Novel Approach for the Remediation of Radioactive Cesium Contaminated Soil with nano-Fe/Ca/CaO Dispersion Mixture in Dry Condition

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Abstract. Present study, first time we developed a nano-Fe/Ca/CaO dispersion mixture based remediation and volume reduction method of real radioactive cesium contaminated soils. After soil samples treated with 10wt% of nano-Fe/Ca/CaO dispersion mixtures, emitting radiation intensity was reduced from 4.00 μSv/h to 0.95 μSv/h in non-magnetic fraction soils. While, after treatment, about 30wt% magnetic and 70wt% non-magnetic fraction soils were separated, and its condensed radioactive cesium concentration was about 80% and 20%, respectively. By this way, cesium contaminated soil volume can be reduced. These preliminary results appear to be very promising and the simple mixing with the addition of nano-Fe/Ca/CaO may be considered potentially applicable for the remediation and separation of radioactive Cs contaminated soil in dry conditions.

Key words: Contaminated soil, Nano-Fe/Ca/CaO, Radioactive cesium, Remediation, Volume reduction.

Introduction

In Japan, the major concern on the radioactive cesium deposition and its soil contamination due to the emission from the Fukushima Daiichi Nuclear Power Plant showed up after a massive quake on March 11, 2011 (Nuclear and Industrial Safety Agency, 2011). The Japan government recent radiation map shows high levels of radioactive cesium in Fukushima and surrounding prefectures. Some spots have levels between 100,000 and 600,000 becquerels per square meter (Bq/m²) (148,000 was the standard used for mandatory resettlement after the Chernobyl disaster) (Patel, 2011). According to the Japan science ministry about 8 percent of the country’s land has been contaminated with Cs137 levels higher than 10,000 Bq/m², a threshold that Japan’s science ministry defines as affected by the nuclear accident (Ishizuka, 2011). The soil area around the nuclear power plant in Fukushima has more than 100,000 Bq/m² of Cs137. Miyagi and Ibaraki prefectures have levels higher than 25,000 Bq/m². Using a conversion factor of 53kg of soil per square meter, they calculate that soils in the eastern fraction of Fukushima prefecture exceed the government’s safe limit of 2,500 Bq/kg of Cs137 (Yasunari et al., 2011). Soil contamination with radioactive cesium has a long term radiological impact due to its long physical half-life (2 years for Cs134 and 30 years for Cs137) and its high biological availability (Staunton and Roubaud 1997). Consumption of agricultural produce contaminated with radio cesium represents the principal route of human exposure to this radionuclide. Therefore, the remediation and volume reduction of radioactive cesium contaminated soil is recognized to be one of the most important problems, which should be solved by taking advantage of suitable technologies. Many diverse methods are used such as solidification, embedding or encapsulation to immobilize the radioactive wastes in a solidified form. Immobilization technique consists of entrapping within a solid matrix i.e cement, cement-based material, polymer or ceramic (Shi and Day, 1996; Plecas et al., 2003). However, among the drawbacks of these processes, the modification of original soil properties, the progressive reduction of the efficiency due to the degradation and/or saturation of reactants, forming of secondary effluents and their treatment for additional cost. Therefore, the treatment under dry and water free conditions should be considered. We have recently shown that the nano-metallic Ca/CaO/NaH2PO4 desorption mixture is most effective for cesium immobilization (about 96%) in soil.
by ball milling treatment in dry condition (Srinivasa Reddy et al., 2012). It is well understood that the most effective measure for immediate relief from radioactive cesium is topsoil removal/separation. We assumed that with nano-Fe/Ca/CaO dispersion mixer by simple mixing/grinding, Ca/CaO can make immobile salts with moisture and CO$_2$ in atmosphere, including pozzolanic cement and hydraulic property, cesium would be brought into the immobile Ca salts. Therefore, the calcium carbonate (CaCO$_3$/calcium hydroxide Ca(OH)$_2$) layer in soil surface could be produced, hence the cesium on soil surface can enclosed/bided (Srinivasa Reddy et al., 2012). Further by magnetic separation, encapsulated nano-Fe particles in solidified soil can also be separated with magnetic, by this way radioactive cesium contaminated soil volume can be reduced. Therefore, in present study, we developed a nano-Fe/Ca/CaO dispersion mixture based solidification method to immobilization of radioactive cesium and its separation form real radioactive cesium contaminated soils in Fukushima, Japan.

Materials and Methods

Initially, the radioactive cesium contaminated surface soil samples were collected in two places in Fukushima prefecture, first one is 15 km far from nuclear power station and second one is about 80 km far from power station, and marked as F1 and F2, both soil samples are having 1% moisture content and, other physical-chemical properties are similar. While, nano-Fe/Ca/CaO was prepared with Fe/metallic Ca/CaO through mechanochemical process. Granular particles of metallic calcium were purchased from Kishida Chemical Co. Ltd. (99%, particle size distribution: 2–2.5 mm, surface area: 0.43–0.48 m$^2$ g$^{-1}$). Similarly, Fe powder (size 0.15 mm) was purchased from Kishida Chemical Co. Ltd. CaO was also commercially obtained. At room temperature under Ar, dry CaO (825°C for 2h) and dry Fe/metallic Ca composition (Fe/Ca/CaO=2/2.5) were introduced in planetary ball mill (20 pieces SUS, 32g/ball). Stirred at 600 rpm for 1h, samples collected in a glass bottles, Ar gas filled and used. Metallic Ca content to total weight was 2.8mmol/g mixture. Radioactive cesium contaminated soil samples were mixed along with dry 10wt% of nano-Fe/Ca/CaO (ratio: 1.00/0.11) at 150-200 rpm in a small ceramic tumbling mill (500mL size, without ceramic balls) at room temperature under air for 1 hr. After mixing, separated magnetic and non-magnetic fractions. The radioactive cesium (Cs$^{137}$ and Cs$^{134}$) concentrations in soil samples was measured with high purity germanium detector in gamma-ray spectrometry (SEIKO EG & G GMX-20P4-70) with standard method. While, radioactive cesium contaminated soils emitting radiation was measured with Geiger–Müller counter from upper 5 cm distance soil surface. In order to elucidate cesium immobilization/separation mechanisms and verify soil alterations during treatment with nano-Fe/Ca/CaO. Further, synthetically cesium (Cs$^{135}$) contaminated soil was prepared in laboratory, 0.2g cesium chloride (CsCl) was dissolved in 5 ml water, next, solutions was thoroughly mixed to 1kg of sandy soil having 7% moisture (non-polluted, commercially availablemica/isinglass/fibrolite soil type, produced in Okayama prefecture, Japan) and used. Scanning electron microscopy combined with electron dispersive spectroscopy (SEM-EDS; JEOL, JSM6510A equipped with a Si (Li) probe at resolution of 138 eV) microanalysis and semi-quantitative analyses was applied.

Results and Discussion

Initial measured emitting radiation in contaminated soil was about 4.00 µSv/h in both (F1 and F2) samples. While, radioactive cesium (Cs$^{134}$ and Cs$^{137}$) concentrations sum of magnetic and non-magnetic fraction soils was 290,970 and 413,440 Bq/kg in F1 sample, and about 293,170 and 420,600 Bq/kg in F2 sample respectively. After soil samples treated with 10wt% of nano-Fe/Ca/CaO dispersion mixtures, in magnetic fraction soil continued the high emitting value about 2.9 and 2 µSv/h, while decreased the value about 0.95 and 0.65 µSv/h in non-magnetic fraction in both soil (F1 and F2) samples. Correspondingly, the concentration of cesium (Cs$^{134}$/Cs$^{137}$) in magnetic fraction soil was reduced, while magnetic fraction soil increased (Fig. 1). After treatment with nano-Fe/Ca/CaO, about 30wt% magnetic and 70wt% of non-magnetic fraction soils was separated by magnetic, and the radioactive cesium concentration was condensed (about 80%) in this magnetic fraction soil, while only 20% remained in non-magnetic fraction soil (Fig. 1). The SEM images for syntactically prepared Cs contaminated soil before and after grinding are shown in Fig. 2. After blending the Cs contaminated syntactic soil with nano-Fe/Ca/CaO decreased the production and development of the core fractionicles and would be caused the hydration products to coat the aggregates extremely, as shown in Fig. 2b. By SEM EDS maps (Fig. 3) and semi- quantitative analysis, it is possible to observe that the amount of Cs mass and atom percent detectable on soil fractionicle surface increases after nano-Fe/Ca/CaO treatment. Decreased Cs mass percent form 0.15 to 0.04 in magnetic fraction, while non-magnetic fraction, 0 % of Cs was observed. The reduction of Cs amount on fractionicle surface is a possible explanation of the increase of immobilization. However, in magnetic fraction slight concentration was observed. Further, schematic representation of possible mechanisms, which may determine the immobilization and separation of radioactive cesium contaminated soils after simple mixing with nano-Fe/Ca/CaO dispersion mixture in dry conditions was shown in Fig. 4. It is possible to assume that when soil is contaminated, Cs is adsorbed onto soil fractionicles through a surface coordination process. When mixing starts, soil fractionicles are subjected to collisions that may promote aggregation in presence of nano-Fe/Ca/CaO. This fact probably indicates that a certain amount of Cs entrapped inside new aggregates during the nano-Fe/Ca/CaO treatment (Fig. 4a). On the other hand, we also assumed that reduction of Cs would be another possible mechanism in case of nano-Ca addition. The standard redox potential of Cs$^{+}$ (~3.026 V, 25 °C) is very
close to that of Ca$^{2+}$ (−2.868 V, 25°C), and thus, the removal of Cs$^+$ ions by nano Ca is due to sorption/replacement with Ca$^{2+}$. Cs either reduced at the nano-Fe/Ca/CaO surface or directly adsorbed to the surface where they are rendered immobile. On the other hand, nano-Fe/Ca/PO$_4$ on the surface of soil can enclosed/bind with the Cs in the presence of moisture due to its electron sources and the CaCO$_3$/Ca(OH)$_2$ layer in soil surface could be produced with them (Fig. 4a). Further by magnetic separation, encapsulated nano-Fe fractionicles in solidified soil can also be separated, by this way contaminated soil volume can be reduced (Fig. 4b).

Conclusions

Present study, first time we developed a nano-Fe/Ca/CaO dispersion mixture based remediation and volume reduction method of real radioactive cesium contaminated soils in Fukushima. After soil samples
treated with 10wt% of nano-Fe/Ca/CaO dispersion mixtures, in non-magnetic fraction soil emitting radiation intensity was reduced from 4 µSv/h to 0.95 µSv/h. While, after treatment, about 30wt% magnetic and 70wt% of non-magnetic fraction soils was separated, and the condensed radioactive cesium concentration was about 80% and 20% in magnetic and non-magnetic fraction soils respectively. By this way, cesium contaminated soil volume can be reduced. Thus, we anticipated, the simple mixing technique with the addition of nano-Fe/Ca/CaO may be considered potentially applicable for the remediation and separation of radioactive cesium contaminated soil in dry conditions.

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References


