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Highlights

- Alkanolamines promoted the hydration of steel slag and with TIPA showing the best performance.
- The UCS of treated HM-contaminated soil at 28 days was more than tripled using 0.1% TIPA-activated SS compared to the non-activated SS.
- TCLP leached concentrations of Cd, Cu, Ni, Pb, and Zn were reduced by 87.2%, 78.8%, 62.4%, 73.6% and 64.5% using 0.1% TIPA-activated SS at 28 days.
- Alkanolamines-activated SS is a sustainable alternative to PC in S/S for heavily-contaminated soil

Supplementary Material

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Credit Author Statement

Bin Zeng: Conceptualization, Experiment, Writing - review & editing, Supervision, Project administration, Investigation.

Zhi Zhang: Methodology, Investigation, Data analysis, Writing - review & editing.

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Fei Jin: Resources, Writing - review & editing, Supervision, Project administration, Investigation.

Declaration of Interest Statement

The authors declared that they have no conflicts of interest to this work. We declare that we do not have any commercial or associative interest that represents a conflict of interest in connection with the work submitted.

1	Alkanolamines-activated steel slag for
2	stabilization/solidification of heavy metal contaminated soil
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11 12 13	 Highlights: Alkanolamines promoted the hydration of steel slag and with TIPA showing the best performance.
14 15	• The UCS of treated HM-contaminated soil at 28 days was more than tripled using 0.1% TIPA- activated SS compared to the non-activated SS.
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20 21 22 23	Abbreviations:SS: Steel slagS/S: Stabilization/SolidificationTEA: TriethanolamineTIPA: TriisopropanolamineDEIPA: DiethanolisopropanolamineUCS: Unconfine strength
24	β -C ₂ S: β-larnite CaCO ₃ : Calcite C ₂ F: Srebrodolskite C ₁₂ A ₇ : Mayenite
25 26 27 28 29	C3A: Tricalcium aluminateC-S-H: Calcium silicate hydrateMc: MonocarboaluminateCH: PortlanditeC-A-H: Calcium aluminate hydrateC-F-H: Calcium ferrite hydrateIC: Isothermal calorimetryXRD: X-ray diffractionTGA: Thermogravimetric analysisTCLP: Toxicity Characteristic Leaching Procedure
30	Abstract: Steel slag (SS) is a byproduct discharged from steel-making industry with less than 25%
31	utilization rate in China. The low utilisation rate of SS is associated with its low hydration activity
32	in cement and concrete. In this study, four different alkanolamines (TEA, TIPA, EDIPA and DEIPA)
33	were used to activate SS to improve its cementitious properties and metal binding performance, and
34	hence its capacity on treating heavy metal-contaminated soils containing Cd, Cu, Ni, Pb and Zn.

35 Compared with the reference SS without activators, concentrations of leached Cd, Cu, Ni, Pb, and 36 Zn have reduced by 87.2%, 78.8%, 62.4%, 73.6% and 64.5% by using 0.1% TIPA-activated SS 37 after 28 days, and they were all below their respective regulatory limits by Standard for Pollution 38 Control on the Hazardous Waste Landfill (GB 18598-2019) in China, and the unconfined 39 compressive strength (UCS) of the treated soil at 28 days was enhanced by 237.7% using 0.1% 40 TIPA-activated SS. To elucidate the activation mechanism, the hydration process of SS was 41 thoroughly followed via isothermal calorimetry (IC) and conductivity analysis, and the nature of 42 hydration products was studied by X-ray diffraction (XRD) and thermogravimetric analysis (TGA). 43 It was concluded that alkanolamines facilitated the dissolution of minerals in SS and formation of 44 hydration products (e.g., C-S-H, C-A-H, C-F-H and Mc), and hence significantly enhanced the 45 microstructural development and engineering properties of SS. This work demonstrated a promising 46 way of upcycling SS as an effective and sustainable S/S agent for handling complex heavy metal 47 contaminated soil, with the potential of enhancing the SS utilization significantly.

48 Keywords: Alkanolamines, Steel slag activation, Heavy Metals, Soil Stabilization/Solidification

49 1. Introduction

50 Globally, heavy metals (HM) discharged from metal casting industries, fossil fuel burning, and the 51 ever-growing use of gasoline, paint, chemical fertilizer and pesticide have been accumulating in 52 soils in the past few decades[1]. Heavy metal-contaminated soil has become one of the most serious 53 environmental issues all over the world, threatening human health [2-4]. A national soil survey 54 found that in China 16.1% of the surveyed land exceeded national standards of soil contamination, 55 within which 19.4% of agricultural land and 34.9% of former industrial land were regarded as 56 contaminated [5-7], with HM as the most prevalent contaminants. HM such as cadmium (Cd), 57 copper (Cu), nickel (Ni), lead (Pb), and zinc (Zn) are highly toxic [8, 9]. Effective and sustainable 58 soil remediation technologies have been developed to treat HM-contaminated soil and achieved 59 great successes in the past few years[10].

60 Stabilization/solidification (S/S) is the most widely-used technology in China to treat HM-61 contaminated soil (48.5% adoption rate in the 2017-2018 year) [11-13]. The method involves using 62 binders to immobilize heavy metals in contaminated soil through physical encapsulation, adsorption 63 and chemical reactions, which decrease the bioavailability/ecotoxicity of the contaminants and 64 improve the engineering properties of the contaminated soils [14-16]. Previous studies had 65 highlighted the effectiveness of using highly alkaline cementitious materials in S/S, such as Portland 66 cement (PC), MgO-based materials and lime-fly ash blends [17-19]. However, the production of 67 these traditional binders was associated with intensive consumption of energy and nonrenewable 68 resources, and contributed to $\sim 10\%$ of anthropogenic greenhouse gas emissions [20]. Furthermore, 69 the high-alkaline binders may have adverse effects including incompatibility with HM, elevated soil 70 pH and high HM leachability, particularly under aggressive environmental conditions [21-24], 71 which limited the effectiveness of them in treating heavily-contaminated soils [25]. Therefore, it is 72 always desirable to develop alternative binders with higher efficiency, better stability, low-cost and 73 more environmentally friendly to remediate contaminated soils [12].

74 Steel slag (SS) is an alkaline industrial waste produced during the steelmaking process with an 75 annual production of approximately 15-20 wt% of the total steel output worldwide [26, 27]. In China, 76 the annual output of SS exceeded 100 million tons which accounted for approximately 24% of 77 Chinese total industrial solid waste; nonetheless, its utilization rate is less than 25% [28]. Therefore, 78 the large-scale utilization of SS is urgently needed as its disposal caused serious environmental 79 pollution and occupied valuable lands [29]. Depending on the steelmaking method, the main 80 chemical composition of SS is SiO₂, CaO, Al₂O₃, Fe₂O₃ and MgO. In terms of mineral forms, SS 81 mainly consists of tricalcium silicate (C₃S), dicalcium silicate (C₂S), C₄AF, C₁₂A₇, C₂F, RO phase 82 (metal oxides solid solution), free CaO and free MgO [30-33]. The composition of SS is similar to 83 PC which shows its potential to be utilized as an alternative green binder in S/S for treating HM-84 contaminated soil. However, the hydration activity of SS is much lower than that of PC [34], which 85 necessitates its activation prior to its application in S/S.

Alkali activation has been widely used to improve the hydraulic properties of SS using water glass, sodium hydroxide, sodium silicate and sodium sulfate, etc. [35, 36]. However, the production of those strong alkalis is not only associated with huge CO₂ emissions but also costly for large-scale SS utilization [37]. Additionally, owing to the ultra-high alkalinity of the alkali activators, the alkaliactivated SS would also elevate the soil alkalinity and hence adversely impact the ecological balance of the environment. Therefore, developing a low-cost, environment friendly and effective activator for SS would pave the way for its application in HM-contaminated soil remediation.

93 Recently, it has been reported that alkanolamines could affect the structure of hydration products in

94 PC, and different types of alkanolamines exhibited different impacts [38-40]. Other researchers also 95 found that alkanolamines could promote the hydration and chelating solubilization of SS [41, 42]. 96 Thus, alkanolamines activated SS may serve as a promising alternative binder to remediate HM-97 contaminated soil considering: (i) the potential large-scale utilisation of SS which would reduce the 98 negative environmental impact of SS accumulation; (ii) enhancement of the hydration of SS for 99 improved S/S performance, particularly the early-age properties; (iii) the low cost and low carbon 100 footprint of SS compared to PC. However, none has examined the performance of alkanolamines-101 activated SS for HM-contaminated soil remediation yet and there is a lack of understanding on the 102 activation mechanism and optimal dosage of this new type of activator (i.e. alkanolamines) for SS. 103 In this study, detailed analyses were conducted on the hydration kinetics, hydration products and 104 strength of the hydrated SS by different alkanolamines including triethanolamine (TEA), 105 triisopropanolamine (TIPA), ethyldiisopropylamine (EDIPA) and diethanolisopropanolamine 106 (DEIPA) using characterization methods such as X-ray diffraction (XRD), isothermal calorimetry, 107 thermogravimetric analysis (TGA) and Fourier transform infrared spectroscopy (FT-IR). The 108 performance (i.e., strength and leachability of HM) of activated-SS treated HM-contaminated soils 109 was assessed within 180 days to investigate the temporal effect of the type and dosage of the 110 activators.

111 2. Materials and methods

112 2.1 Preparation of binders

The SS used in this study was derived from the Meishan Iron & Steel plant in China. Raw SS lumps were crushed, ball milled and then passed through a 20-mesh screen to obtain the SS powder, with its chemical compositions presented in Table 1. Both the concentrations of Cr and V ions in the TCLP leachates were below the detection limits indicating the low leachability of the two potential contaminants from the SS. The mineral components of the SS were examined by XRD (Fig. 1), which shows that the it mainly consists of β -larnite (β -C₂S), Calcite (CaCO₃), srebrodolskite (C₂F),

119 mayenite $(C_{12}A_7)$, tricalcium aluminate (C_3A) .

120 Fig. 2 shows the chemical structures of the four types of alkanolamines (AR grade) used in this

121 work, of which the TEA and TIPA were provided by Aladdin corporation, and EDIPA and DEIPA

122 were provided by Hongbaoli Co., Ltd. The activators were combined with SS powders according to

the proportions in Table 2 and then milled in a planetary ball mill at a speed of 220 r/min for 30 min

to obtain the activated SS.

125 Table 1

126 Chemical compositions of SS measured by XRF.





131 Table 2

127 128

129

132 Th	e type and content	(by weight of SS)	of alkanolamines u	sed as activators
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	Alkanolamines	Content (%)
Control	None	-
TIPA-0.02	TIPA	0.02
TIPA-0.05	TIPA	0.05
TIPA-0.08	TIPA	0.08
TIPA-0.1	TIPA	0.1
TEA-0.05	TEA	0.05

EDIPA-0.05	EDIPA	0.05
DEIPA-0.05	DEIPA	0.05

133 *2.2 Mechanistic study on the activation of SS by alkanolamines*

134 2.2.1 Compressive strength tests of activated SS paste

The SS pastes activated by alkanolamines with water to solid ratio of 0.2 were prepared and cast in 20 mm \times 20 mm \times 20 mm moulds. These samples were cured in a moist cabinet under the condition of 20 \pm 1 °C, 95 \pm 1% relative humidity for 24h, and then demoulded and placed in the isothermal curing cabinet under the same condition until the testing age (1d, 3d, 7d, 28d, 90d and 180d). The mean compressive strength of six cement pastes was recorded for each mix. The crushed samples were ground, sieved and then immersed in ethanol for terminating hydration, followed by drying at 60°C for 24 h in a vacuum oven until further characterization.

142 2.2.2 Isothermal calorimetry

The hydration heat of activated SS was measured by a Thermometric TAM Air isothermal calorimeter (TA Instruments) at 20 ± 0.02 °C. Approximately 4 g of dry powders were loaded in glass ampoules, and syringes were loaded with 2 g of water. When a steady baseline was reached, the solution was injected into glass ampoules and externally stirred for 20 s. Then the glass ampoule was sealed and placed into the isothermal calorimeter. The heat of hydration was measured for 3 days to study the effect of activator on the hydration of SS.

149 2.2.3 Liquid phase conductivity

The dissolution rates of the reference and activated SS were followed by measuring the conductivities of the SS slurries. The slurries were prepared by mixing the SS and water with a liquid/solid ratio of 5. The magnetic stirrer was used for the mixing process at 250 rpm and stirred for 48h. The conductivities of the slurries were continuously recorded by a conductivity meter (DDSJ-308A, made in Shanghai Yueping).

155 2.2.4 X-ray diffraction (XRD)

- 156 XRD analysis was conducted on D/max-2500 X-ray diffraction of Rigaku, Japan, with CuKα
- 157 radiation, 40 kV voltage, 200 mA current, 2θ between 5° and 80°, 0.02°/s scan speed and 0.02° step
- 158 size to characterize the mineralogical phases of SS at different ages.
- 159 2.2.5 Thermogravimetry/differential scanning calorimetry (TG-DTG)
- 160 Thermogravimetric and differential thermogravimetric analysis (TG/DTG) was operated under N₂

161 flow with heating rate of 10°C/min from the ambient temperature to 1000°C using STA409C

162 instrument of NETZSCH.

163 2.3 Preparation of contaminated soils

164 A clean soil was obtained by sampling a surface soil up to 50 cm depth from Xinxiang of Henan 165 Province, China, and dried in an oven at 105°C for 6 hours. The dried soil was crushed, ground and 166 then passed through a 1 mm sieve and stored in a polyethylene container for the subsequent 167 physicochemical properties tests and the results were shown in Table 3. HMs were not leached from 168 clean soil and SS used in the experiments by the TCLP method. In this work, a heavily HM-169 contaminated soil was prepared via doping Cd, Cu, Ni, Pb and Zn (in the form of Cd(NO₃)₂·4H₂O, 170 Cu(NO₃)₂ · 3H₂O, Ni(NO₃)₂ · 6H₂O, Pb(NO₃)₂, Zn(NO₃)₂ · 6H₂O respectively, AR grade from 171 Sinopharm Chemical Reagent Co., Ltd.) to the clean soil. Predetermined amounts of HM (i.e., 24 172 mg/kg of Cd, 10000 mg/kg of Cu, 500 mg/kg of Ni, 280 mg/kg of Pb and 12500 mg/kg of Zn by 173 weight of dry soils) were firstly dissolved into the solution and added to the dry soil (to keep the 174 moisture content at 20%) and then stirred vigorously for 30 min to prepare the co-contaminated soil. 175 The mixture was sealed and kept for 24 hours to ensure the adequate distribution of HM in soil. The 176 binder and water were then added and mixed to achieve homogeneity. The weight ratio of binder to soil is 2:8 and the final weight ratio of water to solid (including soil and binder) was determined as 177 178 25:100 since preliminary results showed that consistencies of the mixtures were optimal at this level 179 (i.e., neither too dry nor too wet for handling). After that, approximately 200 g of the mixture was 180 statically compacted by a stainless steel cylindrical mold with 50 mm diameter and 50 mm height. 181 Then, the specimen was carefully extruded from the mold using a hydraulic jack and sealed in a 182 polyethylene bag for curing under the standard condition (temperature 20 ± 2 °C, relative humidity 183 99%). Specimens were collected for various tests at ages of 1, 3, 7, 28, 90 and 180 days.

- 184 Table 3
- 185 Basic physicochemical properties of the soil

Property	Value ^b	Test method	
Specific gravity, G_s	2.59	ASTM D854-14	
Liquid limit, w_L (%)	33.4	ASTM D4318-10	
Plastic limit, w_P (%)	17.2	ASTM D4318-10	
Optimum water content, w_{opt} (%)	21.8	ASTM D698-12	

Maximum dry density, ρ_d (g/cm ³)	1.82	ASTM D698-12
Average soil pH	8.19	ASTM D4972-13
Soil classification	CL	ASTM D2487-11
Grain size distribution (%) ^a		-
Clay (<0.002 mm)	22.5	
Silt (0.002–0.075 mm)	51.7	
Sand (0.075–2 mm)	25.8	

^a Measured using a laser particle size analyzer Mastersizer 2000 (Malvern, USA).

187 ^b Number of replicate = 3, and coefficient of variance (COV) < 5%.

188 2.4 Unconfined compressive strength (UCS) tests for contaminated soils

The microcomputer-controlled electronic universal testing machine of Shanghai Yihuan Instrument Technology Co., Ltd was used to assess the UCS of contaminated soils according to JTG E51-2009[43], at a speed of 1 mm·min⁻¹. The mean UCS of six soil samples was recorded for each mix and presented here. The crushed samples were collected, ground and sieved through a 4 mm mesh before leaching tests.

194 2.5 Leaching test for contaminated soils

195 After incubation for designated time periods, specimens were firstly dried at 65 °C to achieve 196 constant weights. The leachabilities of Cd, Cu, Ni, Pb and Zn in the samples was evaluated 197 according to US EPA Method 1311 - Toxicity Characteristic Leaching Procedure (TCLP). Briefly, 198 approximately 5 g of the crushed sample was added to ~96.5 mL deionized water and stirred for 5 199 min and the pH value (which determines buffer solution chose) of the mixture was recoded with a 200 pH meter (Rex PHS-3E). The soil and buffer solution (HOAc/NaOAc, pH 4.93) were mixed with a 201 solid/liquid ratio of 1:20 in a 2 L polyethylene bottle and shaken at 250 rpm for 18 h. Cd, Cu, Ni, 202 Pb and Zn concentrations in the filtrates were measured by ICP-OES after filtration with 0.45µm 203 filter, dilution (if necessary) and acidification to pH < 2.

204 2.6 Statistical analysis

205 All compressive strength experiments were carried out in sextuplicate, and leaching experiments

- were carried out in triplicate. The mean and standard deviations of each experiment were presented.
- 207 The significance of differences between groups was determined by one-way ANOVA using Duncan
- 208 method with a significance level of 0.05 using SPSS 17, and indicated by different lowercase letters
- in the figures.

210 **3. Results**

211 *3.1 Hydration properties of alkanolamine-activated SS*

As the capacity to solidify and stabilize HM was closely related to the binder's hydrationcharacteristics, the following sections will focus on the hydration mechanisms and evolution of

214 mineral compositions of the alkanolamine-activated SS.

215 *3.1.1 Heat evolution*

The reaction of SS with water was a thermodynamic reaction accompanied by exothermic behavior, which could be recorded using conduction calorimetry. The heat release curves were characterized by the presence of two peaks. The initial peak was attributed to the dissolution of f-CaO, C₃A and $C_{12}A_7$ [44, 45], leading to the continuous accumulation of ions. When the ion concentrations reached a certain level, the second exothermic peak appeared, ascribed to the precipitation of the CH (portlandite) and C-S-H (calcium silicate hydrate) phases [46] and the dissolution of β -C₂S and C₂F [47]. The hydration process gradually stabilized after the second exothermic peak.

Fig. 3 shows the effect of different alkanolamines on the hydration heat evolution rate and cumulative hydration heat of SS. All the alkanolamines increased the heat generation significantly. No significant difference was observed for the appearance time of the first exothermic peak, while the second peak appeared the earliest for TEA and latest for TIPA. The second peak was not observed for the control sample, which is likely due to the low hydration reactivity of the SS used in this work and hence the 2nd peak may appear beyond 72 hours.

Fig. 4 presents the effect of TIPA dosage on the hydration heat evolution rate and cumulative hydration heat. The first peak increased significantly with the increased TIPA content indicated that a higher dosage of TIPA improved the dissolution of mineral phases. Moreover, higher TIPA dosage shortened the induction period for the second peak, implying that time had reduced for the ion concentrations to reach supersaturation with the addition of more TIPA. The cumulative hydration heat increased proportionally with the dosage of TIPA.



236 237 Fig. 3 Hydration heat evolution of different alkanolamines-activated SS: (a) Heat flow of the whole hydration

process, (b) heat flow in the first 9 h (Peak 1) and (c) cumulative hydration heat.

Gontrol Control Peak 1 Control TIPA-0.05 TIPA-0.05





Fig. 4 Hydration heat evolution of TIPA-activated SS at different TIPA dosages: (a) Heat flow of the whole hydration process, (b) heat flow in the first 9 h (Peak 1) and (c) cumulative hydration heat.

240

241 *3.1.2 Conductivity of SS slurries*

242 The ion concentrations in the aqueous phase could be indirectly assessed by measuring the 243 conductivity of the solution, which could be used to evaluate the dissolution rate of SS [48]. Fig. 5 244 shows the evolution of conductivity in the SS slurries with and without alkanolamines. The 245 conductivity reached the peak at the first 5 min due to the dissolution of the high solubility mineral 246 phases (e.g., free lime and aluminates). This was followed by the ionic reactions in the solution 247 leading to rapidly decreased conductivity. After approximately 60 min, the conductivities of the SS 248 solutions with activators rose again and then dropped gradually over time, while this second peak 249 was not observed in the control sample. Apparently, the activators promoted the dissolution of SS 250 again when the ion concentration dropped, which induced the second peak.

251 For the control sample, the precipitation of the hydration products continuously consumed the 252 dissolved ions. Meanwhile, the dissolved cations adsorbed on the surface of SS particles and formed 253 a coating layer, which was not conducive to further dissolution [49]. This explains the gradually 254 decreased conductivity and higher residual conductivity over time due to the diffusion-controlled 255 dissolution of SS particles over time, ascribed to the surface coating. With the addition of 256 alkanolamines, the chelating of cations led to exposed SS surface for further dissolution, which was 257 evidenced by the second peak. Moreover, chelation likely resulted in higher supersaturation degrees 258 of the ions in the solution, which induced more hydration/precipitation products. This is the reason 259 of the much lower residual conductivities of the SS samples with activators. Among the four 260 alkanolamines, TIPA showed the highest second peak and lowest residual conductivity, indicating

its best performance as the chelating agent for promoting the hydration of SS. Fig. 5 (b) shows that
the effect of TIPA is highly correlated with its dosage. Increasing the dosage from 0.02% to 0.1%
proportionally increased the conductivity of the second peak and reduced the residual conductivity
of the solution.



265

Fig. 5 Evolution of conductivity in the SS solutions with and without alkanolamines: (a) SS with four different
alkanolamines at the same dosage of 0.05%, (b) SS with 0.02%, 0.05%, 0.08% and 0.1% TIPA.

268 3.1.3 XRD

The hydration products were characterized by XRD in Fig. 6. Compared with unhydrated SS (Fig. 269 270 1), the peaks of C₃A disappeared and the peaks of β -C₂S, C₂F, C₁₂A₇ and CaCO₃ weakened. 271 Moreover, the peaks of hydration products such as Ca₃Fe₂Si_{1.15}O_{4.6}(OH)_{7.4}, Ca₃Al₂(O₄H₄)₃ and 272 $C_4(A,F)$ ČH₁₁ appeared, which could be formed by the hydration of C₃A, C₁₂A₇, C₂F and CaCO₃. 273 Moreover, hydration of β -C₂S would form C-S-H and these hydration products would potentially 274 contribute to the cementitious and metal-binding capability in soil S/S. As shown in Fig. 6 (a), the 275 peaks of hydration products of activated SS were stronger than those of SS without activators. The enlarge view ($2\theta = 31.5^{\circ}-34.0^{\circ}$) clearly exhibits that the characteristic peaks of β -C₂S and C₂F 276 277 weakened further after the addition of activators, indicating higher dissolution/reaction rates of these 278 mineral phases. Consistent with the conductivity results, SS activated by TIPA exhibited the lowest 279 peaks of β-C₂S, C₂F and highest peaks of the hydration products compared with other activators. 280 Fig. 6 (b) further demonstrated that the dissolution of the minerals in SS and the precipitation of the 281 hydration products were promoted by the increased dosage of TIPA from 0.02% to 0.1%.



200

285

284

alkanolamines, (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

Fig. 6 XRD patterns of phase compositions in hardened SS pastes at 28 days activated by: (a) four different

287 During SS hydration, mineral phases reacted with water to form hydration products, and hydration 288 products were readily carbonated to form corresponding carbonates. In order to investigate the 289 effects of activator type and dosage on the hydration of SS, the hydration products of SS paste were 290 characterized by TGA as shown in Fig. 7. It could be seen from the derivative thermogravimetric 291 (DTG) curve that there were mainly three mass loss peaks. The peak at \sim 140 °C is mainly due to the 292 dehydration of C-S-H gel and Mc (monocarboaluminate). The peak at ~280 °C is mainly ascribed to 293 the dehydration of C-A-H (calcium aluminate hydrate) and C-F-H (calcium ferrite hydrate) [29, 50]. 294 In the range of 650-750 °C, the peak is mainly due to the decarbonation of CaCO₃ [51]. C-S-H gel 295 was generated from the hydration of silicate phases, C-A-H and Mc from aluminates, and C-F-H

^{286 3.1.4} Thermogravimetric analysis (TGA)

from C_2F . Moreover, a portion of CaCO₃ was reacted and produced Mc.

297 Fig. 7(a) showed the TG and DTG curves of the SS pastes at the age of 28 days activated by the 298 four alkanolamines. Compared with the control sample, the first two peaks were much more 299 pronounced in the samples with activators, while the one associated with $CaCO_3$ was weaker. 300 Moreover, TIPA-activated SS showed the maximum weight losses due to C-S-H, Mc, C-A-H and 301 C-F-H decomposition and the minimum weight loss of CaCO₃ decomposition. In Fig. 7(b), higher 302 dosage of TIPA increased the peaks associated with hydration products while decreased the one with 303 CaCO₃. The TGA results agreed well with other tests on the best activation performance of TIPA 304 and the enhanced effect by increasing its dosage.



Fig. 7 TG-DTG curves of hardened SS pastes at 28d activated by: (a) four different alkanolamines at the same
addition dosage of 0.05%, (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

308 *3.2 Compressive strength of alkanolamine-activated SS pastes*

305

309 The effects of four different alkanolamines on the mechanical properties of hardened SS pastes were 310 evaluated by measuring the compressive strength of SS pastes at different ages as shown in Fig. 311 8(a). It could be seen that the compressive strength gradually increased with curing time with the 312 rate decreased. Moreover, the activators significantly improved the compressive strength of the 313 hardened SS pastes with the effectiveness decreasing in the order of TIPA>EDIPA>DEIPA>TEA. 314 Specifically, at 28d, the compressive strength of the SS paste without activators was only 14.8 MPa, 315 which was increased to 27.1, 34.6, 30.9 and 28.6 MPa after adding 0.05 % TEA, TIPA, EDIPA and 316 DEIPA, respectively. Fig. 8(b) showed the effect of TIPA dosage on the compressive strength, 317 demonstrating the positive correlation between TIPA dosage and the improvement of compressive 318 strength, agreeing well with the increased quantities of hydration products. It is worth noting that 319 adding only 0.1% of TIPA to SS enhanced the compressive strength by \sim 190% and \sim 140% at 28d





335

Fig. 8 Effect of activators on compressive strength of activated SS pastes: (a) four alkanolamines (TEA, TIPA, EDIPA and DEIPA) at the same dosage of 0.05% and (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

324 3.3 UCS of activated-SS treated HM-contaminated soil

325 The UCS test has been widely used to describe the mechanical properties of S/S soils. The type of 326 binder and curing age have significant impacts on the physicochemical properties of the HM-327 contaminated soil treated with S/S. In general, the development of UCS of treated soils was 328 consistent with the results of the paste samples (comparing Fig.9 and Fig. 8). All the activators 329 showed excellent performance in the presence of high concentrations of multiple HMs in the soils. 330 TIPA-activated SS exhibited the highest strength among all the alkanolamines (Fig. 9a) and its effect 331 on UCS was positively correlated with the dosage (Fig. 9b). The UCS results demonstrated that by 332 adding as low as 0.02% of TIPA to SS, the strength could be improved by more than 100% at all 333 ages, which showed their excellent compatibility with HMs and great promise to reduce the use of 334



- **336** Fig. 9 Effect of activated-SS on the UCS of HM-contaminated soil: (a) four alkanolamines (TEA, TIPA, EDIPA
- 337

and DEIPA) at the same dosage of 0.05% and (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

- 338 3.4 TCLP results of activated-SS treated HM-contaminated soil
- 339 *3.4.1 Effect of different alkanolamines*

340 The concentrations of leached HM, i.e., Cd, Cu, Ni, Pb and Zn in contaminated soils treated by 341 alkanolamine-activated SS were shown in Fig. 10. Apparently, the leachability of all the HM were 342 significantly reduced after S/S over time, which indicated that SS exhibited superior capacity in 343 solidifying and stabilizing the HM in the contaminated soil. With activators, the effectiveness of 344 HM stabilization was improved, while the type of activator did not seem to influence the early-age 345 leachability of HM significantly up to 7 days. Above 28 days, TIPA and EDIPA showed the best 346 performance among all the alkanolamines for the TCLP results. Moreover, it is worth noting that 347 0.5% of TIPA successfully lowered the leached concentrations of all the HM (except for Ni, which 348 barely exceeded the limit) to below their regulatory limits (see Standard for Pollution Control on 349 the Hazardous Waste Landfill (GB 18598-2019) [52]) in China after curing of 28 days.





353 Fig. 10 Effect of different alkanolamines (TEA, TIPA, EDIPA and DEIPA at 0.05% dosage) activated SS on the
354 leachability of HM: (a) Cd, (b) Cu, (c) Ni, (d) Pb and (e) Zn.

355 3.4.2 Effect of TIPA dosage

According to the above results, TIPA showed the highest effectiveness to solidify and stabilize HM in contaminated soil. Thus, the effect of its dosage on the leachability of HM was investigated with TCLP results shown in Fig. 11. All the HM showed decreased leached concentrations over time. Consistent with the hydration and mechanical results, more TIPA showed improved HM immobilization capacity due to the enhanced hydration of SS. At the dosage > 0.8% the leachability of Ni also dropped to below its regulatory limit at 90 days.



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Fig. 11 Effect of dosage of TIPA (0.02%, 0.05%, 0.08% and 0.1%) activated SS on the leachability of HM: (a) Cd,
(b) Cu, (c) Ni, (d) Pb and (e) Zn.

367 4. Discussion

368 Based on the experimental results presented above, TIPA was found to be the best activator among 369 the four types of alkanolamines, which was studied in detail. Hence the following sections will use 370 TIPA as the representative alkanolamine to elucidate its effect on SS hydration and soil S/S 371 performance.

372 *4.1 Effect on the hydration process*

The hydration process of TIPA-activated SS was characterized by the evolution of hydration heat and conductivity over time, and the results agreed well (see Figs. 4 & 5). The hydration process is hence divided into two stages. The first stage is the continuous dissolution of minerals and gradual accumulation of ions in the pore solution. TIPA effectively promotes the dissolution of SS, which was evidenced by the enhancement of the first exothermic peak of hydration heat and the first conductivity peak. This effect is more pronounced with the increased dosage of TIPA. Moreover, the dormant periods between the first and second peak (not observed in the reference SS samples 380 within the test time) in both the heat and conductivity evolution curves were significantly reduced, 381 indicating the fast accumulation of ions to achieve the supersaturations for hydration products, 382 which induced the second stage. The second stage is the precipitation of hydration products. The 383 enhancement of second peak of hydration heat and the increased difference between the peak and 384 residual conductivity showed that TIPA led to the formation of more hydration products. Similarly, 385 this effect is positively correlated with the dosage of TIPA. This was further evidence by the XRD 386 and TGA results. Hence, the strength of the SS pastes improved remarkably with the increase of 387 TIPA content (Fig. 8).

388 *4.2 Effect on the hydration products*

As shown in the XRD and TGA (see Figs. 6 &7), TIPA promoted the dissolution of minerals in SS and the formation of hydration products. This effect is mainly exerted in two ways. The first one is the promotion of the hydration of C_2S , aluminates and C_2F , resulting in the formation of C-S-H, C-A-H and C-F-H. The other way is the enhanced reaction between C-A-H, C-F-H and CaCO₃, thereby producing a large amount of Mc.

The mineral phases in SS first dissolve into ions, which accumulate in the cement pore solution (Equations 1-3), and they react with each other to form hydration products. The alkanolamines chelated with the metal ions, which increased the solubility of the minerals and the concentrations of ions [29, 42].

$$398 \quad 2(2CaO \cdot SiO_2) + 4H_2O \to 3CaO \cdot 2SiO_2 \cdot 3H_2O + Ca^{2+} + 2OH^-$$
(1)

$$369 \quad 3CaO \cdot Al_2O_3 + 6H_2O \to 3Ca^{2+} + 2Al(OH)_4^- + 4OH^-$$
(2)

$$400 \quad 2CaO \cdot Fe_2O_3 + H_2O + 2OH^- \to 2Ca^{2+} + 2Fe(OH)_6^{3-} \tag{3}$$

The ions in the pore solution were chelated by TIPA and would not immediately participate in the
precipitation reaction. According to Le Chatelier's principle, the forward progress of reactions (1)(3) were facilitated until a new value of the dissolution equilibrium constant K was reached [53].
Thus, TIPA promoted the dissolution of mineral phases and increased the amount of various
dissolved metal ions in the solution. The precipitation reactions of ions are described in Equations
(4) - (8).

$$407 \quad 3Ca^{2+} + 2Al(OH)_{4}^{-} + 4OH^{-} \rightarrow 3CaO \cdot Al_{2}O_{3} \cdot 6H_{2}O \tag{4}$$

$$408 \qquad 3Ca^{2+} + 2Fe(OH)_6^{3-} \to 3CaO \cdot Fe_2O_3 \cdot 6H_2O \tag{5}$$

$$409 \quad 3Ca^{2+} + 2Al(OH)_4^- + CaCO_3 + 4OH^- + 5H_2O \to 3CaO \cdot Al_2O_3 \cdot CaCO_3 \cdot 11H_2O \tag{6}$$

410
$$3Ca^{2+} + 2Fe(OH)_6^{3-} + CaCO_3 + 5H_2O \rightarrow 3CaO \cdot Fe_2O_3 \cdot CaCO_3 \cdot 11H_2O$$
 (7)

411
$$4Ca^{2+} + (2x)Al(OH)_{4}^{-} + [2(1-x)]Fe(OH)_{6}^{3-} + CO_{3}^{2-} + 4xOH^{-} + 5H_2O$$

$$412 \quad \rightarrow \ 3CaO \cdot xAl_2O_3 \cdot (1-x)Fe_2O_3 \cdot CaCO_3 \cdot 11H_2O \tag{8}$$

The precipitation led to the reduction of metal ion concentrations in the pore solution, resulting in the release of ions from their chelated forms, further promoting the precipitation of more hydration products. The dissolution and precipitation reactions are hence both promoted by TIPA which acts like a "catalyst" in the process. In summary, TIPA promoted the dissolution of mineral phases such as C₂S, C₃A, C₁₂A₇, and C₂F, thereby increased the amount of metal ions in the solution, and hence promoted the subsequent precipitation reaction of generating C-S-H, C-A-H, C-F-H and Mc. Nevertheless, how different alkanolamines interacted with the ions and affected the hydration of SS

420 differently remain unclear.

421 4.3 Effect on the S/S performance for HM-contaminated soil

422 TIPA-activated SS showed excellent compatibility with the heavily contaminated soil, which was 423 evidenced by the UCS of the mixtures (see Fig.9). Moreover, the leachability of all HM (except for 424 Ni) can be successfully reduced to below their regulatory limits with only 0.5% of TIPA in SS at the 425 curing age of 28 days, while Ni was successfully treated at 90 days with 0.8% TIPA-activated SS. 426 The outstanding S/S performance is partly due to the enhanced solidification due to better 427 cementitious properties of TIPA-activated SS and also due to the enhanced chemical stabilisation of 428 HM in the hydration products such as C-S-H, C-A-H, C-F-H and Mc [54-56]. Moreover, TIPA may 429 also play a role in stabilizing HM by chelation, which warrant further studies. Leaching effect of 430 HM in the hydration products of alkanolamines activated SS was shown in Fig. S1 and Fig. S2. The 431 leaching of each heavy metal remained relatively constant after the age of 28d, which demonstrates 432 that the solidification and stabilization of heavy metals by SS hydration products is firm and stable. 433 This study demonstrated that alkanolamines could improve the capacity of SS to solidify and 434 stabilize HM-contaminated soil. Firstly, it provides a sustainable alternative to Portland cement in 435 S/S to reduce energy consumption and carbon emission in cement production. Secondly, it will

enhance the utilization of SS to reduce the environmental impact and waste of land resources.
However, there are many challenges and opportunities to apply this novel binder in the field which
include but not limited to: (i) optimisation of the alkanolamines/SS and SS/soil ratios depending on
the type, concentration of the contaminants and the soil type; (ii) long-term performance of the
treated soils under different climatic conditions; and (iii) variability of the SS composition in the
world and hence the potential of leaching inherent contaminants in SS in the long term.

442 **5.** Conclusion

Four types of alkanolamines (TEA, TIPA, EDIPA and DEIPA) were used to activate SS and then remediate HM-contaminated soil. The effects of alkanolamines on the hydration process and hydration products of SS were thoroughly investigated via a series of techniques. Moreover, their effects on the stabilization/solidification performance of HM-contaminated soil were assessed via strength and leaching tests. The following conclusions could be drawn:

1. The addition of alkanolamines promoted the hydration process of SS and hence facilitated the formation of hydration products. The chelation of dissolved metal ions promoted the dissolution of mineral phases in SS and the subsequent precipitation reaction. The hydration products such as C-S-H, C-A-H, C-F-H and Mc were beneficial to solidify and stabilize heavy metal contaminated soil. 2. TIPA showed the best activation effect among all the investigated alkanolamines and better promotion effect could be obtained with higher dosage. Compared to the reference SS paste, the compressive strength of the 0.1% TIPA-activated SS was enhanced by 190.5% at 28 days.

455 3. TIPA-activated SS showed excellent performance in HM-contaminated soil 456 stabilization/solidification. Leached concentrations of Cd, Cu, Ni, Pb, and Zn have reduced by 457 87.2%, 78.8%, 62.4%, 73.6% and 64.5% using 0.1% TIPA-activated SS after 28 days, and they were 458 all below their respective regulatory limits by Standard for Pollution Control on the Hazardous 459 Waste Landfill (GB 18598-2019) in China. Compared to the reference SS, the UCS of the treated 460 soil at 28 days was enhanced by 237.7% using 0.1% TIPA-activated SS.

461 This study proposed a promising way of upcycling SS as a soil remediation agent to promote circular 462 economy, which would not only alleviate the environmental problems of disposing SS but also 463 provide a sustainable alternative to Portland cement in stabilization/solidification, contributing to a 464 low-carbon future. 465

466 **Declaration of Competing Interest** 467 We declare that we do not have any commercial or associative interest that represents a conflict of 468 interest in connection with the work submitted. 469 470 Acknowledgments 471 We are grateful to the financial supports by State Key Laboratory of Materials-Oriented Chemical 472 Engineering (No.SKL-MCE-22A07), National Natural Science Foundation of China (52272018), 473 National Key R&D Program of China (2021YFB3802002), a Project Funded by the Priority 474 Academic Program Development (PAPD) of Jiangsu Higher Education Institutions. 475 476 References 477 [1] B. Zeng, O. Wang, L. Mo, F. Jin, J. Zhu, M. Tang, Synthesis of Mg-Al LDH and its calcined form 478 with natural materials for efficient Cr(VI) removal, Journal of Environmental Chemical Engineering, 479 10 (2022). 480 [2] M.-K. Zhang, Z.-Y. Liu, H. Wang, Use of Single Extraction Methods to Predict Bioavailability of 481 Heavy Metals in Polluted Soils to Rice, Communications in Soil Science and Plant Analysis, 41 482 (2010) 820-831. 483 [3] Y.-L. Yang, K.R. Reddy, Y.-J. Du, R.-D. Fan, Sodium hexametaphosphate (SHMP)-amended calcium 484 bentonite for slurry trench cutoff walls: workability and microstructure characteristics, Canadian 485 Geotechnical Journal, 55 (2018) 528-537. 486 [4] Y. Yang, Reddy, KR, Du, YJ, Fan, RD, Short-Term Hydraulic Conductivity and Consolidation 487 Properties of Soil-Bentonite Backfills Exposed to CCR-Impacted Groundwater., JOURNAL OF 488 GEOTECHNICAL AND GEOENVIRONMENTAL ENGINEERING, 144 (2018). 489 [5] D. Hou, F. Li, Complexities Surrounding China's Soil Action Plan, Land Degradation & Development, 490 28 (2017) 2315-2320. 491 [6] Y. Song, N. Kirkwood, C. Maksimovic, X. Zheng, D. O'Connor, Y. Jin, D. Hou, Nature based 492 solutions for contaminated land remediation and brownfield redevelopment in cities: A review, Sci 493 Total Environ, 663 (2019) 568-579. 494 [7] C. Qu, W. Shi, J. Guo, B. Fang, S. Wang, J.P. Giesy, P.E. Holm, China's Soil Pollution Control: 495 Choices and Challenges, Environmental science & technology, 50 (2016) 13181-13183. 496 [8] Z. Shen, D. Hou, B. Zhao, W. Xu, Y.S. Ok, N.S. Bolan, D.S. Alessi, Stability of heavy metals in soil 497 washing residue with and without biochar addition under accelerated ageing, Sci Total Environ, 619-498 620 (2018) 185-193. 499 [9] Y.J. Du, M.L. Wei, K.R. Reddy, Z.P. Liu, F. Jin, Effect of acid rain pH on leaching behavior of cement 500 stabilized lead-contaminated soil, J Hazard Mater, 271 (2014) 131-140. 501 [10] D. Hou, A. Al-Tabbaa, D. O'Connor, Q. Hu, Y.-G. Zhu, L. Wang, N. Kirkwood, Y.S. Ok, D.C.W. 502 Tsang, N.S. Bolan, J. Rinklebe, Sustainable remediation and redevelopment of brownfield sites,

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1	Alkanolamines-activated steel slag for
2	stabilization/solidification of heavy metal contaminated soil
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11 12 13	 Highlights: Alkanolamines promoted the hydration of steel slag and with TIPA showing the best performance.
14 15	• The UCS of treated HM-contaminated soil at 28 days was more than tripled using 0.1% TIPA- activated SS compared to the non-activated SS.
16 17	• TCLP leached concentrations of Cd, Cu, Ni, Pb, and Zn were reduced by 87.2%, 78.8%, 62.4%, 73.6% and 64.5% using 0.1% TIPA-activated SS at 28 days.
18 19	• Alkanolamines-activated SS is a sustainable alternative to PC in S/S for heavily-contaminated soil
20	Abbreviations:
21 22	SS: Steel slagS/S: Stabilization/SolidificationHM: Heavy metalsTEA: TriothanolominaTIBA: TriigenrongeneleminaFDIBA: Ethyldiisenrongenegene
22	DEIPA : Diethanolisopropanolamine UCS : Unconfined compressive strength
24	β -C ₂ S: β-larnite CaCO ₃ : Calcite C ₂ F: Srebrodolskite C ₁₂ A ₇ : Mayenite
25 26 27 28 29	C3A: Tricalcium aluminateC-S-H: Calcium silicate hydrateMc: MonocarboaluminateCH: PortlanditeC-A-H: Calcium aluminate hydrateC-F-H: Calcium ferrite hydrateIC: Isothermal calorimetryXRD: X-ray diffractionTGA: Thermogravimetric analysisTCLP: Toxicity Characteristic Leaching Procedure
30	Abstract: Steel slag (SS) is a byproduct discharged from steel-making industry and has not been
31	well utilized in contaminated soil stabilization/solidification (S/S) due to its low hydration activity.
32	In this study, four different alkanolamines (TEA, TIPA, EDIPA and DEIPA) were used to activate
33	SS and improve its cementitious properties and metal binding performance. The activated SS was
34	used to treat contaminated soils containing HM (heavy metals) of Cd, Cu, Ni, Pb and Zn. Compared

35 with the reference SS without activators, alkanolamines activated SS showed better S/S 36 performance in terms of strength and leaching, and the performance was improved by increasing 37 the activator dosage. For instance, concentrations of leached Cd, Cu, Ni, Pb, and Zn have reduced by 87.2%, 78.8%, 62.4%, 73.6% and 64.5% by using 0.1% TIPA-activated SS after 28 days, and 38 they were all below their respective regulatory limits by Standard for Pollution Control on the 39 40 Hazardous Waste Landfill (GB 18598-2019) in China. Compared to the reference SS, the unconfined compressive strength (UCS) of the treated soil at 28 days was enhanced by 237.7% 41 using 0.1% TIPA-activated SS. To elucidate the activation mechanism, the hydration process of SS 42 43 was thoroughly followed via isothermal calorimetry (IC) and conductivity analysis, and the nature of hydration products was studied by X-ray diffraction (XRD) and thermogravimetric analysis 44 45 (TGA). It was concluded that alkanolamines facilitated the dissolution of minerals in SS and 46 formation of hydration products (e.g., C-S-H, C-A-H, C-F-H and Mc), and hence significantly 47 enhanced the microstructural development and engineering properties of SS. This work 48 demonstrated a promising way of upcycling SS as an effective S/S agent for handling complex 49 heavy metal contaminated soil.

50 Abstract: Steel slag (SS) is a byproduct discharged from steel-making industry with less than 25% 51 utilization rate in China. The low utilisation rate of SS is associated with its low hydration activity 52 in cement and concrete. In this study, four different alkanolamines (TEA, TIPA, EDIPA and DEIPA) 53 were used to activate SS to improve its cementitious properties and metal binding performance, and 54 hence its capacity on treating heavy metal-contaminated soils containing Cd, Cu, Ni, Pb and Zn. 55 Compared with the reference SS without activators, concentrations of leached Cd, Cu, Ni, Pb, and 56 Zn have reduced by 87.2%, 78.8%, 62.4%, 73.6% and 64.5% by using 0.1% TIPA-activated SS 57 after 28 days, and they were all below their respective regulatory limits by Standard for Pollution 58 Control on the Hazardous Waste Landfill (GB 18598-2019) in China, and the unconfined 59 compressive strength (UCS) of the treated soil at 28 days was enhanced by 237.7% using 0.1% 60 TIPA-activated SS. To elucidate the activation mechanism, the hydration process of SS was 61 thoroughly followed via isothermal calorimetry (IC) and conductivity analysis, and the nature of 62 hydration products was studied by X-ray diffraction (XRD) and thermogravimetric analysis (TGA). 63 It was concluded that alkanolamines facilitated the dissolution of minerals in SS and formation of

hydration products (e.g., C-S-H, C-A-H, C-F-H and Mc), and hence significantly enhanced the
microstructural development and engineering properties of SS. This work demonstrated a promising
way of upcycling SS as an effective and sustainable S/S agent for handling complex heavy metal
contaminated soil, with the potential of enhancing the SS utilization significantly.

68 Keywords: Alkanolamines, Steel slag activation, Heavy Metals, Soil Stabilization/Solidification

69 1. Introduction

70 Globally, heavy metals (HM) discharged from metal casting industries, fossil fuel burning, and the 71 ever-growing use of gasoline, paint, chemical fertilizer and pesticide have been accumulating in 72 soils in the past few decades[1]. Heavy metal-contaminated soil has become one of the most serious 73 environmental issues all over the world, threatening human health [2-4]. A national soil survey 74 found that in China 16.1% of the surveyed land exceeded national standards of soil contamination, 75 within which 19.4% of agricultural land and 34.9% of former industrial land were regarded as 76 contaminated [5-7], with HM as the most prevalent contaminants. HM such as cadmium (Cd), 77 copper (Cu), nickel (Ni), lead (Pb), and zinc (Zn) are highly toxic [8, 9]. Effective and sustainable 78 soil remediation technologies have been developed to treat HM-contaminated soil and achieved 79 great successes in the past few years[10].

80 Stabilization/solidification (S/S) is the most widely-used technology in China to treat HM-81 contaminated soil (48.5% adoption rate in the 2017-2018 year) [11-13]. The method involves using 82 binders to immobilize heavy metals in contaminated soil through physical encapsulation, adsorption 83 and chemical reactions, which decrease the bioavailability/ecotoxicity of the contaminants and 84 improve the engineering properties of the contaminated soils [14-16]. Previous studies had 85 highlighted the effectiveness of using highly alkaline cementitious materials in S/S, such as Portland 86 cement (PC), MgO-based materials and lime-fly ash blends [17-19]. However, the production of 87 these traditional binders was associated with intensive consumption of energy and nonrenewable 88 resources, and contributed to $\sim 10\%$ of anthropogenic greenhouse gas emissions [20]. Furthermore, 89 the high-alkaline binders may have adverse effects including incompatibility with HM, elevated soil 90 pH and high HM leachability, particularly under aggressive environmental conditions [21-24], 91 which limited the effectiveness of them in treating heavily-contaminated soils [25]. Therefore, it is 92 always desirable to develop alternative binders with higher efficiency, better stability, low-cost and 93 more environmentally friendly to remediate contaminated soils [12].

94 Steel slag (SS) is an alkaline industrial waste produced during the steelmaking process with an 95 annual production of approximately 15-20 wt% of the total steel output worldwide [26, 27]. In China, 96 the annual output of SS exceeded 100 million tons which accounted for approximately 24% of 97 Chinese total industrial solid waste; nonetheless, its utilization rate is less than 25% [28]. Therefore, 98 the large-scale utilization of SS is urgently needed as its disposal caused serious environmental 99 pollution and occupied valuable lands [29]. Depending on the steelmaking method, the main 100 chemical composition of SS is SiO₂, CaO, Al₂O₃, Fe₂O₃ and MgO. In terms of mineral forms, SS 101 mainly consists of tricalcium silicate (C₃S), dicalcium silicate (C₂S), C₄AF, C₁₂A7, C₂F, RO phase 102 (metal oxides solid solution), free CaO and free MgO [30-33]. The composition of SS is similar to 103 PC which shows its potential to be utilized as an alternative green binder in S/S for treating HM-104 contaminated soil. However, the hydration activity of SS is much lower than that of PC [34], which 105 necessitates its activation prior to its application in S/S.

Alkali activation has been widely used to improve the hydraulic properties of SS using water glass, sodium hydroxide, sodium silicate and sodium sulfate, etc. [35, 36]. However, the production of those strong alkalis is not only associated with huge CO₂ emissions but also costly for large-scale SS utilization [37]. Additionally, owing to the ultra-high alkalinity of the alkali activators, the alkaliactivated SS would also elevate the soil alkalinity and hence adversely impact the ecological balance of the environment. Therefore, developing a low-cost, environment friendly and effective activator for SS would pave the way for its application in HM-contaminated soil remediation.

113 Recently, it has been reported that alkanolamines could affect the structure of hydration products in 114 PC, and different types of alkanolamines exhibited different impacts [38-40]. Other researchers also 115 found that alkanolamines could promote the hydration and chelating solubilization of SS [41, 42]. 116 Thus, alkanolamines activated SS may serve as a promising alternative binder to remediate HM-117 contaminated soil considering: (i) the potential large-scale utilisation of SS which would reduce the 118 negative environmental impact of SS accumulation; (ii) enhancement of the hydration of SS for 119 improved S/S performance, particularly the early-age properties; (iii) the low cost and low carbon 120 footprint of SS compared to PC. However, none has examined the performance of alkanolamines-121 activated SS for HM-contaminated soil remediation yet and there is a lack of understanding on the 122 activation mechanism and optimal dosage of this new type of activator (i.e. alkanolamines) for SS.

123 In this study, detailed analyses were conducted on the hydration kinetics, hydration products and 124 strength of the hydrated SS by different alkanolamines including triethanolamine (TEA), 125 triisopropanolamine (TIPA), ethyldiisopropylamine (EDIPA) and diethanolisopropanolamine 126 (DEIPA) using characterization methods such as X-ray diffraction (XRD), isothermal calorimetry, 127 thermogravimetric analysis (TGA) and Fourier transform infrared spectroscopy (FT-IR). The 128 performance (i.e., strength and leachability of HM) of activated-SS treated HM-contaminated soils 129 was assessed within 180 days to investigate the temporal effect of the type and dosage of the 130 activators.

131 2. Materials and methods

132 2.1 Preparation of binders

The SS used in this study was derived from the Meishan Iron & Steel plant in China. Raw SS lumps were crushed, ball milled and then passed through a 20-mesh screen to obtain the SS powder, with its chemical compositions presented in Table 1. Both the concentrations of Cr and V ions in the TCLP leachates were below the detection limits indicating the low leachability of the two potential contaminants from the SS. The mineral components of the SS were examined by XRD (Fig. 1),

which shows that the it mainly consists of β -larnite (β -C₂S), Calcite (CaCO₃), srebrodolskite (C₂F),

139 mayenite $(C_{12}A_7)$, tricalcium aluminate (C_3A) .

140 Fig. 2 shows the chemical structures of the four types of alkanolamines (AR grade) used in this

141 work, of which the TEA and TIPA were provided by Aladdin corporation, and EDIPA and DEIPA

142 were provided by Hongbaoli Co., Ltd. The activators were combined with SS powders according to

the proportions in Table 2 and then milled in a planetary ball mill at a speed of 220 r/min for 30 min

- to obtain the activated SS.
- 145 Table 1
- 146

Chemical compositions of SS measured by XRF.

Chemical											
composition	CaO	Fe ₂ O ₃	SiO ₂	Al ₂ O ₃	MgO	MnO	P_2O_5	TiO ₂	Cr ₂ O ₃	V_2O_5	LOI
(wt%)											
Steel slag	35.89	25.49	14.77	8.23	6.30	3.55	2.32	0.97	0.24	0.21	1.12

150 Fig. 2 Schematic representation of the molecular structures of the four alkanolamines used in this study

151 Table 2

152	The type and content	(by weight of	f SS) of alkand	plamines used	as activators
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	Alkanolamines	Content (%)
Control	None	-
TIPA-0.02	TIPA	0.02
TIPA-0.05	TIPA	0.05
TIPA-0.08	TIPA	0.08
TIPA-0.1	TIPA	0.1
TEA-0.05	TEA	0.05
EDIPA-0.05	EDIPA	0.05
DEIPA-0.05	DEIPA	0.05

153 2.2 Mechanistic study on the activation of SS by alkanolamines

154 2.2.1 Compressive strength tests of activated SS paste

157 of 20 ± 1 °C, $95 \pm 1\%$ relative humidity for 24h, and then demoulded and placed in the isothermal

¹⁵⁵ The SS pastes activated by alkanolamines with water to solid ratio of 0.2 were prepared and cast in

¹⁵⁶ $20 \text{ mm} \times 20 \text{ mm} \times 20 \text{ mm}$ moulds. These samples were cured in a moist cabinet under the condition

- curing cabinet under the same condition until the testing age (1d, 3d, 7d, 28d, 90d and 180d). The
 mean compressive strength of six cement pastes was recorded for each mix. The crushed samples
 were ground, sieved and then immersed in ethanol for terminating hydration, followed by drying at
 60°C for 24 h in a vacuum oven until further characterization.
- 162 *2.2.2 Isothermal calorimetry*
- The hydration heat of activated SS was measured by a Thermometric TAM Air isothermal calorimeter (TA Instruments) at 20 ± 0.02 °C. Approximately 4 g of dry powders were loaded in glass ampoules, and syringes were loaded with 2 g of water. When a steady baseline was reached, the solution was injected into glass ampoules and externally stirred for 20 s. Then the glass ampoule was sealed and placed into the isothermal calorimeter. The heat of hydration was measured for 3 days to study the effect of activator on the hydration of SS.
- 169 *2.2.3 Liquid phase conductivity*
- The dissolution rates of the reference and activated SS were followed by measuring the conductivities of the SS slurries. The slurries were prepared by mixing the SS and water with a liquid/solid ratio of 5. The magnetic stirrer was used for the mixing process at 250 rpm and stirred for 48h. The conductivities of the slurries were continuously recorded by a conductivity meter
- 174 (DDSJ-308A, made in Shanghai Yueping).
- 175 2.2.4 X-ray diffraction (XRD)
- 176 XRD analysis was conducted on D/max-2500 X-ray diffraction of Rigaku, Japan, with CuKα
- 177 radiation, 40 kV voltage, 200 mA current, 2θ between 5° and 80°, 0.02°/s scan speed and 0.02° step
- 178 size to characterize the mineralogical phases of SS at different ages.
- 179 2.2.5 Thermogravimetry/differential scanning calorimetry (TG-DTG)
- 180 Thermogravimetric and differential thermogravimetric analysis (TG/DTG) was operated under N₂
- 181 flow with heating rate of 10°C/min from the ambient temperature to 1000°C using STA409C
- 182 instrument of NETZSCH.
- 183 2.3 Preparation of contaminated soils
- 184 A clean soil was obtained by sampling a surface soil up to 50 cm depth from Xinxiang of Henan
- 185 Province, China, and dried in an oven at 105°C for 6 hours. The dried soil was crushed, ground and
- 186 then passed through a 1 mm sieve and stored in a polyethylene container for the subsequent

187	physicochemical properties tests and the results were shown in Table 3HMs were not leached from
188	clean soil and SS used in the experiments by the TCLP method. In this work, a heavily HM-
189	contaminated soil was prepared via doping Cd, Cu, Ni, Pb and Zn (in the form of Cd(NO ₃) ₂ ·4H ₂ O,
190	$Cu(NO_3)_2 \cdot 3H_2O$, $Ni(NO_3)_2 \cdot 6H_2O$, $Pb(NO_3)_2$, $Zn(NO_3)_2 \cdot 6H_2O$ respectively, AR grade from
191	Sinopharm Chemical Reagent Co., Ltd.) to the clean soil. Predetermined amounts of HM (i.e., 24
192	mg/kg of Cd, 10000 mg/kg of Cu, 500 mg/kg of Ni, 280 mg/kg of Pb and 12500 mg/kg of Zn by
193	weight of dry soils) were firstly dissolved into the solution and added to the dry soil (to keep the
194	moisture content at 20%) and then stirred vigorously for 30 min to prepare the co-contaminated soil.
195	The mixture was sealed and kept for 24 hours to ensure the adequate distribution of HM in soil. The
196	binder and water were then added and mixed to achieve homogeneity. The weight ratio of binder to
197	soil is 2:8 and the final weight ratio of water to solid (including soil and binder) was determined as
198	25:100 since preliminary results showed that consistencies of the mixtures were optimal at this level
199	(i.e., neither too dry nor too wet for handling). After that, approximately 200 g of the mixture was
200	statically compacted by a stainless steel cylindrical mold with 50 mm diameter and 50 mm height.
201	Then, the specimen was carefully extruded from the mold using a hydraulic jack and sealed in a
202	polyethylene bag for curing under the standard condition (temperature 20 ± 2 °C, relative humidity
203	99%). Specimens were collected for various tests at ages of 1, 3, 7, 28, 90 and 180 days.

- 204 **Table 3**
- 205 Basic physicochemical properties of the soil

Property	Value ^b	Test method
Specific gravity, G_s	2.59	ASTM D854-14
Liquid limit, w_L (%)	33.4	ASTM D4318-10
Plastic limit, w_P (%)	17.2	ASTM D4318-10
Optimum water content, <i>w</i> _{opt} (%)	21.8	ASTM D698-12
Maximum dry density, ρ_d (g/cm ³)	1.82	ASTM D698-12
Average soil pH	8.19	ASTM D4972-13
Soil classification	CL	ASTM D2487-11
Grain size distribution (%) ^a		-
Clay (<0.002 mm)	22.5	
Silt (0.002–0.075 mm)	51.7	
Sand (0.075–2 mm)	25.8	

206 ^a Measured using a laser particle size analyzer Mastersizer 2000 (Malvern, USA).

207 ^b Number of replicate = 3, and coefficient of variance (COV) < 5%.

208 2.4 Unconfined compressive strength (UCS) tests for contaminated soils

209 The microcomputer-controlled electronic universal testing machine of Shanghai Yihuan Instrument

210 Technology Co., Ltd was used to assess the UCS of contaminated soils according to JTG E51-

2009[43], at a speed of 1 mm \cdot min⁻¹. The mean UCS of six soil samples was recorded for each mix

- and presented here. The crushed samples were collected, ground and sieved through a 4 mm mesh
- 213 before leaching tests.
- 214 2.5 Leaching test for contaminated soils

215 After incubation for designated time periods, specimens were firstly dried at 65 °C to achieve 216 constant weights. The leachabilities of Cd, Cu, Ni, Pb and Zn in the samples was evaluated 217 according to US EPA Method 1311 - Toxicity Characteristic Leaching Procedure (TCLP). Briefly, 218 approximately 5 g of the crushed sample was added to ~96.5 mL deionized water and stirred for 5 219 min and the pH value (which determines buffer solution chose) of the mixture was recoded with a 220 pH meter (Rex PHS-3E). The soil and buffer solution (HOAc/NaOAc, pH 4.93) were mixed with a 221 solid/liquid ratio of 1:20 in a 2 L polyethylene bottle and shaken at 250 rpm for 18 h. Cd, Cu, Ni, 222 Pb and Zn concentrations in the filtrates were measured by ICP-OES after filtration with 0.45µm 223 filter, dilution (if necessary) and acidification to pH < 2.

- 224 2.6 Statistical analysis
- 225 All compressive strength experiments were carried out in sextuplicate, and leaching experiments
- were carried out in triplicate. The mean and standard deviations of each experiment were presented.
- 227 The significance of differences between groups was determined by one-way ANOVA using Duncan
- 228 method with a significance level of 0.05 using SPSS 17, and indicated by different lowercase letters
- in the figures.
- 230 **3. Results**
- 231 3.1 Hydration properties of alkanolamine-activated SS
- As the capacity to solidify and stabilize HM was closely related to the binder's hydration
 characteristics, the following sections will focus on the hydration mechanisms and evolution of
 mineral compositions of the alkanolamine-activated SS.
- 235 3.1.1 Heat evolution
- 236 The reaction of SS with water was a thermodynamic reaction accompanied by exothermic behavior,

which could be recorded using conduction calorimetry. The heat release curves were characterized by the presence of two peaks. The initial peak was attributed to the dissolution of f-CaO, C₃A and $C_{12}A_7$ [44, 45], leading to the continuous accumulation of ions. When the ion concentrations reached a certain level, the second exothermic peak appeared, ascribed to the precipitation of the CH (portlandite) and C-S-H (calcium silicate hydrate) phases [46] and the dissolution of β -C₂S and C₂F [47]. The hydration process gradually stabilized after the second exothermic peak.

Fig. 3 shows the effect of different alkanolamines on the hydration heat evolution rate and cumulative hydration heat of SS. All the alkanolamines increased the heat generation significantly. No significant difference was observed for the appearance time of the first exothermic peak, while the second peak appeared the earliest for TEA and latest for TIPA. The second peak was not observed for the control sample, which is likely due to the low hydration reactivity of the SS used in this work and hence the 2nd peak may appear beyond 72 hours.

Fig. 4 presents the effect of TIPA dosage on the hydration heat evolution rate and cumulative hydration heat. The first peak increased significantly with the increased TIPA content indicated that a higher dosage of TIPA improved the dissolution of mineral phases. Moreover, higher TIPA dosage shortened the induction period for the second peak, implying that time had reduced for the ion concentrations to reach supersaturation with the addition of more TIPA. The cumulative hydration heat increased proportionally with the dosage of TIPA.

256 Fig. 3 Hydration heat evolution of different alkanolamines-activated SS: (a) Heat flow of the whole hydration

process, (b) heat flow in the first 9 h (Peak 1) and (c) cumulative hydration heat.

- 258
- 259 260

Fig. 4 Hydration heat evolution of TIPA-activated SS at different TIPA dosages: (a) Heat flow of the whole hydration process, (b) heat flow in the first 9 h (Peak 1) and (c) cumulative hydration heat.

261 *3.1.2 Conductivity of SS slurries*

262 The ion concentrations in the aqueous phase could be indirectly assessed by measuring the 263 conductivity of the solution, which could be used to evaluate the dissolution rate of SS [48]. Fig. 5 264 shows the evolution of conductivity in the SS slurries with and without alkanolamines. The 265 conductivity reached the peak at the first 5 min due to the dissolution of the high solubility mineral 266 phases (e.g., free lime and aluminates). This was followed by the ionic reactions in the solution 267 leading to rapidly decreased conductivity. After approximately 60 min, the conductivities of the SS 268 solutions with activators rose again and then dropped gradually over time, while this second peak 269 was not observed in the control sample. Apparently, the activators promoted the dissolution of SS 270 again when the ion concentration dropped, which induced the second peak.

271 For the control sample, the precipitation of the hydration products continuously consumed the 272 dissolved ions. Meanwhile, the dissolved cations adsorbed on the surface of SS particles and formed 273 a coating layer, which was not conducive to further dissolution [49]. This explains the gradually 274 decreased conductivity and higher residual conductivity over time due to the diffusion-controlled 275 dissolution of SS particles over time, ascribed to the surface coating. With the addition of 276 alkanolamines, the chelating of cations led to exposed SS surface for further dissolution, which was 277 evidenced by the second peak. Moreover, chelation likely resulted in higher supersaturation degrees 278 of the ions in the solution, which induced more hydration/precipitation products. This is the reason 279 of the much lower residual conductivities of the SS samples with activators. Among the four 280 alkanolamines, TIPA showed the highest second peak and lowest residual conductivity, indicating 281 its best performance as the chelating agent for promoting the hydration of SS. Fig. 5 (b) shows that 282 the effect of TIPA is highly correlated with its dosage. Increasing the dosage from 0.02% to 0.1%

283 proportionally increased the conductivity of the second peak and reduced the residual conductivity

Fig. 5 Evolution of conductivity in the SS solutions with and without alkanolamines: (a) SS with four different
alkanolamines at the same dosage of 0.05%, (b) SS with 0.02%, 0.05%, 0.08% and 0.1% TIPA.

288 3.1.3 XRD

285

289 The hydration products were characterized by XRD in Fig. 6. Compared with unhydrated SS (Fig. 290 1), the peaks of C₃A disappeared and the peaks of β -C₂S, C₂F, C₁₂A₇ and CaCO₃ weakened. 291 Moreover, the peaks of hydration products such as Ca₃Fe₂Si_{1.15}O_{4.6}(OH)_{7.4}, Ca₃Al₂(O₄H₄)₃ and 292 $C_4(A,F)$ ČH₁₁ appeared, which could be formed by the hydration of C₃A, C₁₂A₇, C₂F and CaCO₃. 293 Moreover, hydration of β -C₂S would form C-S-H and these hydration products would potentially 294 contribute to the cementitious and metal-binding capability in soil S/S. As shown in Fig. 6 (a), the 295 peaks of hydration products of activated SS were stronger than those of SS without activators. The 296 enlarge view ($2\theta = 31.5^{\circ}-34.0^{\circ}$) clearly exhibits that the characteristic peaks of β -C₂S and C₂F 297 weakened further after the addition of activators, indicating higher dissolution/reaction rates of these 298 mineral phases. Consistent with the conductivity results, SS activated by TIPA exhibited the lowest 299 peaks of β -C₂S, C₂F and highest peaks of the hydration products compared with other activators. 300 Fig. 6 (b) further demonstrated that the dissolution of the minerals in SS and the precipitation of the 301 hydration products were promoted by the increased dosage of TIPA from 0.02% to 0.1%.

305

304

alkanolamines, (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

Fig. 6 XRD patterns of phase compositions in hardened SS pastes at 28 days activated by: (a) four different

307 During SS hydration, mineral phases reacted with water to form hydration products, and hydration 308 products were readily carbonated to form corresponding carbonates. In order to investigate the 309 effects of activator type and dosage on the hydration of SS, the hydration products of SS paste were 310 characterized by TGA as shown in Fig. 7. It could be seen from the derivative thermogravimetric 311 (DTG) curve that there were mainly three mass loss peaks. The peak at \sim 140 °C is mainly due to the 312 dehydration of C-S-H gel and Mc (monocarboaluminate). The peak at ~280 °C is mainly ascribed to 313 the dehydration of C-A-H (calcium aluminate hydrate) and C-F-H (calcium ferrite hydrate) [29, 50]. 314 In the range of 650-750 °C, the peak is mainly due to the decarbonation of CaCO₃ [51]. C-S-H gel 315 was generated from the hydration of silicate phases, C-A-H and Mc from aluminates, and C-F-H

³⁰⁶ *3.1.4 Thermogravimetric analysis (TGA)*

316 from C_2F . Moreover, a portion of CaCO₃ was reacted and produced Mc.

317 Fig. 7(a) showed the TG and DTG curves of the SS pastes at the age of 28 days activated by the 318 four alkanolamines. Compared with the control sample, the first two peaks were much more 319 pronounced in the samples with activators, while the one associated with $CaCO_3$ was weaker. 320 Moreover, TIPA-activated SS showed the maximum weight losses due to C-S-H, Mc, C-A-H and 321 C-F-H decomposition and the minimum weight loss of CaCO₃ decomposition. In Fig. 7(b), higher 322 dosage of TIPA increased the peaks associated with hydration products while decreased the one with 323 CaCO₃. The TGA results agreed well with other tests on the best activation performance of TIPA 324 and the enhanced effect by increasing its dosage.

Fig. 7 TG-DTG curves of hardened SS pastes at 28d activated by: (a) four different alkanolamines at the same
addition dosage of 0.05%, (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

328 *3.2 Compressive strength of alkanolamine-activated SS pastes*

325

329 The effects of four different alkanolamines on the mechanical properties of hardened SS pastes were 330 evaluated by measuring the compressive strength of SS pastes at different ages as shown in Fig. 331 8(a). It could be seen that the compressive strength gradually increased with curing time with the 332 rate decreased. Moreover, the activators significantly improved the compressive strength of the 333 hardened SS pastes with the effectiveness decreasing in the order of TIPA>EDIPA>DEIPA>TEA. 334 Specifically, at 28d, the compressive strength of the SS paste without activators was only 14.8 MPa, 335 which was increased to 27.1, 34.6, 30.9 and 28.6 MPa after adding 0.05 % TEA, TIPA, EDIPA and 336 DEIPA, respectively. Fig. 8(b) showed the effect of TIPA dosage on the compressive strength, 337 demonstrating the positive correlation between TIPA dosage and the improvement of compressive 338 strength, agreeing well with the increased quantities of hydration products. It is worth noting that 339 adding only 0.1% of TIPA to SS enhanced the compressive strength by \sim 190% and \sim 140% at 28d

355

Fig. 8 Effect of activators on compressive strength of activated SS pastes: (a) four alkanolamines (TEA, TIPA,
EDIPA and DEIPA) at the same dosage of 0.05% and (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

344 3.3 UCS of activated-SS treated HM-contaminated soil

345 The UCS test has been widely used to describe the mechanical properties of S/S soils. The type of 346 binder and curing age have significant impacts on the physicochemical properties of the HM-347 contaminated soil treated with S/S. In general, the development of UCS of treated soils was 348 consistent with the results of the paste samples (comparing Fig.9 and Fig. 8). All the activators 349 showed excellent performance in the presence of high concentrations of multiple HMs in the soils. 350 TIPA-activated SS exhibited the highest strength among all the alkanolamines (Fig. 9a) and its effect 351 on UCS was positively correlated with the dosage (Fig. 9b). The UCS results demonstrated that by 352 adding as low as 0.02% of TIPA to SS, the strength could be improved by more than 100% at all 353 ages, which showed their excellent compatibility with HMs and great promise to reduce the use of 354

- 356 Fig. 9 Effect of activated-SS on the UCS of HM-contaminated soil: (a) four alkanolamines (TEA, TIPA, EDIPA
- 357

and DEIPA) at the same dosage of 0.05% and (b) 0.02%, 0.05%, 0.08% and 0.1% TIPA.

- 358 3.4 TCLP results of activated-SS treated HM-contaminated soil
- 359 *3.4.1 Effect of different alkanolamines*

360 The concentrations of leached HM, i.e., Cd, Cu, Ni, Pb and Zn in contaminated soils treated by 361 alkanolamine-activated SS were shown in Fig. 10. Apparently, the leachability of all the HM were 362 significantly reduced after S/S over time, which indicated that SS exhibited superior capacity in 363 solidifying and stabilizing the HM in the contaminated soil. With activators, the effectiveness of 364 HM stabilization was improved, while the type of activator did not seem to influence the early-age 365 leachability of HM significantly up to 7 days. Above 28 days, TIPA and EDIPA showed the best 366 performance among all the alkanolamines for the TCLP results. Moreover, it is worth noting that 367 0.5% of TIPA successfully lowered the leached concentrations of all the HM (except for Ni, which 368 barely exceeded the limit) to below their regulatory limits (see Standard for Pollution Control on 369 the Hazardous Waste Landfill (GB 18598-2019) [52]) in China after curing of 28 days.

Fig. 10 Effect of different alkanolamines (TEA, TIPA, EDIPA and DEIPA at 0.05% dosage) activated SS on the
leachability of HM: (a) Cd, (b) Cu, (c) Ni, (d) Pb and (e) Zn.

375 3.4.2 Effect of TIPA dosage

372

According to the above results, TIPA showed the highest effectiveness to solidify and stabilize HM
in contaminated soil. Thus, the effect of its dosage on the leachability of HM was investigated with
TCLP results shown in Fig. 11. All the HM showed decreased leached concentrations over time.
Consistent with the hydration and mechanical results, more TIPA showed improved HM
immobilization capacity due to the enhanced hydration of SS. At the dosage > 0.8% the leachability
of Ni also dropped to below its regulatory limit at 90 days.

382

Fig. 11 Effect of dosage of TIPA (0.02%, 0.05%, 0.08% and 0.1%) activated SS on the leachability of HM: (a) Cd,
(b) Cu, (c) Ni, (d) Pb and (e) Zn.

387 4. Discussion

Based on the experimental results presented above, TIPA was found to be the best activator among
the four types of alkanolamines, which was studied in detail. Hence the following sections will use
TIPA as the representative alkanolamine to elucidate its effect on SS hydration and soil S/S
performance.

392 *4.1 Effect on the hydration process*

The hydration process of TIPA-activated SS was characterized by the evolution of hydration heat and conductivity over time, and the results agreed well (see Figs. 4 & 5). The hydration process is hence divided into two stages. The first stage is the continuous dissolution of minerals and gradual accumulation of ions in the pore solution. TIPA effectively promotes the dissolution of SS, which was evidenced by the enhancement of the first exothermic peak of hydration heat and the first conductivity peak. This effect is more pronounced with the increased dosage of TIPA. Moreover, the dormant periods between the first and second peak (not observed in the reference SS samples 400 within the test time) in both the heat and conductivity evolution curves were significantly reduced. 401 indicating the fast accumulation of ions to achieve the supersaturations for hydration products, 402 which induced the second stage. The second stage is the precipitation of hydration products. The 403 enhancement of second peak of hydration heat and the increased difference between the peak and 404 residual conductivity showed that TIPA led to the formation of more hydration products. Similarly, 405 this effect is positively correlated with the dosage of TIPA. This was further evidence by the XRD 406 and TGA results. Hence, the strength of the SS pastes improved remarkably with the increase of 407 TIPA content (Fig. 8).

408 *4.2 Effect on the hydration products*

As shown in the XRD and TGA (see Figs. 6 &7), TIPA promoted the dissolution of minerals in SS
and the formation of hydration products. This effect is mainly exerted in two ways. The first one is
the promotion of the hydration of C₂S, aluminates and C₂F, resulting in the formation of C-S-H, CA-H and C-F-H. The other way is the enhanced reaction between C-A-H, C-F-H and CaCO₃, thereby
producing a large amount of Mc.

The mineral phases in SS first dissolve into ions, which accumulate in the cement pore solution (Equations 1-3), and they react with each other to form hydration products. The alkanolamines chelated with the metal ions, which increased the solubility of the minerals and the concentrations of ions [29, 42].

418
$$2(2CaO \cdot SiO_2) + 4H_2O \rightarrow 3CaO \cdot 2SiO_2 \cdot 3H_2O + Ca^{2+} + 2OH^-$$
 (1)

419
$$3CaO \cdot Al_2O_3 + 6H_2O \to 3Ca^{2+} + 2Al(OH)_4^- + 4OH^-$$
 (2)

420
$$2CaO \cdot Fe_2O_3 + H_2O + 2OH^- \rightarrow 2Ca^{2+} + 2Fe(OH)_6^{3-}$$
 (3)

The ions in the pore solution were chelated by TIPA and would not immediately participate in the precipitation reaction. According to Le Chatelier's principle, the forward progress of reactions (1)(3) were facilitated until a new value of the dissolution equilibrium constant K was reached [53].
Thus, TIPA promoted the dissolution of mineral phases and increased the amount of various dissolved metal ions in the solution. The precipitation reactions of ions are described in Equations
(4) - (8).

427
$$3Ca^{2+} + 2Al(OH)_{4}^{-} + 4OH^{-} \rightarrow 3CaO \cdot Al_{2}O_{3} \cdot 6H_{2}O$$
 (4)

19

$$428 \qquad 3Ca^{2+} + 2Fe(OH)_6^{3-} \to 3CaO \cdot Fe_2O_3 \cdot 6H_2O \tag{5}$$

$$429 \quad 3Ca^{2+} + 2Al(OH)_4^- + CaCO_3 + 4OH^- + 5H_2O \to 3CaO \cdot Al_2O_3 \cdot CaCO_3 \cdot 11H_2O \tag{6}$$

430
$$3Ca^{2+} + 2Fe(OH)_6^{3-} + CaCO_3 + 5H_2O \rightarrow 3CaO \cdot Fe_2O_3 \cdot CaCO_3 \cdot 11H_2O$$
 (7)

431
$$4Ca^{2+} + (2x)Al(OH)_{4}^{-} + [2(1-x)]Fe(OH)_{6}^{3-} + CO_{3}^{2-} + 4xOH^{-} + 5H_2O$$

$$432 \quad \rightarrow \ 3CaO \cdot xAl_2O_3 \cdot (1-x)Fe_2O_3 \cdot CaCO_3 \cdot 11H_2O \tag{8}$$

The precipitation led to the reduction of metal ion concentrations in the pore solution, resulting in the release of ions from their chelated forms, further promoting the precipitation of more hydration products. The dissolution and precipitation reactions are hence both promoted by TIPA which acts like a "catalyst" in the process. In summary, TIPA promoted the dissolution of mineral phases such as C₂S, C₃A, C₁₂A₇, and C₂F, thereby increased the amount of metal ions in the solution, and hence promoted the subsequent precipitation reaction of generating C-S-H, C-A-H, C-F-H and Mc. Nevertheless, how different alkanolamines interacted with the ions and affected the hydration of SS

440 differently remain unclear.

441 4.3 Effect on the S/S performance for HM-contaminated soil

442 TIPA-activated SS showed excellent compatibility with the heavily contaminated soil, which was 443 evidenced by the UCS of the mixtures (see Fig.9). Moreover, the leachability of all HM (except for 444 Ni) can be successfully reduced to below their regulatory limits with only 0.5% of TIPA in SS at the 445 curing age of 28 days, while Ni was successfully treated at 90 days with 0.8% TIPA-activated SS. 446 The outstanding S/S performance is partly due to the enhanced solidification due to better 447 cementitious properties of TIPA-activated SS and also due to the enhanced chemical stabilisation of 448 HM in the hydration products such as C-S-H, C-A-H, C-F-H and Mc [54-56]. Moreover, TIPA may 449 also play a role in stabilizing HM by chelation, which warrant further studies. Leaching effect of 450 HM in the hydration products of alkanolamines activated SS was shown in Fig. S1 and Fig. S2. The 451 leaching of each heavy metal remained relatively constant after the age of 28d, which demonstrates 452 that the solidification and stabilization of heavy metals by SS hydration products is firm and stable. 453 This study demonstrated that alkanolamines could improve the capacity of SS to solidify and 454 stabilize HM-contaminated soil. Firstly, it provides a sustainable alternative to Portland cement in 455 S/S to reduce energy consumption and carbon emission in cement production. Secondly, it will

456 enhance the utilization of SS to reduce the environmental impact and waste of land resources.

457 However, there are many challenges and opportunities to apply this novel binder in the field which

458 include but not limited to: (i) optimisation of the alkanolamines/SS and SS/soil ratios depending on

the type, concentration of the contaminants and the soil type; (ii) long-term performance of the

460 treated soils under different climatic conditions; and (iii) variability of the SS composition in the

461 world and hence the potential of leaching inherent contaminants in SS in the long term.

462 **5.** Conclusion

463 Four types of alkanolamines (TEA, TIPA, EDIPA and DEIPA) were used to activate SS and then 464 remediate HM-contaminated soil. The effects of alkanolamines on the hydration process and 465 hydration products of SS were thoroughly investigated via a series of techniques. Moreover, their 466 effects on the stabilization/solidification performance of HM-contaminated soil were assessed via 467 strength and leaching tests. The following conclusions could be drawn:

1. The addition of alkanolamines promoted the hydration process of SS and hence facilitated the formation of hydration products. The chelation of dissolved metal ions promoted the dissolution of mineral phases in SS and the subsequent precipitation reaction. The hydration products such as C-S-H, C-A-H, C-F-H and Mc were beneficial to solidify and stabilize heavy metal contaminated soil. 2. TIPA showed the best activation effect among all the investigated alkanolamines and better promotion effect could be obtained with higher dosage. Compared to the reference SS paste, the compressive strength of the 0.1% TIPA-activated SS was enhanced by 190.5% at 28 days.

475 3. TIPA-activated SS showed excellent performance in HM-contaminated soil 476 stabilization/solidification. Leached concentrations of Cd, Cu, Ni, Pb, and Zn have reduced by 477 87.2%, 78.8%, 62.4%, 73.6% and 64.5% using 0.1% TIPA-activated SS after 28 days, and they were 478 all below their respective regulatory limits by Standard for Pollution Control on the Hazardous 479 Waste Landfill (GB 18598-2019) in China. Compared to the reference SS, the UCS of the treated 480 soil at 28 days was enhanced by 237.7% using 0.1% TIPA-activated SS.

481 This study proposed a promising way of upcycling SS as a soil remediation agent to promote circular 482 economy, which would not only alleviate the environmental problems of disposing SS but also 483 provide a sustainable alternative to Portland cement in stabilization/solidification, contributing to a 484 low-carbon future. 485

486 **Declaration of Competing Interest** 487 We declare that we do not have any commercial or associative interest that represents a conflict of 488 interest in connection with the work submitted. 489 490 Acknowledgments 491 We are grateful to the financial supports by State Key Laboratory of Materials-Oriented Chemical 492 Engineering (No.SKL-MCE-22A07), National Natural Science Foundation of China (52272018), 493 National Key R&D Program of China (2021YFB3802002), a Project Funded by the Priority 494 Academic Program Development (PAPD) of Jiangsu Higher Education Institutions. 495 496 References 497 [1] B. Zeng, O. Wang, L. Mo, F. Jin, J. Zhu, M. Tang, Synthesis of Mg-Al LDH and its calcined form 498 with natural materials for efficient Cr(VI) removal, Journal of Environmental Chemical Engineering, 499 10 (2022). 500 [2] M.-K. Zhang, Z.-Y. Liu, H. Wang, Use of Single Extraction Methods to Predict Bioavailability of 501 Heavy Metals in Polluted Soils to Rice, Communications in Soil Science and Plant Analysis, 41 502 (2010) 820-831. 503 [3] Y.-L. Yang, K.R. Reddy, Y.-J. Du, R.-D. Fan, Sodium hexametaphosphate (SHMP)-amended calcium 504 bentonite for slurry trench cutoff walls: workability and microstructure characteristics, Canadian 505 Geotechnical Journal, 55 (2018) 528-537. 506 [4] Y. Yang, Reddy, KR, Du, YJ, Fan, RD, Short-Term Hydraulic Conductivity and Consolidation 507 Properties of Soil-Bentonite Backfills Exposed to CCR-Impacted Groundwater., JOURNAL OF 508 GEOTECHNICAL AND GEOENVIRONMENTAL ENGINEERING, 144 (2018). 509 [5] D. Hou, F. Li, Complexities Surrounding China's Soil Action Plan, Land Degradation & Development, 510 28 (2017) 2315-2320. 511 [6] Y. Song, N. Kirkwood, C. Maksimovic, X. Zheng, D. O'Connor, Y. Jin, D. Hou, Nature based 512 solutions for contaminated land remediation and brownfield redevelopment in cities: A review, Sci 513 Total Environ, 663 (2019) 568-579. 514 [7] C. Qu, W. Shi, J. Guo, B. Fang, S. Wang, J.P. Giesy, P.E. Holm, China's Soil Pollution Control: 515 Choices and Challenges, Environmental science & technology, 50 (2016) 13181-13183. 516 [8] Z. Shen, D. Hou, B. Zhao, W. Xu, Y.S. Ok, N.S. Bolan, D.S. Alessi, Stability of heavy metals in soil 517 washing residue with and without biochar addition under accelerated ageing, Sci Total Environ, 619-518 620 (2018) 185-193. 519 [9] Y.J. Du, M.L. Wei, K.R. Reddy, Z.P. Liu, F. Jin, Effect of acid rain pH on leaching behavior of cement 520 stabilized lead-contaminated soil, J Hazard Mater, 271 (2014) 131-140. 521 [10] D. Hou, A. Al-Tabbaa, D. O'Connor, Q. Hu, Y.-G. Zhu, L. Wang, N. Kirkwood, Y.S. Ok, D.C.W.

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