# **Removal of Pb<sup>2+</sup> and Cd<sup>2+</sup> from contaminated water using low-cost**

2

# materials

# 3 ABSTRACT

- 4 Heavy metals that exist in municipal wastewater can cause many problems for human hygiene and environment. 5 Therefore, it should be removed from wastewater before being used in irrigation. Materials of high surface reactivity; 6 such as alginit, shale and iron oxide are used as a potential sorbents to eliminate Pb and Cd from polluted water. In 7 remediation studies, these materials were added to Pb and Cd polluted water at addition ratios of 1:10000, 1:1000 and 8 1:100 (remedy agents: polluted water). The mixtures were then gently agitated and submitted to different equilibrium 9 periods of 1, 5 and 24 h. The results showed the efficiency of tested agents (shale, alginit, and iron oxide) in the remove 10 of Pb and Cd from polluted water containing various concentrations of 5, 10 and 50 mg/l. Shale was able to reduce Pb 11 and Cd concentration from 5 to 1.14 and 0.34 mg/l, respectively, in a reaction period of one hour. Shale, alginit and 12 iron oxide, reduced the initial concentration of; 10 mg Pb/l to 0.98, 0.46 and 0.57 mg/l; and of 50 mg Pb/l to 0.21, 6.5 13 and 1.68 mg/l; respectively. Shale was the most effective material in decontamination of heavy metals polluted water 14 and it could be recommended to be used to decontaminate wastewater. This research aims to use a non expensive, 15 environmentally safe, and efficient technique to remove heavy metals from industrial wastewater to leave them free and
- 16 suitable for discharging to sanitary sewer system.

17 Keywords: contaminated water; remediation, heavy metals; alternative low-cost materials

# Introduction

18 The most dangerous toxic elements listed by the European Economic Community on a "Black List", were 19 Hg and Cd, while the less dangerous substances forming the "Grey List" were Zinc, Copper, Nickel, 20 Chromium, Lead, Selenium, Arsenic, Antimony, Molybdenum and Titanium [1]. Cadmium is present in 21 wastewaters from metallurgical alloying, ceramics, electroplating, photography, pigment works, textile 22 printing, chemical industries and lead mine drainage. The application of phosphate fertilizers or sewage 23 sludge may increase cadmium levels in soil, which can cause increases in cadmium levels in crops [2]. The 24 average cadmium content of sea water is about 0.1ug/1 or less [4]. While river water contains dissolved 25 cadmium at concentration of < 1.1 - 13.5 ng/l, Cadmium levels of up to 5 mg/kg have been reported in river and lake sediments and from 0.03 to 1mg/kg in marine sediments [5], A drinking water guideline value of 26

Comment [c1]: This should be deleted

**Comment [c2]:** Replace this with "in the removal"

0.003 mg/l has been set for cadmium by WHO. In addition, the provisional tolerable weekly cadmium
intake must not exceed 7 µg/kg of body weight [6]. The guideline value for lead in drinking water given by
WHO is 0.01 mg/l [6].

The present study aimed to achieve an efficient non-expensive and environmentally safe method to decontaminate heavy metals (Pb and Cd) from polluted wastewater. Natural and non expensive materials, shale, alginit, and iron oxide, were used to decontaminate heavy metals polluted water to be suitable for discharging in drains and sanitary sewer system. At the end of remediation trails, the remedy agents can be removed easily, then recycled and utilized in manufacturing of building materials.

### 35 Material and methods

#### 36 Synthesized polluted water

Synthesized polluted water were prepared by accurate dilution of standard heavy metals solution of 1000 mg/l to known concentrations using distilled water. Lead polluted water was prepared using lead stock solution (1000 mg Pb/l in 0.5 M HNO<sub>3</sub> as matrix). A series of standard lead solutions of 5, 10, 50 and 100 mg Pb/l were prepared and used to test the ability of remedy agents in Pb removal. Cadmium polluted water was prepared using cadmium stock solution (1000 mg Cd/l in 0.5 M HNO<sub>3</sub> as matrix). A series of standard cadmium solutions of 5, 10, 20 and 50 mg Cd/l were prepared and used to test the efficiency of remedy agents in Cd removal.

#### 44 Remedy Agents

Three remedy agents, alginit, shale, and iron oxide, were used for polluted water remediation trials. These agents are common, inexpensive and easy to obtain. Besides, they could be separated easily from treated water at the end of the remediation process. These agents were selected based on their negatively charged surface and high adsorption capacity, which was attributed to their high surface area. The important Characteristics of the remedy agents are summarized in the following paragraphs.

- Alginit: Alginit is a natural rock out of the oil shale family. It originated from fossil algae biomass and
  pumice, descents from the mine in Gerce, Hungary. The essential ingredients of alginit [14] are high of
  organic matter (19%), clay (54%) and lime content (22%). The clay is rich in montmurillionte (52%).
- 53 Shale: Shale is a naturally occurring material exists in many places in Egypt at different depths. It mainly
  54 consists of clay (55%). The clay is rich in Montomorillonit. Chemical analysis showed that the shale

contains high amount of salt, the electrical conductivity (EC) of 1:2.5 water extract equals 10.63 dS/m and pH = 7.31. Sodium was the dominate cation.

57 Iron Oxide (60 % Fe): The sample of iron oxide is imported from Roseland Kazreti. The Chemical
58 composition of the iron Oxide is Fe<sub>d</sub> (60%), Fe<sub>o</sub> (19%), Al (0.16 mg/ kg), Zn (12.8 mg/kg) and Cu
59 (9.94mg/kg).

# 60 **Remediation studies**

Remedy agents of Alginit, shale, and iron oxide, were added to heavy metals polluted water at different solid: solution ratios of 1:10000, 1:1000 and 1:100. The mixtures were then gently agitated and subjected to different equilibrium periods of 1, 5 and 24 hrs. At the end of each equilibrium period, the supernatant solution was obtained by centrifuging the mixtures at 3000 rpm for 10 minutes. Concentrations of studied heavy metals (Pb and Cd) were measured before adding the remedy agent and at the end of equilibrium period as well as the pH and electrical Conductivity (Ec). All trials were done in three replicates.

# 67 Analytical procedures

Total concentrations of heavy metals (Pb, Cd, Ni, Co, Zn, Mn, Cu and Cr) were determined in both
suspended matter and clear water. Suspended matter was digested using Aqua Regia method [15].
Concentrations of Pb and Cd in polluted water as well as treated ones were determined using Atomic
Absorption Spectrophotometer (UNICAM, 969) (APHA, 1998). The pH was measured using digital Orion
pH meter (model 420A).

73 The electrical conductivity (EC) of the reacted solutions was measured using digital YSIEC meter (model 74 35), Cation exchange capacity (CEC) of remedy agents were determined using ammonium acetate method 75 as described by [16].Specific surface area of remedy agents were determined using O- phenanthroline 76 method [17].

# 77 Statistical analyses

- All statistical analyses were carried out by SAS version 9 software for all data of remediation trials. RSquared values (R<sup>2</sup>) and Equation were considered significant (*p*-values <0.05) for the analysis of variance</li>
  test (ANOVA).
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- 83 Results and discussion

Efficiency of remedy agents (shale, alginit and iron oxides) to remove Pb and Cd from synthetically polluted water containing various concentrations of 5, 10 and 50 mg/l were examined using different addition ratios (1:10000, 1:1000 and 1:100) of remedy agents: polluted water, at different reaction periods varying between 1 to 24 hour. The results obtained are as the follows.

88 Shale

89 The results (Table 1) showed that, although at the low addition ratio (shale: heavy metal polluted water) of 1: 10000 had no pronounced ability to remove Pb from polluted water, it was effective in removing Cd 90 91 from solutions containing low concentration of 5 mg/l. Shale reduced the initial concentration from 5 to 92 0.55 mg Cd/l in a reaction period of one hour. As the addition ratio increased to 1:1000, the removal 93 efficiency increased, particularly for Pb. Shale was able to reduce the initial concentration of Pb and Cd 94 from 5 to 1.14 and 0.34 mg/l, respectively, in one hour reaction time. As the addition ratio increased to 1: 95 100, the efficiency greatly increased. Shale successfully reduced the initial Pb concentration of 5 and 10 to 96 0.4 and 0.7 mg/l, in a reaction period of 1 hour which were lower than the permissible level (5 mg/l) for 97 irrigation water [18]. The corresponding values for Cd were 0.22 and 0.74 mg/l. Although these values are 98 much lower than the initial one, they were higher than the permissible level (0.01mg/l) for irrigation water. 99 Shale proved efficient in the remediation of higher concentration of 20 mg Cd /l at high addition ratio of 100 1:100. It reduces Cd concentration from 20 to 1.5 mg /l in a matter of 2 hour reaction time. Shale has 101 negligible ability to remove Pb and Cd from solutions containing high concentrations of 50 mg/l of Pb and 102 Cd. In general, the obtained results showed that shale has higher efficiency to remove Cd rather than Pb. 103 This efficiency could be attributed to high CEC values of shale (Table 2) and at the same time to the 104 character of Cd which is attracted to the negatively charged sites and exist in diffuse ion swarm as 105 exchangeable cation rather than forming inner sphere complex with surface functional groups which is a 106 characteristic of Pb.

**Comment [c3]:** Replace this with "The results obtained were as follows: "

107 Table 1. Concentrations of Pb and Cd in polluted water before and after treating with shale at addition ratios of 1:10000, 1:1000 and 1:100 for different reaction times.

Addition	Heavy metal	5(mg/l)			R <sup>2</sup>		10(mg/	)	R <sup>2</sup>		50(mg/l)		P <sup>2</sup>	LSD(5%) for
ratio	neavy meta	1(h)	5(h)	24(h)		1(h)	5(h)	24(h)		1(h)	5(h)	24(h)	ĸ	24h
1:10000	Pb	4.11	4.20	4.18	0.8450	8.10	7.70	9.24	0.8364	40.30	38.70	45.04	0.8422	0.26
1:10000	Cd	0.55	3.80	3.40	0.8465	8.80	5.40	9.20	0.9408	46.50	33.00	52.36	0.9202	0.22
1:1000	Pb	1.14	1.14	1.68	0.9735	2.70	2.60	4.40	0.9553	32.70	32.70	39.77	0.9735	0.30
1:1000	Cd	0.34	1.11	1.43	0.8793	5.00	3.00	4.40	0.9113	41.50	28.80	50.00	0.911	0.26
1:100	Pb	0.40	0.40	0.20	0.9735	0.70	0.51	0.98	0.9640	14.50	15.30	21.59	0.9964	0.32
1:100	Cd	0.22	0.21	0.07	0.9893	0.74	0.77	0.68	0.8607	17.80	13.60	14.00	0.9976	0.31

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# $116\quad$ Table 2. Cation exchange capacity (CEC) and surface area of remedy agents.

				Surface area							
		Remedy Agent	CEC {Cmol <sub>(c)</sub> /kg}								
				(m²/g)							
		Shale	60.13	165							
		Alginit	34.99	81							
		Iron oxide	7.49	150							
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121	The results (Table 1) showed that the efficiency of shale in removing Pb and Cd from polluted										
122	water	was higher at shorter <mark>equilit</mark>	prium period of one hour rather	than longer ones which could	be						
123	attribu	ted to the release of exchange	able cations initially existing in th	e interlayer of clay minerals, tl	hen						
124	replaci	ng Pb and Cd readily excha	nged to the surface. So that, hig	her reaction time of 24 h is	not						
125	recom	mended for shale. Similar trails	s have been done [20] using alkali-t	reated oil shale ash as adsorben	t to						
126	remove	e lead and cadmium ions from	aqueous solutions. They reported th	at adsorption of lead and cadmi	um						
127	ions by	y the modified oil shale ash de	pended on adsorbent concentration,	, ash particle size, contact time a	and						
128	pH of solution. At initial concentration of an aqueous solution of 10mg/L and that of the adsorbent 5g/L,										
129	91% of lead and cadmium ions was removed from the solution. These results proved lower efficiency of										
130	the modified oil shale ash compared with the shale sample utilized in our research										
131	Alginit										
132	The lowest addition ratio of 1:10000 was not effective even for low concentration of 5 mg Pb/l. The higher										
133	addition ratio of 1:1000 (Table 3) was effective only for relatively low concentration of 5mg /l, which is										
134	considerably reduced to 0.52 mg/l in a reaction time of 24 h. The efficiency of alginit in remediation of Pb										

polluted water increased as the addition ratio increased. Addition ratio of 1:100 significantly eliminates Pb

**Comment [c4]:** This should be replace with " These results proved of lower efficiency when the modified oil shale ash was compared with " from polluted water containing 5 and 10 mg Pb/l in a reaction time of 24h, Pb concentration in the previous solutions reduced to 0.21 and 0.46 mg/l, respectively. For water highly polluted with 50 mg Pb /l, addition ratio of 1:100 was able to reduce Pb to concentration (6.5 mg/l) little higher than the permissible level.

139 Regarding Cd, addition ratios of 1:10000 and 1:1000, gave low efficiency in removing Cd from polluted 140 water. However, addition ratio of 1:100 proved more efficient in eliminating Cd from polluted water. It 141 reduces the initial Cd concentrations of 5 and 10 to 0.21 and 1.57 mg/l, respectively. The results showed 142 that, removal of Cd by alginit was time dependent since the changes in pollutant concentrations with time 143 were significant (LSD<sub>0.05</sub> ranged from 0.20 - 0.33). Based on Cd permissible level for irrigation water (0.01 144 mg/l), alginit showed low efficiency in Cd removal. It works only with low Cd concentration at high 145 addition ratio of 1:100. Based on these results, it could be concluded that, unlike shale, alginit has higher 146 efficiency in the removal of Pb rather than Cd, which could be attributed to the relatively low CEC value 147 (34.99 Cmol (c/kg) of alginit (Table 2) comparing with shale (60.13 Cmol(c) /kg), i.e. low electrostatic 148 attraction between Cd and alginit surface which led to low Cd removal via exchange process. Therefore, 149 specific adsorption of Pb is more likely to be dominant on alginit surface than the exchange process

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151 Table 3. Concentrations of Pb and Cd in polluted water before and after treating with alginit at addition ratios of 1:10000, 1:1000 and 1:100 for different reaction times

Heering	5(mg/l)					10(mg/l)				50(mg/l)		LSD	
metal	1(h)	5(h)	24(h)	R <sup>2</sup>	1(h)	5(h)	24(h)	R <sup>2</sup>	1(h)	5(h)	24(h)	- R <sup>2</sup>	(5%) for 24 h
Pb	5.00	4.73	4.70	0.8733	10.00	9.50	9.50	0.8018	40.10	40.80	40.30	0.9985	0.26
Cd	5.00	5.00	5.00	Nd	8.90	10.00	5.50	0.8483	47.40	46.50	39.60	0.9967	0.32
Pb	1.85	1.2	0.52	0.8843	9.01	8.10	9.68	0.9117	40.20	39.70	38.50	0.9352	0.20
Cd	2.80	4.40	3.20	0.9691	9.10	8.00	5.20	0.9872	50.00	48.90	50.00	0.9859	0.33
Pb	0.04	0.21	0.21	0.8803	0.40	0.42	0.46	0.9868	19.80	17.90	6.50	0.9990	0.31
Cd	0.29	0.23	0.21	0.2845	2.00	1.70	1.57	0.9307	47.40	39.90	30.00	0.9233	0.31
	Heavy metal Pb Cd Pb Cd Pb Cd	Heavy metal         1(h)           Pb         5.00           Cd         5.00           Pb         1.85           Cd         2.80           Pb         0.04           Cd         0.29	Heavy metal         5(mg/l)           Pb         5.00         4.73           Cd         5.00         5.00           Pb         1.85         1.2           Cd         2.80         4.40           Pb         0.04         0.21           Cd         0.29         0.23	Heavy metal         5(mg/l)           1(h)         5(h)         24(h)           Pb         5.00         4.73         4.70           Cd         5.00         5.00         5.00           Pb         1.85         1.2         0.52           Cd         2.80         4.40         3.20           Pb         0.04         0.21         0.21           Cd         0.29         0.23         0.21	5(mg/l)           Heavy metal         5(mg/l)         24(h)         R <sup>2</sup> Pb         5.00         4.73         4.70         0.8733           Cd         5.00         5.00         5.00         Nd           Pb         1.85         1.2         0.52         0.8843           Cd         2.80         4.40         3.20         0.9691           Pb         0.04         0.21         0.21         0.8803           Cd         0.29         0.23         0.21         0.2845	5(mg/l)           Heavy metal         5(h)         24(h)         R <sup>2</sup> 1(h)           Pb         5.00         4.73         4.70         0.8733         10.00           Cd         5.00         5.00         5.00         Nd         8.90           Pb         1.85         1.2         0.52         0.8843         9.01           Cd         2.80         4.40         3.20         0.9691         9.10           Pb         0.04         0.21         0.21         0.8803         0.40           Cd         0.29         0.23         0.21         0.2845         2.00	5(mg/l)         10(mg/l)           Heavy metal $1(h)$ $5(h)$ $24(h)$ $R^2$ $1(h)$ $5(h)$ Pb $5.00$ $4.73$ $4.70$ $0.8733$ $10.00$ $9.50$ Cd $5.00$ $5.00$ $5.00$ Nd $8.90$ $10.00$ Pb $1.85$ $1.2$ $0.52$ $0.8843$ $9.01$ $8.10$ Cd $2.80$ $4.40$ $3.20$ $0.9691$ $9.10$ $8.00$ Pb $0.04$ $0.21$ $0.21$ $0.8803$ $0.40$ $0.42$ Cd $0.29$ $0.23$ $0.21$ $0.2845$ $2.00$ $1.70$	Heavy metal $5(mg/l)$ $R^2$ $10(mg/l)$ Pb $5(h)$ $24(h)$ $R^2$ $1(h)$ $5(h)$ $24(h)$ Pb $5.00$ $4.73$ $4.70$ $0.8733$ $10.00$ $9.50$ $9.50$ Cd $5.00$ $5.00$ $5.00$ Nd $8.90$ $10.00$ $5.50$ Pb $1.85$ $1.2$ $0.52$ $0.8843$ $9.01$ $8.10$ $9.68$ Cd $2.80$ $4.40$ $3.20$ $0.9691$ $9.10$ $8.00$ $5.20$ Pb $0.04$ $0.21$ $0.21$ $0.8803$ $0.40$ $0.42$ $0.46$ Cd $0.29$ $0.23$ $0.21$ $0.2845$ $2.00$ $1.70$ $1.57$	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c c c c c c c c c c c c $

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# 157 Iron oxide

158 Table (4) shows the concentration of Pb and Cd in polluted water before and after treating with iron oxide 159 at additio ratios of 1:10000, 1:1000 and 1:100 for different reaction times. At the lowest addition ratio of 160 1:10000, iron oxide proved not efficiency to remove Pb even from water polluted with low concentrations 161 of 5 mg/l. As the addition ratio increased to 1:1000, iron oxide worked well with low Pb concentration of 5 162 mg /l and reduced the initial concentration to 0.67 mg/l. However, for water polluted with higher Pb 163 concentration of 10 mg/l, the efficiency dropped, in which the initial concentration reduced to be 3.5 mg/l 164 in a reaction time of 24 hour. Iron oxide can be potentially efficient in removing Pb from polluted water 165 only at high addition ratio of 1:100. This efficiency increased even for high Pb concentration of 50 mg/l. 166 Also the removal efficiency increased significantly ( $R^2$ =0.888) as the reaction time increased. In a reaction 167 time of 24 hour, it eliminated Pb from solutions of initial concentration of 5 and 10, with the concentrations 168 reduced to undetectable value (Table 4). Also, it reduced the concentration of 50 mg Pb/l to 1.67 mg/l 169 where is lower than the permissible level (5 mg/l) for irrigation water.

170 The efficiency of Fe-oxide in removing Cd was significantly (LSD<sub>0.05</sub>=0.24-0.32) time dependent. As the 171 reaction period increased, Cd concentration in the equilibrium solution decreased. High addition ratio of 172 1:100 was the most effective in removing Cd. At addition ratio of 1:100, iron oxide efficiently reduced the 173 initial concentrations of 5 and 10 mg Cd/l to 0.1 and 0.4 mg/l, respectively, in a reaction period of 24 h. 174 These concentrations (0.1 and 0.4 mg/l) were higher than the permissible level (0.01 mg/l) for irrigation 175 water. Although, several methods have been adopted to remove heavy metals from polluted water, these 176 methods succeeded only with water of high pH values and low concentrations of pollutants. At acidic 177 conditions and relatively high concentrations of pollutants, the efficiency of these methods were limited. 178 On the other hand, the techniques utilized in our research succeeded in decontaminate water of low pH 179 values and high concentrations of Pb and Cd. Among these methods [21] stated, that waste iron 180 (III)/chromium (III) hydroxide has been used as an adsorbent for the effective removal of Pb(II) from 181 aqueous solution at pH greater than 7.0. The percent adsorption of Pb(II) increased with a decrease in 182 concentration of Pb(11) and an increase in temperature. [22], suggested a process for removing lead from 183 battery industry wastewater by neutralization with NaOH, in the presence of Fe(III) salts which the lead concentration of the treated effluent is below O.2 mg/l. [23], proposed a procedure for purifying waters 184

**Comment [c5]:** Please replace this with "efficient in the removal of "

185	polluted	with	metal	ions	[	Al(I1I),	V(V),	Ni(II),	Cd(II),	Co(II),	Pb(II),	Hg(II),	Cr(II),
186	Sn(II),Bi(	II),Zn(	(II),and (	Cu(II)],	by	precipitat	tion of m	ietals as	magnetic	ferrite fro	om the all	kalinized	solution
187	containin	g iron(	II) was l	based o	on tl	he precipi	tation of	metals a	as magnet	ic ferrite.	The may	kimal pur	ification
188	efficiency	(99.9	9%) was	s achiev	ved	when wa	ste wate	r sample	s are treat	ed for 3	hours at	50°C and	pH 10,
189	Fe(1I)/To	tal	metal	ratio		was 1	5.0 a	und d	ifferent	concent	rations	of	KMnO <sub>4</sub> .

192	Addition	Норм	5(mg/l)				10(mg/l)				50(mg/l)				LSD
193	ratio	metal	1(h)	5(h)	24(h)	R <sup>2</sup>	1(h)	5(h)	24(h)	R <sup>2</sup>	1(h)	5(h)	24(h)	- R <sup>2</sup>	for 24h
194	1:10000	Pb	2.76	5.00	5.00	0.851	10.10	10.00	10.00	0.906	42.60	10.78	44.04	0.853	0.25
	1:10000	Cd	4.50	2.60	3.00	0.984	0.30	10.40	8.00	0.850	47.10	52.80	42.00	0.926	0.25
	1:10000	Pb	2.76	2.60	0.67	0.991	9.60	8.53	3.50	1.000	42.60	39.82	44.83	0.879	0.29
	1:10000	Cd	3.70	2.00	2.00	0.842	2.10	9.20	8.80	0.796	49.80	49.20	49.40	0.962	0.24
	1:100	Pb	0.27	Nd	Nd	0.402	0.20	0.28	Nd	0.815	29.01	15.80	1.68	0.888	-
	1:100	Cd	0.70	0.34	0.10	0.790	4.00	2.40	0.40	0.915	38.80	44.00	34.00	0.880	0.32

Table 4. Concentrations of Pb and Cd in polluted water before and after treating with iron oxide at addition ratios of 1:10000, 1:1000 and 1:100 for different 190

reaction times. 191

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210	Conclusion	
211	All remedy agents, shale, alginit and iron oxides, proved potential efficiency in the removal of Pb and Cd	<b>Comment [c6]:</b> Replace this with "proved potentially efficient"
212	from water polluted of wide range of Pb and Cd varied between 5 to 50 mg/l. Generally, their efficiency	
213	increased as the addition ratio between remedy agents and polluted water increased from 1:10000 - 1:100.	
214	Among all tested agents, shale had highest efficiency for the removal of Cd. It had high potential ability to	<b>Comment [c7]:</b> Replace this with "had the"
215	remediate higher concentration of 20 mg Cd /l at addition ratio of 1:100. However, shorter equilibrium	
216	period of 1 hour was more effective than the higher one of 24 h. Alginit proved high efficiency in removing	
217	Pb and Cd from polluted water when added at high ratio of 1:100. Unlike shale, alginit had a higher	
218	removal efficiency for Pb rather than Cd. Iron oxide had potential efficiency in removing Pb from polluted	
219	water only at high addition ratio of 1:100. This efficiency was shown even for high Pb concentration of 50	

- 220 mg/l. In a reaction period of 24 hour, it eliminated Pb from solutions of initial concentration of 5 and 10,
- 221 mg/l, while reducing the concentration of 50 to a level (1.67 mg/l) lower than irrigation water permissible
- 222 level (5 mg/l).
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