Relationship between organic matter humification and bioavailability of sludge-borne copper and cadmium during long-term sludge amendment to soil

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HIGHLIGHTS

- Organic matter in sludge degraded with time goes after sludge was recycled to soil.
- DP in sludge is well coupled with incremental uptaken fraction of its borne copper.
- Profiles of Cadmium fractions in sludge exhibit an independent manner.

GRAPHICAL ABSTRACT

ABSTRACT

Recycling of sludge as soil amendment poses certain risk of heavy metals contamination. This study investigated the relationship between organic matter in composted sludge and its heavy metals bioavailability over 7 years. Periodic monitoring indicated a gradual increase in organic matter degradation, accompanied by changing degrees of polymerization, i.e., ratio of humic acid (HA)/fulvic acid (FA) coupled with incremental exchangeable fraction of copper (Cu) in sludge, with a growing rate of 74.7%, rather than that in soil. However, cadmium (Cd) in composted sludge exhibited an independent manner. Linear-regression analysis revealed that the total proportion of the Cu active fraction (exchangeable plus carbonate bound) was better correlated with the degree of polymerization (DP) and humification ratio (HR) than the degradation ratio of organic matter. Overall, amount of uptaken Cu was more dependent on the humification degree of organic matter, especially the proportion of HA in humus.

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1. Introduction

There are currently many ways to reuse sewage sludge, including co-incineration with cement, anaerobic digestion to produce methane, and carbonization to enable use in building materials. Among available treatment and disposal methods, application to soil as an amendment or fertilizer is regarded as a good option because it uses little energy and results in only minor secondary pollution of the environment (Liu and Zhang, 2013; Yoshida, et al., 2013; Mills, et al., 2014). Phyto-nutrients in sewage sludge, such as nitrogen, phosphorus and organic matter, can be adequately recycled and valorized. However, this method results in introduction of contaminants to the soil, such as organic contaminants and heavy metals. Although organic contaminants gradually degrade to some extent when sludge is applied to soil (Butler, et al., 2012; Rhind, et al., 2013; Stanczyk-Mazanek, et al., 2009; Wu, et al., 2011), levels of heavy metals in sludge do not decrease after soil application. Indeed, heavy metals are the most important environmental concern associated with the application of sludge to soil. Therefore, standards have been set to limit the amount of heavy metals in sludge that is applied to soil. However, current standards limit total heavy metals rather than the active fraction (General Administration of Quality Supervision, Inspection and Quarantine of China, 2009; Ministry of Housing and Urban-Rural Development of China, 2009; 2011; United States Environmental Protection Agency, 1993), especially the forms characterized as bioactivity of heavy metal, which can objectively reflect the its amounts from sludge to edible tissues uptaken by crop when sludge was reused as an amendment. In other words, the abundance of total heavy metals cannot account for the potential caused by the proportion of mobile heavy metals in sludge.

From 2007 to 2014, many studies have reported the heavy metals risk associated with the amendment of sludge to soil (Alcantara, et al., 2009; Antoniadis, et al., 2007; Brazauskiene, et al., 2008; Carbonell, et al., 2009; Lag-Brotets, et al., 2014; Oladeji, et al., 2012; Veganeh, et al., 2010). However, there is certain boundedness on the conclusions obtained by these researchers mentioned above. For example, (i) the risk has only been assessed by monitoring the results of heavy metals at one point in time after application of sludge to soil rather than consecutive detection. (ii) There is consensus regarding the relationship between potential risk and heavy metals bioavailability, but rarely involving organic matter. (iii) The relationship was investigated; however, the results were due to the simulating and laboratory level, in other words, in the absence of the impact by natural process and the support by the data obtained in field condition.

This study elaborated the connection between changes in organic matter transformation and heavy metal bioavailability in sludge that was applied to soil as an amendment over 7 years based on experimental results under field conditions. Significant differences were found in the shift of Cu and Cd from the inactive fraction to the active fraction in sludge, and the results implied that Cu is more susceptible to the transformation between organic matter and humus.

2. Materials and methods

Sludge used in this study was collected from a municipal wastewater treatment plant located in Qinhuangdao (Hebei Province, China), then mixed with wheat straw (bulking agent) at a ratio of 5:4 by volume. The resulting mixture was composted in a static forced aeration device with a capacity of 180 m³. The sludge was then transported by road from Qinhuangdao to Beijing. The physicochemical properties of the sludge compost are shown in Table 1. The experiment was performed for 7 consecutive years (2008–2014) at an experimental site located on the north side of Baima Road in Shunyi District, Beijing, which is a sod base of the Chinese Seed company. First, the soil was turned over twice, after which the sludge was sprayed onto the soil at depths of 8–10 cm at an amended dose of 90 Mg·ha⁻¹ in early October 2008. The experimental region was designed and divided into two plots (200 m x 20 m every plot), a treated one and another un-amended plot as a comparative background. A nylon mesh fabric (50 mesh·ft⁻²) was employed to keep unmixed and amended soil and sludge relatively separate without impacting the interaction between soil and sludge. Periodic sampling was conducted in mid-October every year, and each plot was sampled repeatedly six times. Following sampling, the location was marked with a flag to enable it to be resampled. All collected soil and sludge samples were brought back to the laboratory for indicator analysis.

All samples were dried and passed through a 2-mm sieve. Ash content was determined by combustion at 550 °C for 8 h. Organic matter was determined by subtraction of ash content, and total organic carbon was calculated as 58% of the organic matter (Zmora-Nahum, et al., 2005). Total extractable carbon (TEC) content was obtained by extraction from the sample with 0.1 mol·L⁻¹ of Na₂P₂O₇ (pH 9.7) using a sample: extraction ratio of 1:10 and subsequent colorimetric determination (Sims and Haby, 1971). An aliquot of the extract was acidified with HCl to pH 1.0, left to stand for 24 h in a refrigerator for the complete precipitation of HA, and then centrifuged. FA content was determined by measuring the C content in the supernatant in the same way as explained above for TEC. HA content was calculated by subtracting the C content in the acidified supernatant from the TEC content. The HR percentage of HA and DP was calculated from these parameters as (TEC/TOC) x 100, HA/TOC x 100 and HA/FA, respectively (Senesi, 1989).

The Cu and Cd fractions were analyzed by the Tessier Sequential Extraction method to determine the fractions in soil and sludge samples (Tessier, et al., 1979; Zheng, et al., 2007). Five solutions were generated successively: exchangeable (1 mol·L⁻¹ MgCl₂, pH 7); carbonated (1 mol·L⁻¹ NaOAc/HOAc, pH 5); Fe—Mn oxide bound (0.04 mol·L⁻¹ NH₄OH-HCl in 25% HOAc); organically bound (0.02 mol·L⁻¹ HNO₃ in 50% HNO₃; pH 2); 3.2 mol·L⁻¹ NH₄OAc in 20% HNO₃; residual (digested with concentrated HNO₃ + HClO₄). To perform quality control, the samples of certified standard reference materials were simultaneously digested for soils and sludge (GSS-1) from the China National Standard Materials Center with the experimental samples. The concentrations of Cu and Cd were determined by flame atomic absorption spectrophotometry (ContraAA700, Jena, Germany). All figures were developed using OriginPro 8.0, and all data were analyzed using SPSS v.13.0. Linear regression, based on p-value (p < 0.05), was used to simulate the correlations between organic matter degradation, HR, DP, and Cu active fraction to explain their characteristics in sludge.

3. Results

3.1. Tendency for organic matter degradation in soil and sludge

Fig. 1 (A) and (B) display the organic matter and its degradation profile in soil over seven years, respectively. There was no significant fluctuation in organic matter content in soil from the beginning of sludge amendment to the end. Correspondingly, the degradation ratio of organic matter also exhibited no abrupt change, with a maximum increase of only 5.9%. However, as shown in Fig. 1 (C) and (D), the content of

| Indicator                  | Soil   | Sludge |
|----------------------------|--------|--------|
| Moisture (%)               | 21     | 44     |
| Organic matter content (%) | 18.1   | 17.5   |
| Cu content (mg·kg⁻¹)       | 785.4  | 1345.6 |
| Cd content (mg·kg⁻¹)       | 1.3    | 2.9    |
| Total extractable carbon (g·kg⁻¹) | 28.6  | 61.1   |
| Humic acid carbon (g·kg⁻¹) | 1.2    | 26.4   |
| Fulvic acid carbon (g·kg⁻¹) | 0.9   | 22.7   |
organic matter in sludge decreased gradually with time from 2008 to 2014, while the change in the degradation ratio of organic matter in sludge showed the inverse pattern. As shown in Fig. 1(D), the degradation ratio increased to 3.6% in the second year, after which it was gradually elevated from 19.4% to 35.3%, indicating that the majority of organic matter in sludge decomposed with the aid of soil enzymes and functional microorganisms.

3.2. Change in humification ratio (HR) and degree of polymerization (DP) in soil and sludge

After sludge was amended to the soil, the HR in soil showed no obvious change with time, with an initial value of 18.25% and a final value of 20.42% being observed (Fig. 2(A)). Conversely, the HR in sludge presented an increasing appearance from 2008 to 2014, with the maximum ratio reaching 28.37% (Fig. 2(B)). These results are in accordance with those shown in Fig. 1, implying that most of degradable organic matter was converted to humus.

The degree of polymerization in soil and sludge was further analyzed. As shown in Fig. 3(A), a similar tendency of DP in soil was observed compared with HR, ranging from 1.35% to 1.67%. Additionally, there was an increasing trend in DP in sludge (Fig. 3(B)) approaching that of the HR, implying that the humus converted from the organic material all repolymerized to stabilized macromolecule organic compounds, which usually function as regulators of soil or biosolids improvement.

3.3. Change in bioavailability of Cu and Cd and their transformation among different fractions in soil and sludge

As shown in Fig. 4, the levels of the five Cu fractions (exchangeable, carbonate bound, Fe/Mn-oxide bound, organically bound, residual) in soil all remained constant during the studied period. However, the proportion of the Cu exchangeable faction in sludge increased from 16.2% to 28.3% with time accompanied by a gradual decline in the residual fraction. The total exchangeable plus carbonate bound fractions are usually regarded as the Cu active fraction, which can be taken up by plants. From this viewpoint, the level of the Cu active fraction increased with time.
The levels of exchangeable Cd in both soil and sludge showed similar unchangeable tendencies as Cu, although these changes seemed to be independent of time. These changes were clearly not associated with the changes in organic matter degradation, HR or DP. Moreover, the level of the Cd active fraction and residual fraction were also relatively stable from 2008 to 2014 (Fig. 5).

3.4. Linear-regression analyses between the proportion in Cu active fractions and organic matter relative indicators

Linear-regression analysis of the Cu active fraction and organic matter degradation revealed a determination coefficient of \( R^2 = 0.787 \) \((p < 0.05)\), while analysis of the Cu active fraction and HR revealed a determination coefficient of \( R^2 = 0.887 \) \((p < 0.05, \text{ as shown in Fig. 6 (B)})\). Surprisingly, the determination coefficient between the Cu active fraction and DP was up to \( R^2 = 0.91 \) \((p < 0.05, \text{ as shown in Fig. 6 (C)})\). These findings indicate that DP should reflect the level of active Cu more directly than organic matter degradation, even the HR.

4. Discussion

Previous reports showed that organic matter was involved in conversion of heavy metal fractions in sludge when it was added to soil for a relatively long duration. Accordingly, there are two theories regarding this regulation. One is known as the ‘time-bomb’ viewpoint, which indicates that the change in bioavailability of heavy metals in sludge accounted for the bondability of organic matter to different heavy metal fractions (Bergkvist et al., 2005; McBride, 2003; Stietiya and Wang, 2011). In this theory, the proportion of active fractions of heavy metal increases after sludge is recycled to soil as the sludge is gradually degraded. Another concept is the ‘Protect’ theory, in which the bioavailability of heavy metals is controlled by the inorganic fraction in sludge, which is adsorbed onto soil particles in an invariable ratio once sludge is added to soil as an organic fertilizer. Therefore, changes in the active form of heavy metal in sludge and its release potential to soil are independent of time (Frost and Ketchum, 2000; McBride, 2003).

It is generally believed that organic matter in sludge decreases with time after application to soil (Diagboya et al., 2015; Lu et al., 2014; Stietiya and Wang, 2011). This is a consequence of stabilization and humification (Fernández et al., 2007). In the presence of related soil enzymes, small molecules of organic matter are gradually transformed to complicated and stable macromolecular humus. Based on the results obtained in this study, the incremental change in the proportion of the Cu active fraction was positively associated with organic matter degradation, but not as closely as with HR, especially DP. In other words, a more accurate indicator can be employed to characterize the incremental change in Cu active fraction on the basis of the affirmative conclusion that uptake of Cu can be reflected by organic matter degradation (Mamindy-Pajany et al., 2014). Accordingly, the DP of sludge, which is responsible for the development of aggregate structure, also induces transformation of the Cu inactive fraction to the active fraction, to some extent, implying that the fraction of Cu is mainly regulated by sludge-borne micromolecular organic matters prior to its stabilization. Incremental HA, which functions as strong aggregation, weakens the affinity to Cu, accounting for the difference in competitive sorption caused by functional groups (Adani and Tambone, 2005; Oliver et al., 2005). In addition, high HA in sludge can be characterized by the lower acidic functional group content relative to soil (Pedra et al., 2008), further confirming that control of Cu was lost when sludge was mixed together with soil. Therefore, the degradation ratio of organic matter was the only superficial indicator that could be used to predict the proportion of the active fraction in the total amount of Cu, indicating that DP would be a more effective and accurate indicator. Sludge-borne organic matter stabilized gradually after application, and its amount in the
humified fraction increased. Consequently, the uptaken Cu, which is defined as the phyto-absorbed fraction, also presented an increasing tendency. This conclusion is referential and helpful to revise or fill the hole in the limit-value of heavy metal issued in the standard, which is responsible for controlling environmental risk from recycling of sludge to soil. Active fraction of some specific heavy metal should be taken to the consideration, particularly the organic matter profile, such as DP, also should be onto the list of emphasized parameters.

It is well-known that phenolic hydroxyl and carbonylic functional groups, which are responsible for the strong adsorption to Cd, are absent from sludge (Lee et al., 1996; Gray et al., 1999). Therefore, Cd mobility usually increases once sludge is amended to soil. It follows that possible environmental risks triggered by Cd should receive more attention.

The proportion of Cd in the active fraction was relatively steady and unchanged (Fig. 5). There is a consensus that the proportion of Cd in the active fraction remains constant during long-term reuse of composted sludge by application to soil (Bergkvist et al., 2005; Tapia et al., 2010), indicating that the pitch point of regulation for active Cd is not dependent on the change of humus in sludge-amended soil, at least in composted sludge. There are two possible reasons for why active Cd did not vary. One possibility is that transformation of Cd fraction occurs in accordance with ‘Protect’ theory, and is therefore determined by an inorganic ion (salt) functional group rather than the organic fraction. Considering this, active Cd may exist and behave in a fixed manner. The other possibility is that active Cd is dependent on the transformation from rapid degradable organic matter to humus, which occurs in and overlaps with a sludge composting period of no > 30 days. After sludge composting terminated, the amount of active Cd remained constant, and once composted sludge was sprayed to the soil, slowly degradable organic matter remained dominant, indicating that long-term humification is likely to last for several years. Nevertheless, transformation of Cd from the inactive fraction to the active fraction was deduced to be regulated by composting process. In sum, these results provide a theoretical framework of availability in sludge-borne Cu and Cd after application of sludge to soil (Fig. 7).
5. Conclusions

Previous studies have concluded that bioavailability of heavy metals is associated with organic matter profile. However, these conclusions were based on simulation experiments in the laboratory, and organic matter degradation occurred via chemical oxidation instead of natural processes. The present study presented field monitoring results, which showed that changes in the active fraction of Cu in sludge better correlated with DP level in sludge rather than organic matter HR and organic matter degradation. However, changes in the active fraction of Cd were independent of these indicators, remaining almost constant over the 7-year experimental period.

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