Study on Remediation of Hexachlorobenzene Contaminated Soil by Mechanochemical Method

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Abstract: The mechanochemical method is a potential way to destroy pollutants such as heavy metals and organic compounds due to its advantages such as complete reaction, adaptation of various pollutants and low energy consumption, etc. Research work was conducted to investigate the feasibility of remediating the persistent organic pollutants (POPs) contaminated soil and how the parameters influence the destruction of the pollutants. In the study, hexachlorobenzene (HCB) was used as a representative of the POPs in soil. Natural minerals such as albite and magnetite were selected as additives to treat HCB contaminated soil with the application of mechanochemical method. The reasonable operation parameters as well as the soil properties on the destruction of HCB were determined. Analysis such as Fourier Transform Infrared Spectroscopy (FTIR), X-ray diffractometer (XRD), X-ray Photoelectron Spectroscopy (XPS) and Raman Spectrometer were conducted for the supplement of mechanism study. A degradation rate of 92.5% for HCB was achieved under the optimal reaction condition. According to the XPS analysis results, the transformed valence state of iron, provided electrons for the destruction of HCB, on the basis of specific structure of albite. The amorphous carbon and graphite carbon were the final products of the destruction of HCB in the process of ball milling. The selected reagents with albite and magnetite would be viable for the damage of other POPs by mechanochemical method.

1 Introduction

A wide range of industrial products, resulting in complex pollution components, including a variety of heavy metals and organic pollutants, and the toxicity of persistent organic pollutants (POPs) to organisms has aroused wide concern [1]. Since the implementation of the Stockholm Convention, the manufacture and application of many kinds of POPs products have been restricted. However, the problem of soil pollution caused by product circulation, production leakage and abandoned factory field is very serious [2-4], then the research of soil remediation has become a hot spot. Moreover, the rapid remediation technologies of soil contaminated by organic pollutants cover extensively, including chemical reduction oxidation [5,6,7], soil leaching technology [8], photocatalytic degradation [9, 10,11], plasma photolysis and so on, with various requirements for the contaminated site. Physical methods, i.e., electro-kinetic remediation [9,12,13,14], soil vapor extraction [15,16], thermal desorption [17,18,19], supercritical fluid extraction [20], etc., also display certain efficiency. It is necessary to restore the nature of soil after remediation. Apart from the mentioned applications, the biological treatments also work [21,22], in spite of long period of remediation time. Therefore, to explore soil remediation technologies with high efficiency and environment protection has become a top priority.

With the investigation of mechanochemistry, researches of applying ball mill as a reactor to treat solid waste get flourished, for the superiorities it preserves, i.e., no-combustion, possible implementation process, less release of byproducts with toxicity [23,24], etc. The application of high efficient co-milling reagents can enhance the destruction reaction, shorten the repair time and realize the complete degradation of pollutants. Since Rowlands found that when CaO was used as the reagent, no original pollutants could be detected in the products of POPs, such as PCBs, chlorobenzene and DDTs by mechanical ball milling, and then CaO became the preferred co-milling reagent for mechanical and chemical degradation of POPs [25]. Many studies have been implemented with CaO as the main grinding assistant.

It was probed that, the PCBs contaminated soil was remediated by planetary ball mill with CaO and SiO₂ as reagents, and found that the total amount and toxicity equivalent of PCBs were reduced by 74% and 78% respectively [26]. Since then, the investigation of additives on organic compounds decomposition has gradually covered alkali metal oxides, neutral substances, oxidants and some reducing agents [27]. For example, it was proved that MgO, Al₂O₃ and La₂O₃ present great effect on the treatment of monochlorobiphenyl by ball milling [28]. As a neutral substance, CaC₂ shows great advantages in the process of degradation of hexachlorobenzene on mechanochemical method [24].

The existing researches on mechanochemical
treatment of POPs contaminated soil showed that exogenous additives applied present impact on soil texture, although they are efficient in the removal of pollutants. For example, the addition of CaO results in the increase of soil alkalinity, inhibits plant growth and has adverse effects on soil reclamation. Therefore, it is of great significance to explore the degradation of POPs contaminated soil by mechanochemistry with natural minerals which are similar to soil components and rich in source, to efficiently and economically remediate POPs contaminated field with full utilization of occupied field resources.

Albite belongs to feldspar group minerals, one of the sources of soil minerals with rich reserves and diverse compositions, such as Al₂O₃, SiO₂ and so on, which also has developed pore structure. And it is expected to realize the efficient degradation of pollutants by combing with magnetite, potential to provide internal electron transfer during the mechanochemistry process. In the study, natural mineral combination was used as co-milling reagents, with albite as main grinding agent and magnetite as grinding assistant, to study the mechanochemical effect on decomposition of hexachlorobenzene (HCB) in contaminated soil. The optimal parameters were determined, as well as the soil texture effect. And the mechanism of decomposition of HCB was studied with supplement of characterized methods.

2 Materials and method

2.1 Chemicals and instruments

Hexachlorobenzene (HCB, purity: 99.6%) was produced by Shenyang Research Institute of Chemical Industry (China), N-hexane (purity: pesticide analysis grade) was purchased from Nanjing Chemical Reagent Company Limited (China), Acetone (purity: pesticide analysis grade) was produced by Anhui Tiandi High Purity Solvent Company Limited (China), Albite (500 mesh) was produced by Yanxi mine (China), Fe₃O₄ (purity: 85%) was purchased from Aladdin Reagent Company limited (China), Montmorillonite (500 mesh) was purchased from Huzhou Chengke New Material Corporation (China), Quartz sand (20-40 mesh) was purchased from Shanghai Taitan Corporation.

A planetary ball mill (F-P400E, FOCUCY, China) with four zirconia pots was used for mechanochemical degradation experiments, An ultrasonic microwave collaborative reaction workstation (XO-SM100, ATPIO, China) was applied to extract target pollutants, A low speed centrifuge (TDZ5-WS, XIANGZHI, China) was employed to separate solid particles from liquids, A Nitrogen blowing concentrator (WD-12, Allsheng, China) was used to concentrate the substance to be tested, A gas chromatography mass spectrometry (A91 PLUS, PANNA, China) was applied to test target pollutants.

2.2 Materials

Fresh garden soil was collected according to the multi-point mixed sampling method, crushed and mixed by hand, and reduced to about 1kg by quartering method. Chemical analysis of soil samples showed that HCB was not detected. The properties of the soil samples such as total nitrogen (TN, g/kg), total phosphorus (TP, g/kg), total potassium (TK, g/kg), pH, organic matter (OM, %), electrical conductivity (EC, µs/cm) and cation exchange capacity (CEC, cmol(+)/kg), were detected and the results are shown in Table 1.

| P  | TN  | TK  | TP  | pH  | OM  | EC  | CEC |
|----|-----|-----|-----|-----|-----|-----|-----|
| C  | 1.63| 22.9| 1.15| 7.28| 3.09| 284 | 18  |

Where P is the abbreviation of parameters and C is the abbreviation of component contents.

2.3 Experiments

2.3.1 Preparation of simulated contaminated soil

The fresh soil sample was crushed and sieved through 100 mesh nylon sieve after air drying, and then put into an amber glass wide mouth packer. The quantitative HCB was dissolved in acetone to prepare HCB solution, fully stirred and mixed with the soil sample, and kept for one month. The above operations were carried out in the fume hood.

2.3.2 Ball milling experiment

Operational parameters of ball milling experiment, such as rotational speed (RS), material ratio (MS), ratio of different ball sizes (BDR), ratio of balls and material (R) and reaction time (t) as well as the soil texture, organic matter and moisture content were investigated on the effect of HCB destruction. For simplicity, the mass ratio of contaminated soil to additives is denoted as MS. The mass ratio of grinding balls with different diameters is denoted as BDR. The ratio of grinding ball to total materials is denoted as R.

The planetary ball milling was set at selected operation conditions according to experimental parameters. And the simulated soil sample, co-milling reagents and grinding balls were put into the grinding pot in a certain proportion. In order to ensure the stable operation of the machine, intermittent operation was set up, with a 15-min interval after every 1h.

2.3.3 Sample treatment and analysis

1g of grinding samples was placed in a beaker with 50mL of extracting agent (VAcetone/VN-hexane=1/1), the extraction was carried out with ultrasonic device (500W) for 15 min and repeated for 3 times. The collected solution was centrifuged at 3000 rpm for 20 min, and filtered with 0.45 µm organic filter membrane. The clarified solution was concentrated by nitrogen blowing and purified by magnesium silicate purification column, and finally
concentrated to 1mL.

The concentration of HCB in soil was analyzed by GC-MS. The heating procedure is as follows. The temperature was set at 120 °C holding for 2 min and then increased at 10 °C/min to 180 °C kept for 5 min. The temperature continued to increase at 8 °C/min to 240 °C holding for 1 min, and at 2 °C/min to 250 °C holding for 2 min; And then increased to 280 °C at 12 °C/min holding for 2 min. Temperature of the injection port was 250 °C with no shunt. The ion source temperature was 230 °C. The scanning mode was chosen as the selective ion scanning.

2.3.4 Characterization of grinding products

The instant changes of grinding products with different reaction time were characterized through X-ray diffractometer (XRD), Fourier Transform Infrared Spectroscopy (FTIR), X-ray Photoelectron Spectroscopy (XPS) and Raman Spectrometer. XRD analysis was conducted at a speed of 5°min-1 in the range of 2θ=10°-70° with Cu Kα radiation (Rigaku Ultima IV, Japan). FTIR analysis was carried out over the wave number ranging from 400cm-1 to 4000cm-1 with the KBr disk method (Thermo Scientific Nicolet 6700, US). XPS was conducted with Al Kα radiation at energy step size of 0.05eV and pass energy of 30eV (Thermo Scientific K-Alpha+, US), and the binding energy was calibrated by the surface contaminated C1s (284.8ev) as the standard. Raman was conducted at 514 nm from 50cm-1 to 2000cm-1 (HORIBA Scientific LabRAM HR Evolution, Japan).

3 Results and discussion

3.1 Effects of different operation parameters

Albite and magnetite were chosen as the co-milling reagents to improve the damage effect of HCB in soil through mechanochemical method. Different operation parameters were taken into consideration: rotational speed, material ratio, ball material ratio, ratio of different ball size and milling time.

Fig. 1 displayed that the residue of HCB in soil decreased with the increment of rotational speed. Low degradation rate was achieved at low rotational speed. Specifically, the degradation rate was 37.9% at a speed of 300rpm, and about 38.5% at 400rpm. The results reflect that the energy from collision and friction between the grinding ball and materials is limited at a low running speed, which does not favor the degradation of the organic pollutants.

![Figure 1. Effect of rotational speed on degradation rate of HCB in soil](image1)

As the rotational speed changed from 500rpm to 550rpm, the degradation rate was greatly enhanced to 72.5% and further increased to 79.3% when the milling speed was increased to 600rpm. Similar results from the literatures such as nearly 100% destruction of perfluorohexane sulfonate (PFHxS) with the reagents of ferrate(VI) and zero-valent iron (ZVI), as the rotational speed was at 600rpm for 4h [29] and less than 0.004% HCB left for 8h operation at 550 rpm [30] indicate that the energy input to reaction system was an important factor for reaction. Increasing rotational speed ensures increased collision frequency between the milling balls and reagents which would quickly impose on the destruction of pollutants and result in high degradation rate of HCB in soil.

![Figure 2. Effect of material ratio on degradation rate of HCB in soil](image2)
As the main energy input system in the experimental process, different BDR exerts an enormous function on the destruction of HCB. An in-depth study was conducted on the ball motion in planetary ball mill through the establishment of energy transfer model, and it was confirmed that the total effective impact energy and R are two important factors to determine the mechanochemical conversion rate of POPs [32].

Six sets of grinding ball ratio parameters were selected in the experiment, which were represented by the mass ratio of zirconia grinding balls of different sizes. Corresponding parameters conditions were shown in Table 2. Fig. 3 presents the optimal BDR to achieve relatively high HCB destruction rate when the size of grinding ball was medium and large that result in the well-distributed energy intensity during the operation.

| Parameter | Φ3mm | Φ5mm | Φ10mm | Φ15mm |
|-----------|------|------|--------|--------|
| BDR1      | 25g  | 50g  | 25g    | 0g     |
| BDR2      | 0g   | 0g   | 50g    | 50g    |
| BDR3      | 0g   | 0g   | 0g     | 100g   |
| BDR4      | 0g   | 0g   | 100g   | 0g     |
| BDR5      | 0g   | 33.3g| 33.3g  | 33.3g  |
| BDR6      | 0g   | 25g  | 50g    | 25g    |

Table 2. Combinations of different ball diameter ratio parameters

The worst destruction effect with BDR of 1, manifested that the desired effect of large grinding ball was better than that of small grinding ball. Due to the high quality of grinding balls, the force exerted on materials increased correspondingly, which accelerated the degradation of pollutants. What’s more, the grinding ball with large diameter can easily engender milling blind area during the reaction process, which meant that the combination of large and small grinding balls can ensure the applied density and reaction area of mechanical force at the same time. It was proved that different grinding ball size ratio could significantly affect the destruction of DDTs in soil with the reagent of CaO, and the effect of applying solely large ball was worst [33]. The ratio of grinding balls with different sizes impacts the collision density and frequency, which can be proved by that the results obtained under the same value of R.

The contact efficiency between materials and milling ball was affected by ball material ratio. An appropriate ratio could achieve the efficient damage of pollutants as well as improve the capacity of single contaminated soil treatment with the appropriate filling ensured.

It was indicated in Fig. 4 that the increasing tendency of HCB degradation rate was gentle with R. The damage rate of HCB was 81.1% as R was 30:1, and a rapid increase in degradation rate of HCB was achieved as R was changed from 15:1 to 20:1. When R was 10:1 and 15:1, the damage of HCB in soil was less than 70%. It was demonstrated that the destruction efficiency of HCB was significantly improved with the increase of R. The degradation rate of HCB increased from 26.7% to 86.3% as R increased from 10:1 to 20:1[34]. While R reaches a certain value, the effective counteraction force between grinding balls increases, and the energy generated by collision is beneficial to the structural damage of pollutants.

Milling time is directly associated with the extent of mechanochemical remediation of contaminated soil. The prolonged milling time would give rise to excessive power consumption and impose a burden to equipment operation and maintenance. In order to determine the suitable time for HCB contaminated soil treatment, six working conditions were set up, including 1h, 2h, 3h, 4h, 5h and 6h. With the extension of milling time, the removal rate of HCB in contaminated soil increased, as shown in Fig. 5. The destruction rate of HCB was rapidly enhanced from 43.8% to 87.67% as t was increased from 1h to 4h. As the milling time continued to extend, the destruction rate increased gently. When t was 6h, the damage rate of HCB could reach 92.52%. With the continuous accumulation of mechanical collision, the complex chemical reactions occurred between HCB and co-milling reagents. Considering the treatment effect and economic benefits, the suitable milling time would be 5h, while the destruction rate of HCB is more than 90%, a more economical and efficient choice, slightly lower than the destruction effect of 6h.
3.2 Effects of soil texture

The adhesion force between pollutants and soil was closely related to soil quality, and the interception of pollutants were relevant to the type of soil, which possesses different adsorption capacity. Li et al. probed into the relationship between soil properties and pollutants, such as volatile organic compounds (VOCs) and total petroleum hydrocarbons (TPH), at specific contaminated site. It was manifested that the TPH content was higher in the surface plain fill, while the VOCs was low. In the clay layer, a large amount of TPH and VOCs were adsorbed on the top of the soil layer [35]. Organic pollutants had a strong binding force with clay, which hindered the destruction of pollutants.

Different classifications of HCB contaminated soils, including sand, loam and clay, were selected for the study. Quartz sand, garden soil and montmorillonite powder were utilized as sand source, loam source and clay source respectively. The concentration of HCB in simulated contaminated soil was 50mg/kg.

It was shown in Fig. 6 that the degradation effect of HCB in contaminated sand was the best, in which the residue components was near non-detectable, followed by the contaminated loam, and the destruction extent of contaminated clay was the worst. The quartz sand was lack of organic matter and had the worst adsorption capacity to pollutants. The garden soil had suitable content of nutrient and organic matter, which was closely combined with pollutants and had little impact on pollutant destruction. The clay had small particle size and developed pores, which provided abundant retention sites for pollutants. Van der Waals force and hydrogen bond interaction made the organic and inorganic contaminants bounded to particles tightly [36]. Therefore, the clay was most closely combined with pollutants, resulting in poor ball milling effect, and its HCB degradation rate could only reach 31.4%.

Fig. 7 displayed that with the increment of humic acid in soil, the destruction of HCB in soil initially decreased, then followed by an increase. The degradation rate of HCB was 82.1% when humic acid addition was 15wt.%. As the humic acid addition was low, the binding ability of soil with HCB was higher than that of soil without humic acid addition, which resulted in the decreased degradation rate of HCB during ball milling. Li et al. clarified that the
adsorption capacity of bisphenol A in soil increased with the increment of soil organic matter content [37]. The contaminants accumulated due to the adsorption of humic acid as the increase of humic acid addition, which concentrated the pollutants in soil, and improved the degradation effect of HCB. The organic matter in soil in China is mostly 1wt.% - 5wt.%, and the damage rate of pollutants is less than 70% without exogenous organic matter. Therefore, reasonable application of exogenous nutrients can not only improve the soil nutritional status, but also improve the degradation efficiency of POPs contaminated soil.

Table 3. Distribution of organic matter content in soil samples

| Humic Acid Addition/ % | 2   | 5   | 10  | 15  |
|------------------------|-----|-----|-----|-----|
| Organic Matter/ %      | 4.98| 7.62| 9.46| 10.60|

The effect of moisture content in contaminated soil on the HCB degradation rate is an important factor to balance the pretreatment cost. Drying time of soil and treatment cost would be controlled effectively as the influence of moisture was determined. Four experimental groups of 0wt.%, 3wt.%, 5wt.%, 10wt.% were set up to explore the influence of moisture content on HCB degradation. The experiment results were shown in Fig. 8.

71.5% of HCB damage rate was achieved when the polluted soil was completely dried. Great decrease in degradation rate was seen as the soil moisture increased from 0% to 3%, which reflected the hindrance of water to destroy pollutants by mechanochemical method. With the moisture content further increased to 5wt.%, not much change in the destruction of HCB rate of 65%, while it decreased to less than 60% as the moisture increased to 10wt.%.

![Figure 8. Effect of moisture content on degradation rate of HCB in soil](image)

Figure 8. Effect of moisture content on degradation rate of HCB in soil
(t=4h; RS=550rpm; R=20:1; MR=10:1; BDR(Φ10mm:Φ15mm=1:1))

The planetary ball mill with zirconia pots and balls was used as the reaction system to destroy POPs in soil. The resistance of the ball movement raised with high content of moisture. Continuous hits led to the adhesion of soil to grinding balls and tank walls, resulted in uneven impact frequency and density of pollutants, which was not beneficial to the damage of HCB in soil. Thus, in order to achieve high degradation rate, it is necessary to extend the reaction time, which increase the power consumption and equipment operation burden. Therefore, the moisture content of polluted soil should be less than 5wt.% to ensure the degradation efficiency with low cost in the process of drying.

3.3 Characteristics of milling products

In order to clarify the mechanochemical effect of albite and Fe3O4 on HCB in soil, FTIR was conducted on the products of different milling time to identify the group transformation during ball milling [38].

As shown in Fig. 9, a sharp moderate intensity adsorption peak at 3617.8cm-1 was detected for the original samples (0h) which was caused by –OH stretching vibration. The absorption intensity decreased gradually with the increase of milling time, indicating that –OH has participated in the reaction. The peaks at 3021.9cm-1 and 2897.5cm-1 were =C–H stretching vibration and saturated –C–H antisymmetric stretching vibration, respectively and disappeared during the reaction.

![Figure 9. FTIR spectra of products at different reaction time](image)

Figure 9. FTIR spectra of products at different reaction time (RS=500rpm; R=20:1; MR=1:1; BDR(Φ10mm:Φ15mm=1:1))

The peak at 730.9cm-1 originated from the C–Cl stretching vibration of HCB molecule, and the absorption peak was weakened and disappeared after ball milling, indicating HCB molecule has been destroyed. The enhancement peak of 1868.7cm-1 was caused by phenoxy. The new peaks at 1494.6cm-1, 1159cm-1 and 883.2cm-1 were C=C stretching vibration in aromatic ring, stretching vibration of C–O and wagging vibration of =CH2, respectively. The new absorption peaks were mainly from the intermediate products generated after dechlorination of HCB [37,23,24].

The raw materials had three sharp Raman peaks at 177.1cm-1, 300.3cm-1 and 1097.9cm-1 as shown in Fig. 10. After 2h reaction, the three characteristic peaks became weaker and a new peak engendered at 1354.8cm-1. After 6h reaction, the peaks at 177.1cm-1 and 197.9cm-1 faded away, and a new peak at 1578.9cm-1 emerged, which indicated that the pollutants was damaged gradually and new products appeared. According to Raman characteristic peaks of carbon crystal, D band and G band, are located at approximate 1300cm-1 and 1580cm-1 respectively. D band represents lattice defect of carbon, reflecting the disorder of crystal structure, and G band represents the first-order scattering E2g vibration.
mode, which is applied to characterize the sp2 bond structure of carbon.

![Figure 10](image1.png)

**Figure 10.** Raman spectra of different milling products (RS=500rpm; R=20:1; MR=1:1; BDR(Φ10mm:Φ15mm=1:1))

Upon the previous studies, the peak at 1354.8cm⁻¹ belongs to the defect and disorder of carbon structure, specifically the amorphous carbon, and the response peak at 1578.9cm⁻¹ belongs to the graphite. Raman spectrum clearly showed that carbonization process occurred in the process of mechanochemical reaction [30,24,39].

![Figure 11](image2.png)

**Figure 11.** XRD patterns of the milling products at different reaction time

XRD was applied to verify the crystalline forms of milling mixtures. It was depicted in Fig.11 that the peak of HCB in soil samples were largely decreased after 6h milling, as a result of destruction caused by mechanical stress.

The peak intensity of other inorganic substances, i.e., Fe3O4 and Al2O3, was obviously weakened, and the peaks became broad and weak with the grinding aggravation. It was manifested that the crystallinity decreased and the ball milling mixture became amorphous. The changes happened in structure of mixtures, made the damage reaction of HCB in soil possible.

In order to clarify the effect of Fe3O4, the solid residues were exploited to detect the chemical environment of Fe 2p by XPS. As shown in Fig.12, for the raw mixture, two typical peaks were measured at 711.0eV and 724.5eV, which were attributed to Fe 2p3/2 and Fe 2p1/2, respectively. Peak differentiating and imitating analysis was detected at the peak of Fe 2p3/2, and two fitting peaks were achieved at 711.5eV and 710.4eV [40]. The former stood for Fe3⁺, while Fe2⁺ was represented by the latter. The ratio of peak area between Fe3⁺ and Fe2⁺ was about 2:1, which was consistent with that the theoretical electronic composition of Fe3O4. The results showed that the ratio of peak area between the two kinds of electron increased after 6h ball milling as shown in Fig. 13.

The following reactions must have occurred:

\[
\text{Fe}^{2+} - \text{e}^- = \text{Fe}^{3+} \quad (1)
\]

\[
\text{HCB} + \text{e}^- = \text{HCB}^- \quad (2)
\]

![Figure 12](image3.png)

**Figure 12.** XPS spectra (Fe 2p) of raw materials

![Figure 13](image4.png)

**Figure 13.** XPS spectra (Fe 2p) of milling mixture at 6h

The internal electron transformation between Fe2⁺ and Fe3⁺ in Fe3O4 brought in electrons for the reaction system, which promoted the generation of HCB anion, as shown in Equation 2 [23]. The process of receiving electron of HCB is an endothermic reaction, which requires certain amounts of energy to stimulate it. What’s more, the electronegativity of HCB- was strengthened by extra electrons in the molecule, which caused an increased mean length of C–Cl, about 0.02Å higher than that of HCB, and thus weakened the C–Cl [23]. A series of dissolved reactions would happen during the destruction of HCB by giving rise to stable chlorine.

### 4 Conclusion

The feasibility of destroy HCB in soil with the mechanochemistry method was studied. The parameters such as rotational speed (RS), material ratio (MS), milling time (t), ratio of ball size (BDR), classifications of soil, moisture and organic matter in soil, etc., were investigated
on the influence of HCB destruction rate in contaminated soil. It was manifested that degradation rate of HCB could reach 92.5% after 6h milling with the BDR of 2, RS of 550rpm, MS of 10:1 and R of 10:1, while the enough energy from collision and friction was achieved.

The classifications of soil and content of organic matter present the significant effect on HCB destruction rate in soil because of the difference in absorption from different organic components existing in soil, while the remediation of HCB contaminated soil with few organic matters was complete, compared with loam and clay. Soil moisture exerted a certain blocking effect on the mechanochemical reaction, and resulted in increased movement resistance for grinding balls, leading to an uneven collision density and frequency.

Natural mineral assemblage had significant effect on the damage of HCB in soil. The specific composition and structure of Albite combined with the input internal electron transformed from the magnetite during the reaction, were greatly beneficial to the destruction of POPs in contaminated soil. The work presents an environmental friendly and efficiently method for POPs contaminated soil remediation with few impact on soil texture.

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