Al-Substituted Tobermorites: An Effective Cation Exchanger Synthesized from “End-of-Waste” Materials

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ABSTRACT: The policies to meet the “zero waste” regime and transition to sustainable circular economy can no longer ignore the use of wastes in place of natural resources, and these daunting and vital societal challenges are nowadays being faced by several nations. The main objective of this work was to search waste materials suitable for a quick and environmentally friendly production of a nanoporous geomaterial able to trap toxic metals at the solid/liquid interface. More specifically, the end-of-waste from the thermal inertization of cement−asbestos and glass powder from domestic glass containers have been employed as sources for the hydrothermal synthesis of a tobermorite-rich material (TRM) successfully tested for the selective removal of Pb²⁺, Zn²⁺, Cd²⁺, and Ni²⁺ from aqueous solutions. The synthesis was carried out in alkaline solution under mild hydrothermal conditions (120 °C) within 24 h. The quantitative phase analyses of the TRM carried out by applying the Rietveld method showed the occurrence of a large amount of well-crystallized 11 Å Al-substituted tobermorites and an amorphous phase and a lower content of aragonite and calcite. Chemical analyses and thermogravimetric measurements coupled with simultaneous evolved gas mass spectrometry highlighted that Al³⁺ for Si⁴⁺ substitutions in the wollastonite-like tetrahedral chains of tobermorites are balanced by the occurrence of Ca²⁺, Na⁺, and K⁺ cations in the interlayer rather than by (OH)⁻ for O²⁻ substitutions in the CaO polyhedra. Time-dependent removal tests clearly indicated that metal cations are selectively adsorbed depending on their concentration in solution. Moreover, the kinetic curves showed that the removal of metals from solution is fast and the equilibrium is almost reached in the first 8 h.

1. INTRODUCTION

One of the main challenges for modern society is to reduce the production of waste and limit the consumption of natural raw materials (NRM) whose availability is increasingly difficult and costs are increasing. At once, the growing awareness for the “environmental issues” has led to the development of environment-friendly, sustainable, and cost-effective industrial processes and, in several countries, new production activities can be launched only if fully integrated within the circular economy regime. Nevertheless, these goals are frustrated by the fact that, for a long time, humans will have to live with the hazardous waste produced in the past and the nonhazardous waste generated at a faster rate than their recycling. Two typical examples are asbestos and glass from the recycling of domestic containers.

In countries where asbestos is banned, massive amounts of asbestos-containing materials, especially cement−asbestos slates (CAS), must be removed and disposed. Disposal in landfills cannot be the ultimate solution not only as it conflicts with the directives of the European Commission (2008/98/EC, end-of-waste concept) but also as zero risk of fiber dispersion in air and water in the long run cannot be guaranteed. A very promising alternative to landfill storage is the inertization achieved through thermal treatment as firing of CAS yields an inert fine material, generally referred to as KRY-AS. KRY-AS is an end-of-waste and has been proposed as a replacement for NRM in various applications depending on its mineralogical and chemical composition, demonstrating that the thermal inertization and recycling of the product of transformation are an effective solution to the asbestos problem.

The separate collection of domestic glass containers (DGC) is a political obligation within many nations and it has progressively increased in the last decades as demonstrated by several national reports. Nevertheless, their recovery is
Table 1. Elemental Composition (mol/100 g) and Loss on Ignition (LOI, Weight %) at 1100 °C of KRY·AS, Glass Powder, Zeolitic Tuff, KGT, and TRM:

|       | KRY·AS       | glass powder | zeolitic tuff | KGTb  | KGTc  | KGTd  | TRMb | TRMd |
|-------|--------------|--------------|--------------|-------|-------|-------|------|------|
| Si    | 0.4139       | 1.1538       | 0.8778       | 0.6562| 0.6634| 0.6870| 0.5715| 0.7065|
| Al    | 0.0798       | 0.0557       | 0.3328       | 0.0771| 0.0759| 0.0786| 0.0610| 0.0754|
| Fe    | 0.0341       | 0.0059       | 0.0482       | 0.0255| 0.0262| 0.0271| 0.0207| 0.0256|
| Ti    | 0.0026       | 0.0018       | 0.0054       | 0.0024| 0.0025| 0.0026| 0.0019| 0.0023|
| P     | 0.0000       | 0.0003       | 0.0008       | 0.0001| 0.0001| 0.0001| 0.0001| 0.0002|
| Mn    | 0.0010       | 0.0007       | 0.0021       | 0.0009| 0.0011| 0.0012| 0.0007| 0.0009|
| Mg    | 0.2363       | 0.0620       | 0.0270       | 0.1772| 0.1777| 0.1840| 0.1526| 0.1887|
| Ca    | 0.8622       | 0.1648       | 0.0794       | 0.6269| 0.6123| 0.6444| 0.5359| 0.6626|
| Na    | 0.0083       | 0.3957       | 0.0323       | 0.1308| 0.1384| 0.1330| 0.1591| 0.1967|
| K     | 0.0062       | 0.0164       | 0.1232       | 0.0118| 0.0121| 0.0392| 0.0098| 0.0121|
| LOI   | 9.65         | 2.42         | 13.94        | 7.46  | 6.09  | 0.00  | 19.11 | 0.00  |

*TRM chemical composition is listed here to facilitate the comparison with source materials. bCalculated. cExperimental. dExperimental recalculated as anhydrous.

often frustrated by various aesthetic and technical hindrances which, sometimes, leave landfill disposal as the only solution.15–18 Typical limiting factors are color heterogeneity, high calcium concentration, not being compatible with fiberglass production, occurrence of metals (in several countries, DGC are collected together with iron and aluminum cans), and, often, the anomalous concentration of lead due to the wrong disposal of crystalware in separate collection which drastically impacts the possible reuse applications.17,19

Although the production of aggregates, concretes, ceramics, and abrasives represents an alternative market with few activities,53 and they are mainly composed of calcium carbonate and calcium silicates (KRY·AS) and by amorphous materials with a small amount of crystalline calcium carbonate and metal aluminum (glass powder).

As mentioned, KRY·AS is the product of the thermal inertization of CAS performed in industrial kilns at temperatures in the range of 1200–1300 °C. The obtained product is free of asbestos phases and contains newly formed clinker
phases with cytotoxic effects comparable to those of commercial clinker. More specifically, KRY-AS is composed of subospherial particles of the newly formed crystalline phases but with an excess of aluminum, iron, and magnesium and a compositional variability mostly depending on the provenience of the CAS. Glass powder is from the sleeve filters of the mills used to grind glass from domestic recycling. It should be emphasized that due to the small grain size (less than 50 μm) and the occurrence of a small amount of Mg and Fe, the recovery options are quite limited, and this powder is generally considered a waste.

2.2. Synthesis Procedure. Al-substituted tobermorites can be successfully obtained in a simulated hydrothermal environment from materials whose composition approximately falls within the molar ratios 0.80 < Ca/[Si + Al + Fe] < 0.85 and 0.00 < [Al + Fe]/[Al + Si + Fe] < 0.17.51,52 Even if various approaches have been attempted in the past to optimize the reaction mechanism mainly by varying the reaction time, temperature, and reagents (see the literature cited in the Introduction), to date, the synthesis of a pure form of tobermorite has not yet been obtained. The optimization of the synthesis parameters to gain the maximum amount of tobermorite in the final product goes beyond the aim of this research as our primary goal is maximizing the quantity of waste recovery as opposed to lower energy consumption. The hydrothermal synthesis was carried out in a Teflon-lined Parr bomb by adding 15 mL of NaOH 1 M solution to 1 g of a phillipsite was found to be a catalyst for the synthesis of waste recovery as opposed to lower energy consumption. The tobermorite in the mixture (KGT, Table 1) prepared by dosing KRY bomb by adding 15 mL of NaOH 1 M solution to 1 g of a phillipsite, as it was introduced as a solid phase as its concentration increases in the TRM with respect to KGT. On the contrary, the concentration of the other elements, likely involved in the chemical substitutions in tobermorite, has not changed significantly, suggesting that they contributed almost completely to the formation of crystalline phases or precipitated as amorphous materials rather than remaining in the solution and being removed when the solid is separated from the liquid after the synthesis.

The XRPD measurements of the TRM (Figure 1) showed the occurrence of 11 Å-tobermorite, aragonite, and calcite. The basal spacing defined by the (002) reflection plane in tobermorite is 11.43 Å, thus in good agreement with values previously reported for Al-substituted tobermorites.39,48 There is no evidence of the occurrence of 9 Å- and 14 Å-tobermorites as the main reflections are lacking and/or mismatched. The model from Mitsuda and Taylor (1975) was applied to estimate the crystallinity rate of the synthesized tobermorite. This model defines four levels of crystallinity in Al-substituted tobermorites based on the occurrence and shape of the (002),
large amount of 11 Å-tobermorite and a lesser content of aragonite and calcite. Moreover, the addition of a standard to the raw sample powder to apply a combination of the Rietveld method and using as starting structural models for the refinement those from Yamazaki and Toraya (2001) for Al-substituted 11 Å-tobermorite (COD code 1527001),64 Villiers (1971) for aragonite (COD code 5000085),65 and Sitepu (2009) for calcite (COD code 9016022).66 The QPA results (Table 2 and Figure S2) better detailed the occurrence of a refinement (9.9 wt %, sum of aragonite and calcite, Table 2). Likewise, the moles of Ca hosted in the two carbonates (0.09891 and 0.1219 mol/100 g from QPA and TGA, respectively) can be calculated from the mineral formula. Reactions that involve the release of N (m/z = 30) and SO2 (m/z = 64) were not observed, and the absence of nitrogen and sulfur was confirmed through EA.

The time-dependent removal of metal cations (Figures 3 and S3) was evaluated considering the dependence between the concentration of the elements in solution at room temperature and the unit mass of the adsorbent (TRM). The results shown in Figure 3 clearly indicated that cations are selectively removed and the higher the concentration of metals in solution, the greater the selectivity (Figure 3a–c). Pb2+ is the cation that is preferentially removed followed, in order, by Zn2+, Cd2+, and Ni2+. The removal of Cd2+ is comparable to that of Ni2+ only in the batches prepared with the solutions at concentrations 1 and 5 mmol/L (Figure 3d,e, respectively) while that of Zn2+ and, mostly, of Pb2+ remains high and selective. The kinetic curves shown in Figure 3 also showed that the removal takes place fairly and quickly and the equilibrium is almost reached in the first 8 h (only in the 100 and 50 mmol/L batches, the saturation is reached just faster than in the others). The representation by the weight % of the total amount of toxic metals removed by the TRM may be relevant for applicative purposes and is shown in Figure 3f. This analysis highlights how the overall adsorption depends on
the concentration of the solution, and it varies between 10 and 70 wt% for the highest (100 mmol/L) and lowest (1 mmol/L) concentrations, respectively. An analogue representation for each batch is shown in Figure S3.

4. DISCUSSION

The experimental results presented in this work clearly indicate that it is possible to synthesize tobermorite using end-of-waste materials. The whole compositional range for 11 Å-tobermorite, without accounting Al³⁺ and Fe³⁺ for Si⁴⁺ substitutions, spans between Ca₄Si₆O₁₅(OH)₂·5H₂O when...
no zeolitic Ca occupies the interlayer and Ca$_x$Si$_y$O$_z$·nSH$_2$O when one Ca cation per unit formula is hosted in the interlayer space.\textsuperscript{57} Based on the goodness of fit of the Rietveld refinement (Table 2 and Figure S2) achieved using an Al-substituted tobermorite as the starting model,\textsuperscript{64} it can be argued that in TRM tobermorite, Si$^{4+}$ is partly substituted by Al$^{3+}$ and, likely, by Fe$^{3+}$. This conclusion is in agreement with the chemical data as in the TRM, the [Al + Fe]/[Si + Al + Fe] molar ratio of 0.125 is in the optimal range of 0.00–0.17, typical of substituted tobermorite, as well as the [Al]/[Al + Si] molar ratio of 0.097 is below the maximum value of 0.13–0.14.\textsuperscript{10,14,53,68} In TRM tobermorite, Al$^{3+}$ and Fe$^{3+}$ for Si$^{4+}$ substitutions are balanced by the occurrence in the interlayer of Ca$^{2+}$ and, likely, Na$^+$ and K$^+$ cations rather than by (OH)$^−$ for O$^{2−}$ substitutions, as suggested by the absence of thermal events related to a dehydroxylation reaction which should occur between 550 and 650°C.\textsuperscript{57} The contribution of Na$^+$ and K$^+$ to the charge balance is necessary as a nonnegligible amount of Ca (0.1104 mol/100 g, calculated averaging the concentration values from QPA and TGA) is hosted in the calcite and aragonite lattice. Therefore, the charge deficiency provided by Al$^{3+}$ and Fe$^{3+}$ for Si$^{4+}$ substitutions (0.3030 equiv/100 g, Table S1) cannot be neutralized only by exchangeable Ca$^{2+}$ in tobermorite (ideally 1/5, thus 0.2209 equiv/100 g in TRM tobermorite). Nevertheless, these considerations are based on the constraint that in the amorphous phase there are no compounds with a tobermorite-like structure.

The crystallization of carbonates, as well as that of other compounds such as C–S–H phases (the latter being not found in the TRM), could sound unusual; nevertheless, it has already been observed in other similar synthesis experiments.\textsuperscript{52,70–73} More specifically, the crystallization of calcite and aragonite is due to progressive and time-dependent carbonation reactions and here, it could be favored by the occurrence of calcite in the source materials which could have acted as a catalyst.\textsuperscript{70} The high amount of the amorphous phase is reasonably due to the short times used for the synthesis (although, as mentioned, the synthesis of a pure form of tobermorite has never been reported in the literature to date) and to the presence in the source materials (mostly in KRY–AS) of huge amounts of Mg that can hardly be incorporated into the structure of tobermorite. In contrast, the short time of synthesis does not affect the crystallinity of the tobermorite phase.

The adsorption curves (Figure 3) showed that the TRM, like other synthetic materials with similar mineralogical composition, may remove toxic metals from aqueous solutions. Nevertheless, metal removal must occur not only by cation-exchange reactions but also by surface adsorption. In fact, with the exception of the 1 mmol/L batch, the charge equivalents of metal cations removed are higher than those related to exchangeable Ca$^{2+}$, Na$^+$, and K$^+$ in the TRM (Table S2). Therefore, inner- and outer-sphere bonding may occur either with the exposed oxygens of tobermorite and the hydroxyls forming the amorphous phase.\textsuperscript{74,75} This hypothesis is confirmed by considering the small increase (between 0.1 and 0.3) of pH observed at the end of each treatment and attributed to zeolitic Ca$^{2+}$, Na$^+$, and K$^+$ cations released by tobermorite (metal cations, in view of the partial deprotonation of their aqua complexes, are more acidic). The adsorption rate of the 100 and 50 mmol/L solutions is very close (only a little faster) to that observed for the other concentrations. Although there is no direct experimental evidence, it is possible that this atypical behavior is due to the occurrence of unsaturated negative charges (e.g., those related to amorphous materials) which affects the adsorption process. Regardless of all the considered metals being divalent, the cation-exchange reactions should first involve the release of Na$^+$ which has less affinity for the tobermorite interlayer.\textsuperscript{39,76}

The high affinity of 11 Å-tobermorite toward toxic metals had been observed before,\textsuperscript{39,53,59,63} but the competitive adsorption from multielement solution had rarely been verified.\textsuperscript{59} In the case of the TRM, it should be stressed that the markedly selective removal of metal cations is not opposed by fluctuating time-dependent adsorption paths. Indeed, with the exception of a slight decrease in the removal rate of Cd$^{2+}$ and Ni$^{2+}$ between 16 and 32 h in the 100 mmol/L batch, the equilibrium conditions, once reached, are preserved over a time span of almost 100 h. This behavior distinguishes the TRM from other materials such as zeolitized tufts and clay minerals where consecutive adsorption/desorption reactions may occur over time.\textsuperscript{77–79}

In recent years, waste recovery and exploitation have become options playing a strategic role in drastically reducing final disposal in landfills while ensuring increasing quantities of secondary raw materials. Currently, the main interest for tobermorite is due to its occurrence in cements as it improves thermal and acoustic insulating properties. On the other hand, the use of tobermorite as an adsorbent does not yet have a significant impact in large-scale applications, probably due to the availability of NRM (mainly bentonites and zeolitic tufts) with similar, or even better, adsorption performance. However, it should be emphasized that tobermorite-rich materials (like the TRM but also others mentioned in the Introduction) have a CEC close to that of natural bentonites (i.e., ranging approximately between 60 and 120 meq/100 g) and just below that of several zeolitic tufts (i.e., 100–200 meq/100 g).\textsuperscript{80–82} Even without a detailed market analysis, it is reasonable to assume that the large-scale production of tobermorite using only pure reagents would be impractical as it would cost significantly more than the mining and trade of NRM. Nevertheless, it is also true that the use of NRM has a high impact on the environment (opening/expanding of quarries is always a hotly debated issue), especially if this is countered by a possible green solution to gain materials with similar properties. In addition, it is also conceivable that the environmental benefits of replacing NRM with waste-derived materials outweigh the overall impact of the collection, sorting, transport, and recycling of waste, and the cost of these operations can be recovered by the sale of the synthesized materials.

5. CONCLUSIONS

The main objective of this work was to test KRY–AS and DGC as possible candidates for a fast and environmentally friendly synthesis of a microporous material to be used for the removal of toxic metals from aqueous solutions. Although the tests were carried out at the laboratory scale, we believe that the transfer to a larger scale should not meet major impediments, either in relation to the simple preparatory procedure and the mild conditions under which the synthesis is carried out.

Despite the relevant properties of the TRM, we are aware that the method described here addresses two obstacles that may slow down its large-scale application: (i) the chemical composition of the waste which may vary depending on the source and (ii) the occurrence of NRM with similar properties
available at lower costs. The former, however, is of little impact as it is sufficient to calculate the amount of each waste to meet the synthesis protocol. On the other hand, the availability of analogue NRM (e.g., bentonites and zeolitic tuffs) is a more sensitive issue and, to be critically commented, would require a thorough analysis of the cost-benefit ratio, also properly assessing the overall environmental impact.

**ASSOCIATED CONTENT**

*Supporting Information*

The Supporting Information is available free of charge at [https://pubs.acs.org/10.1021/acsomega.1c04193](https://pubs.acs.org/10.1021/acsomega.1c04193).

Details about analytical methods and sample preparation before measurements, instruments, experimental conditions and data analyses; sketch of the tobermorite structure; graphical results of the Rietveld refinement of the TRM; time-dependent metal cation removal in weight %; and chemical analyses of the TRM considering the repartition of Ca between tobermorite and the sum of calcite and aragonite (PDF).

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**Notes**

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