residues  | Parameter | Mean  | Median | Min   | Max   | Std Dev |
|---------------|-----------|-------|--------|-------|-------|---------|
| Residual as-is| Solid Content | 0.35  | 0.34   | 0.04  | 0.72  | 0.2     |
|               | EC        | 1198  | 1124   | 118   | 3950  | 772     |
|               | pH        | 7.5   | 7.6    | 4.7   | 9.9   | 1.2     |
| Solid Portion | Carbon    | 1.42  | 0.51   | 0.09  | 6.61  | 1.75    |
|               | Nitrogen  | 0.04  | 0.04   | 0.01  | 0.26  | 0.04    |
|               | C:N       | 42.89 | 14.04  | 2.97  | 276.39| 62.62   |
| Water Portion | EC        | 1439  | 1334   | 241   | 4296  | 848     |
|               | pH        | 7.7   | 8      | 4.2   | 10.1  | 1       |

HDD: horizontal directional drilling. Min: minimum value. Max: maximum value. Std Dev: Standard Deviation.

3.2. Characterization of the Separated Liquid Portion

Descriptive statistics of the whole dataset are presented in Table 2. Sodium adsorption ratio (SAR) was determined on the separated liquid portion of every sample. The mean SAR was 13 with a median of 12.5, ranging from 0.5 to 90.2 (Table 2). A soil with a SAR \(\geq 13\) is considered sodic at a pH \(> 8.5\) and an EC \(< 4000\) µS cm\(^{-1}\). High concentrations of Na can negatively impact soil structure and compete with the Ca and Mg for plant uptake [18]. Thus, depending on the amount of residual land applied, it may or may not cause any problem to the soil because sodium will be diluted once added to the soil. Most soils contain large amounts of Ca and Mg. If a residual with high SAR is applied in a large quantity on the surface with no incorporation, it could seal the surface, causing it to be physically dispersed. However, the incorporation of such a residual would prevent this problem from occurring.

Table 2. Descriptive statistics for the elemental concentrations, in mg L\(^{-1}\), and sodium adsorption ratio (SAR) of the separated liquid portion of HDD residual samples. Ceiling concentrations for the EPA Aquatic Life Criteria are shown as a reference. \(n = 58\).

| Element | Mean   | Median | Min   | Max   | Std Dev | EPA Aquatic Life Criteria ¹ |
|---------|--------|--------|-------|-------|---------|----------------------------|
| S       | 326.8  | 248.2  | 5.2   | 2708.3| 400.8   | NA                         |
| Na      | 285.5  | 272.7  | 18.0  | 856.2 | 191.2   | NA                         |
| Ca      | 47.0   | 27.2   | 0.8   | 584.7 | 80.9    | NA                         |
| Fe      | 16.9   | 0.4    | <LOD  | 183.1 | 40.8    | NA                         |
| Mg      | 16.4   | 10.9   | <LOD  | 74.9  | 17.6    | NA                         |
| K       | 8.4    | 5.2    | 0.7   | 78.2  | 11.4    | NA                         |
| Mn      | 1.1    | <LOD   | <LOD  | 23.4  | 3.3     | NA                         |
| P       | 0.6    | 0.1    | <LOD  | 26.6  | 3.5     | NA                         |
| B       | 0.2    | 0.1    | <LOD  | 1.5   | 0.3     | NA                         |
| Mo      | 0.1    | <LOD   | <LOD  | 0.6   | 0.1     | NA                         |
| Zn      | 0.1    | <LOD   | <LOD  | 0.3   | 0.1     | 0.12                       |
| Cr      | <LOD   | <LOD   | <LOD  | 0.5   | 0.1     | 0.57                       |
| Cu      | <LOD   | <LOD   | <LOD  | 0.3   | 0.1     | NA                         |
| Cd      | <LOD   | <LOD   | <LOD  | 0.2   | NA      | 0.002                      |
| Co      | <LOD   | <LOD   | <LOD  | 0.1   | NA      | NA                         |
| As      | <LOD   | <LOD   | <LOD  | <LOD  | NA      | 0.34                       |
| Pb      | <LOD   | <LOD   | <LOD  | <LOD  | NA      | 0.065                      |
| SAR     | 13.0   | 12.5   | 0.5   | 90.2  | 12.8    | NA                         |

Min: minimum value. Max: maximum value. Std Dev: Standard Deviation. ³: Limit of Detection. ⁴: USEPA 2014. National Recommended Water Quality Criteria (http://water.epa.gov/scitech/swguidance/standards/criteria/current/index.cfm#atable). §: Not Applicable.

Elemental concentrations measured on the separated liquid portion were also compared to the EPA Surface Water Quality Criteria for Aquatic Life. One to two elements in 11 samples exceeded this criterion. Eight samples exceeded the criteria for Zn and three for Cd. It is important to note that the
maximum concentration for Cd in this criterion was 2.0 µg L\(^{-1}\), which is less than the detection limit of the ICP-AES used for quantification. Based on the results of the metals measured in the liquid portion of the residuals, there is no risk of causing aquatic toxicity if the material was accidentally disposed of directly into a surface water body, except for Zn and Cd in a few cases. However, such toxicity is unlikely or negligible when the residual is used properly as a soil amendment.

3.3. Nutrients in the Solid Portion

Mehlich 3 extractions were performed on the separated solid portion to determine concentrations of P, K, Ca, and Mg, potentially available for plant uptake. The P, K, Ca, and Mg concentration from all samples widely ranged, respectively, from <LOD to 274.6, 9.2 to 658.4, 181.8 to 39320, and 12.7 to 1315 mg kg\(^{-1}\). When compared to the agronomic optimum, mean concentrations for K (156.5 mg kg\(^{-1}\)), Mg (438.8 mg kg\(^{-1}\)), and Ca (5600 mg kg\(^{-1}\)) were sufficient [19], however, P (23.1 mg kg\(^{-1}\)) would be slightly deficient if the residual was solely used as a medium for plant growth instead of as a soil amendment. A mean Ca level of 5660 mg kg\(^{-1}\) suggests that a residual could be a source of that nutrient among soils that are limited in Ca.

3.4. Metal Contents of the Separated Solids Portion of HDD Residuals

The solid portion of HDD residuals digested using EPA method 3050 and analyzed for trace metals including Mn, Ba, Cu, Zn, Cr, Ni, Pb, Co, As, and Cd were also submitted to descriptive statistics (Table 3). Metal concentrations were compared to EPA Part 503 (EPA 1994), which regulates the concentrations and loading of metals for land application of biosolids. The criteria for ‘Exceptional Quality’ (EQ) biosolids are the strictest standards in this regulation, so the land application of EQ biosolids requires no permit and can be applied based on the N and P contents of the material only. Metal concentrations in every residual were considerably less than the ceiling level for EQ biosolids (Table 3). However, it must be considered that the threshold metal EQ limits were determined based on expected bioavailability in the background matrix of organic matter. This is an important difference from the matrix of the residual samples, which are not rich in organic matter, and will, therefore, have an impact on the bioavailability of the metals. Likely, the metal bioavailability of the residual samples will highly vary as a function of the mineralogy of the material. This difference should be taken into consideration when calculating loading rates for land applying HDD residuals. If all residuals examined in this study were land applied at 112 Mg solid ha\(^{-1}\) (or 50 tons solid acre\(^{-1}\)) and compared to the EPA cumulative loading rates for biosolids, no samples would exceed the thresholds (Tables 4 and 5). One sample would exceed the EPA annual loading rate for Cu.

The concentrations of metals were also compared to the background concentrations found in Oklahoma benchmark soils [21]. Fifteen samples had some elements above these background concentrations, 12 of which were for Cu. Sample concentrations were also compared to typical levels found in soils throughout the United States [20]. Only seven samples were above these levels, all due to Cu (Table 3). When compared to the EPA Soil Screening Levels for residential soils (Table 3), As concentrations of two samples exceeded the maximum level but were far lower than the levels found in Oklahoma soils [21]. The modified SPLP was also performed on the solid portion of all samples (Table 6). This procedure is used as a means to estimate the leachability of the constituents in a material. Sample concentrations were compared to Maximum Contaminant Levels (MCLs) set by the EPA for drinking water. No samples exceeded the MCLs. Recommendation and implications on the use of the HDD residues are discussed next.
Table 3. Descriptive statistics for total elemental concentrations measured in solid residual samples, in mg kg\(^{-1}\). Values shown in comparison to typical levels found in typical soils of the US, benchmark soils in Oklahoma, EPA 503 ceiling concentrations for exceptional quality (EQ) biosolids, and EPA Soil Screening Levels for Residential Soils. \(n = 58\).

| Element | Mean  | Median | Min  | Max   | Std Dev | Typical Levels in Soil \(\dagger\) | Typical Levels in Oklahoma Soils \(\ddagger\) | EPA Ceiling Concentration for EQ Biosolids \(\S\) | EPA SSL for Residential Soils |
|---------|-------|--------|------|-------|---------|-----------------------------------|----------------------------------------|--------------------------------|--------------------------------|
| Ca      | 23,836 | 5751   | 192 | 223,986 | 40,310  | NA                                | NA                                     | NA                               | NA                           |
| Fe      | 16,972 | 14,362 | 570 | 51,185 | 11,299  | NA                                | NA                                     | NA                               | NA                           |
| Al      | 12,814 | 10,764 | 2454| 42,668 | 8960    | NA                                | NA                                     | NA                               | NA                           |
| Mg      | 5565   | 3641   | 102 | 34,074 | 7334    | NA                                | NA                                     | NA                               | NA                           |
| Na      | 2158   | 829    | 51  | 16,221 | 3314    | NA                                | NA                                     | NA                               | NA                           |
| K       | 1342   | 1088   | 85  | 6614   | 1209    | NA                                | NA                                     | NA                               | NA                           |
| S       | 971.8  | 549    | 19  | 6236   | 1187    | NA                                | NA                                     | NA                               | NA                           |
| P       | 460.6  | 287    | 30  | 5793   | 826.2   | NA                                | NA                                     | NA                               | NA                           |
| Mn      | 301    | 238    | <LOD| 1606   | 306     | 80–1300                           | 27–2165                               | NA                               | NA                           |
| Ba      | 105.9  | 57.6   | 9.2 | 974.1  | 142.9   | NA                                | NA                                     | NA                               | 5500                         |
| Cu      | 66.1   | 11     | <LOD| 1863   | 252.2   | 6–80                              | 2–33                                   | 4300                            | NA                           |
| Zn      | 39.9   | 30.5   | <LOD| 341    | 46.4    | 17–125                            | 15–142                                 | 7500                            | 23,000                       |
| B       | 37     | 31.2   | <LOD| 194.6  | 31.7    | NA                                | NA                                     | NA                               | NA                           |
| Cr      | 16.8   | 13.4   | 1.9 | 100.1  | 15.3    | 7–221                             | 4–70                                   | 3000                            | 230                          |
| Ni      | 11     | 8.5    | <LOD| 43.3   | 9.4     | 4–55                              | 2–57                                   | 420                             | 1600                         |
| Pb      | 2.4    | <LOD   | <LOD| 38.4   | 8.3     | 10–94                             | 3–32                                   | 840                             | NA                           |
| Co      | 2.3    | <LOD   | <LOD| 15.6   | 4.1     | 1–22                              | NA                                     | NA                               | NA                           |
| As      | <LOD   | <LOD   | <LOD| 1.6    | 0.2     | 4–9                               | 1–34                                   | 75                              | 0.4                          |
| Cd      | <LOD   | <LOD   | <LOD| <LOD   | 0.06–1.10 | 0.13–0.80                        | 85                                     | 70                              |                               |

Min: minimum value. Max: maximum value. Std Dev: Standard Deviation. \(\dagger\): [20]. \(\ddagger\): [21]. \(\S\): USEPA 1994. Land Application of Biosolids. \#\#: Below Limit of Detection. \(\dagger\dagger\): Not Applicable. (http://water.epa.gov/scitech/wastetech/biosolids/upload/2002_06_28_mlb_biosolids_503pe_503pe_2.pdf). USEPA 2002. GENERIC SSLs FOR THE RESIDENTIAL AND COMMERCIAL/INDUSTRIAL SCENARIOS (http://www.epa.gov/superfund/health/conmedia/soil/pdfs/ssg_appa-c.pdf).
Table 4. Calculated metal loading rates, in kg ha\(^{-1}\), for HDD residual samples tested in the study (1 to 27), assuming application rates of 112 metric tons solid ha\(^{-1}\). Metal loading rates are compared to EPA Cumulative and Annual Pollutant loading rates for biosolids.

| Sample | Zn  | Cu  | Ni  | As  | Cd  | Cr  | Pb  |
|--------|-----|-----|-----|-----|-----|-----|-----|
| 1      | 0.9 | 0.2 | 0.3 | <LOD| <LOD| 3.1 | <LOD|
| 2      | 8.2 | 53.6| 2.2 | <LOD| <LOD| 4.5 | <LOD|
| 3      | 3.7 | 1.1 | 1.1 | <LOD| <LOD| 1.5 | <LOD|
| 4      | 2.5 | 0.9 | 1.2 | <LOD| <LOD| 2   | 0.1 |
| 5      | 7.8 | 5.5 | 3.3 | <LOD| <LOD| 2.8 | 0.1 |
| 6      | 5.5 | 3.2 | 2.6 | <LOD| <LOD| 1.9 | <LOD|
| 7      | 2   | 1.3 | 0.9 | <LOD| <LOD| 1   | <LOD|
| 8      | 2.4 | 1.5 | 0.8 | <LOD| <LOD| 1.8 | 0.5 |
| 9      | 1.6 | 16.8| 0.9 | <LOD| <LOD| 3.8 | <LOD|
| 10     | 0.7 | 11.5| 0.4 | <LOD| <LOD| 2.9 | <LOD|
| 11     | 4.8 | 1.6 | 1.8 | <LOD| <LOD| 1.5 | 0.1 |
| 12     | 4.6 | 1.6 | 1.8 | <LOD| <LOD| 1.8 | <LOD|
| 13     | 1.8 | 2.3 | 0.7 | <LOD| <LOD| 0.7 | <LOD|
| 14     | 3.8 | 1.4 | 1.2 | <LOD| <LOD| 2.6 | <LOD|
| 15     | 7.1 | 2.6 | 3.2 | <LOD| <LOD| 2.7 | <LOD|
| 16     | 4.3 | 2.1 | 1.3 | <LOD| <LOD| 1.3 | <LOD|
| 17     | 1.3 | 0.1 | 0.3 | <LOD| <LOD| 1   | 0.9 |
| 18     | 1.4 | 0.9 | 0.3 | <LOD| <LOD| 0.9 | 0.4 |
| 19     | 2.3 | 0.9 | 1.3 | <LOD| <LOD| 1.2 | <LOD|
| 20     | 2.5 | 1   | 1   | <LOD| <LOD| 1   | <LOD|
| 21     | 8.4 | 49.7| 1.8 | <LOD| <LOD| 1.8 | 2.1 |
| 22     | 7.9 | 230.2| 1.7 | <LOD| <LOD| 1.9 | 2.1 |
| 23     | 8   | 4.2 | 0.3 | <LOD| <LOD| 0.9 | <LOD|
| 24     | 2.3 | 3.7 | 0.2 | <LOD| <LOD| 5.4 | <LOD|
| 25     | 3.6 | 2.7 | 1.3 | <LOD| <LOD| 1.4 | <LOD|
| 26     | 42.2| 4.7 | 2.4 | 0.2 | <LOD| 2.2 | 7.2 |
| 27     | 3.8 | <LOD| 2.2 | <LOD| <LOD| 12.4| <LOD|

EPA Annual Pollutant Loading Rate § 140 75 21 2 1.9 150 15
EPA Cumulative Pollutant Loading Rate § 2800 1500 420 41 39 3000 300

†: Below Limit of Detection. §: USEPA 1994. Land Application of Biosolids. USEPA (http://water.epa.gov/scitech/wastetech/biosolids/upload/2002_06_28_mtb_biosolids_503pe_503pe_2.pdf).

Table 5. Calculated metal loading rates, in kg ha\(^{-1}\), for HDD residual samples tested in the study (28 to 58), assuming application rates of 112 metric tons solid ha\(^{-1}\). Metal loading rates are compared to EPA Cumulative and Annual Pollutant loading rates for biosolids.

| Sample | Zn  | Cu  | Ni  | As  | Cd  | Cr  | Pb  |
|--------|-----|-----|-----|-----|-----|-----|-----|
| 28     | <LOD| <LOD| 0.1 | <LOD| <LOD| 0.7 | 0.3 |
| 29     | 4   | 4   | 1.6 | <LOD| <LOD| 2.3 | <LOD|
| 30     | 1.9 | 1.2 | 0.9 | <LOD| <LOD| 2.3 | <LOD|
| 31     | 1.4 | 0.2 | 0.5 | <LOD| <LOD| 0.7 | <LOD|
| 32     | 8.6 | 1.2 | 0.8 | <LOD| <LOD| 1.1 | <LOD|
| 33     | 1.6 | 1.3 | 0.5 | <LOD| <LOD| 0.5 | 0.5 |
| 34     | 12.3| 4.1 | 0.6 | <LOD| <LOD| 0.7 | 1.1 |
| 35     | 0.6 | 0.1 | 0.3 | <LOD| <LOD| 0.6 | 0.2 |
| 36     | 0.6 | 3.1 | 0.3 | <LOD| <LOD| 0.8 | 0.9 |
| 37     | 6.8 | 4.8 | 5.3 | <LOD| <LOD| 6.1 | <LOD|
| 38     | 5.7 | 1.4 | 1.3 | <LOD| <LOD| 2   | <LOD|
| 39     | 4.3 | 1.2 | 2.4 | <LOD| <LOD| 1.8 | 0.8 |
| 40     | 8.1 | 3   | 3.6 | <LOD| <LOD| 3.5 | <LOD|
| 41     | 10.6| 6.9 | 4.7 | <LOD| <LOD| 4.6 | <LOD|
| 42     | 2.1 | 0.7 | 0.3 | <LOD| <LOD| 0.4 | <LOD|
| 43     | 5.9 | 1.4 | 2.2 | <LOD| <LOD| 2.4 | <LOD|
| 44     | 2.1 | 1   | 0.3 | <LOD| <LOD| 0.9 | <LOD|
| 45     | 6.7 | 0.8 | 1.8 | <LOD| <LOD| 0.8 | <LOD|
Table 5. Cont.

| Sample | Zn | Cu | Ni | As | Cd | Cr | Pb |
|--------|----|----|----|----|----|----|----|
| 46     | 4.7 | 0.3 | 0.5 | <LOD | <LOD | 0.2 | <LOD |
| 47     | 2.9 | 1.3 | 0.5 | <LOD | <LOD | 0.5 | <LOD |
| 48     | 4.5 | 0.4 | 0.9 | <LOD | <LOD | 1   | <LOD |
| 49     | 3.1 | 0.1 | 2.8 | <LOD | <LOD | 3.6 | <LOD |
| 50     | 1.7 | 0.8 | 0.6 | <LOD | <LOD | 1.1 | <LOD |
| 51     | 2.5 | 1   | 0.4 | <LOD | <LOD | 3   | <LOD |
| 52     | 3.3 | 4.6 | 2.1 | <LOD | <LOD | 1.6 | <LOD |
| 53     | 3.6 | 1.4 | 1.7 | <LOD | <LOD | 1.3 | <LOD |
| 54     | 2.7 | 0.3 | 0.4 | <LOD | <LOD | 0.5 | <LOD |
| 55     | 4.9 | 2   | 2.2 | <LOD | <LOD | 2.3 | <LOD |
| 56     | 5.3 | 2.8 | 2.9 | <LOD | <LOD | 1.5 | <LOD |
| 57     | 7.1 | 9.9 | <LOD | <LOD | <LOD | 1.7 | <LOD |
| 58     | 11.4 | 6.8 | <LOD | <LOD | <LOD | 4.3 | <LOD |

EPA Annual Pollutant Loading Rate § 140 75 21 2 1.9 150 15
EPA Cumulative Pollutant Loading Rate § 2800 1500 420 41 3000 300

†: Below Limit of Detection. §: USEPA 1994. Land Application of Biosolids. USEPA (http://water.epa.gov/scitech/wastech/biosolids/upload/2002_06_28_mtb_biosolids_503pe_503pe_2.pdf).

Table 6. Descriptive statistics for elemental concentrations, in mg L\(^{-1}\), for analytes from the modified synthetic precipitation leaching procedure (SPLP) performed on the separated solid portion of HDD residuals. Values compared to maximum levels allowed by EPA Drinking Water Criteria. \(n = 61\).

| Element | Mean | Median | Min | Max | Std Dev | EPA Drinking Water Criteria |
|---------|------|--------|-----|-----|---------|-----------------------------|
| S       | 75.3 | 30.3   | 3.4 | 527.7 | 98.6 | NA |
| Na      | 63.5 | 38.4   | 1.9 | 386.9 | 74.5 | NA |
| Al      | 38.7 | 21.6   | <LOD | 204.1 | 51.0 | NA |
| Fe      | 15.4 | 9.0    | <LOD | 105.0 | 20.6 | NA |
| Ca      | 12.7 | 7.4    | 0.1 | 239.6 | 30.3 | NA |
| Mg      | 6.3  | 4.1    | <LOD | 42.1  | 8.4  | NA |
| K       | 4.1  | 3.6    | <LOD | 21.5  | 5.1  | NA |
| B       | 0.6  | 0.7    | 0.1 | 1.6   | 0.4  | NA |
| P       | 0.3  | 0.1    | <LOD | 5.5   | 0.8  | NA |
| Mn      | 0.1  | <LOD   | <LOD | 2.3   | 0.3  | NA |
| Ba      | 0.1  | 0.1    | <LOD | 0.3   | 0.1  | 2.0 |
| Cu      | <LOD | <LOD   | <LOD | 2.1   | 0.3  | 1.5 # |
| Zn      | <LOD | <LOD   | <LOD | 0.2   | 0.1  | NA |
| Cr      | <LOD | <LOD   | <LOD | 0.1   | NA\(^\d\) | 0.1 |
| Ni      | <LOD | <LOD   | <LOD | 0.2   | NA   | NA |
| Mo      | <LOD | <LOD   | <LOD | 0.1   | NA   | NA |
| Co      | <LOD | <LOD   | <LOD | 0.1   | NA   | NA |
| As      | <LOD | <LOD   | <LOD | NA    | 0.01 | NA |
| Cd      | <LOD | <LOD   | <LOD | NA    | 0.005 | NA |
| Pb      | <LOD | <LOD   | <LOD | NA    | <LOD | NA |

Min: minimum value. Max: maximum value. Std Dev: Standard Deviation. †: USEPA 2014. Drinking Water Contaminants. (https://www.epa.gov/sdwa/drinking-water-regulations-and-contaminants). ‡: Below Limit of Detection. §: Not Applicable. #: At Tap.

4. Discussion

There are typically two criteria used to assess a residual’s “suitability” for land application: (i) the by-product’s ability to enhance, or have no negative impact, on the productivity of the soil and plant growth, and (ii) a lack of negative environmental impacts [22]. Before land applying HDD residuals, one must consider the in-situ soil or geologic formation being bored through, from which the residuals originated. If boring through a site with known contamination, performing analysis for contaminants, such as heavy metals and salinity, is suggested. This could potentially be a problem when boring through the ground with a historic industrial use. For example, during the industrial revolution, industrial processes such as Cd plating and Cu smelting were commonly conducted outside of the major metropolitan areas. These areas are now referred to as suburbs. This is especially true for areas...
along the east coast. If boring takes place in an area where Cu smelting once occurred, drilling residuals would potentially contain high concentrations of Cu. A quick metal detection technique should be developed for onsite screening before deciding how HDD should be disposed of.

The EPA Part 503 criteria for metals concerning the land application of biosolids can be used as a guide for maximum allowable metals in an HDD residual to be land applied. One key difference between biosolids and HDD residuals is the organic matter content. Because of HDD residuals having low organic content, the bioavailability of metals is something to take into consideration. Because bioavailability differs with plant species [23], having an estimate of free metal concentration may be more useful. Separation by diffusion using the Donnan membrane technique is one method suggested to measure inorganic contaminants, or free metal concentrations, available for plant uptake [24]. Other methods include using the chelating agents’ diethylenetriaminepentaacetic acid (DTPA) or ethylenediaminetetraacetic acid (EDTA) to extract potentially bioavailable metals [25]. A study in China used the rhizosphere method, which uses low molecular weight organic acids to extract bioavailable metals, and compared results to the DTPA and EDTA methods. Their results showed the rhizosphere method to be the most robust in predicting the bioavailability of Cr, Cu, Zn, and Cd in soils to barley [26]. However, this study showed that the metals contained in the residuals were poorly soluble in water, thus potentially indicating that bioavailability and mobility would also be minimal.

A residual with a high SAR and/or salt concentration could be a problem, especially if the receiving soil is borderline saline and/or sodic. Unlike oil and gas drilling mud, HDD residuals are not going to contain geologic salts or return water from deep formations. However, this does not mean that HDD residuals do not have the potential to have a negative impact. Measuring the EC of an HDD residual before it is land applied would be beneficial in determining whether it could harm the soil that is receiving it. If the SAR of a residual is known to be above 13, incorporating the residual is recommended.

It is not recommended to use HDD residuals solely as a planting medium, although nothing was chemically limiting among the residual samples included in this study that would limit its use as a soil amendment. Total solids were found to be the most limiting factor for land application. Because of the residuals’ high clay content, applying too much of an HDD residual could smother the receiving soil and plants, resulting in poor plant growth and minimal water infiltration. If applied at a proper rate, the high clay content could improve sandy soil’s ability to retain water and nutrients. To minimize environmental impact to the receiving soil and plant life, acceptable HDD residuals (i.e., low metals and salt concentrations) should be applied at a rate ≤ of 112 metric tons solid ha⁻¹ on the ground with a slope ≤8% to minimize erosion and loss to surface water. Current regulations for oil and gas drilling residuals in Oklahoma allow for nearly 225 metric tons of solids ha⁻¹.

5. Conclusions

The physicochemical characterization of liquid and solid HDD residuals was performed on several samples from a broad geographic area in the US and showed that the total solids content was the most limiting factor for HDD residuals application. Despite that, land application proved itself a viable option for HDD residuals disposal, which encourages further studies investigating the impacts of field application of HDD on soils and plants. Faster and non-destructive methodologies for measuring elements in HDD residues, such as metals, are necessary to quickly qualify the materials for land application in the field. Such a technique could be explored with sensor technology that would eliminate the need to digest samples from the HDD residues in the laboratory.

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