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In situ synchrotron powder diffraction study of Cd intercalation into Chevrel phases: crystal structure and kinetic effect

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Abstract

Chevrel phases are molybdenum chalcogenides of formula MₓMo₆X₈ (M is a cation and X a chalcogen) that present a complex and captivating intercalation chemistry that has drawn the interest of the solid state chemistry community since their discovery1–3. This property has a huge potential for applied science and device development for energy storage4, pollutants removal or detection5,6, but a deeper knowledge of the intercalation processes and chemistry is still necessary. In the present work, the intercalation of Cd²⁺ in aqueous solution has been studied taking advantage of the complementarity of electrochemistry characterization and synchrotron powder diffraction, acquired during an in situ combined experiment. During the experiment, industrially adequate electrochemical conditions (room temperature and reduced process time) have been applied, allowing a better understanding of the intercalation processes. The intercalated phases obtained by electrochemistry have been characterized ex situ, and for the first time, the structures of CdₓMo₆X₈ (X=S, Se) have been determined.
Unexpectedly, Cd₂Mo₆Se₈ presents a trigonal crystal structure with only cavity 2 occupied, which has not been encountered before for Chevrel phases.

**KEYWORDS**

Chevrel phases, Cd removal, powder diffraction, electrochemistry

**Introduction**

MₓMo₆X₈ and the binary compounds Mo₆X₈ belong to the family of insertion materials called Chevrel phases, that are of great interest for numerous purposes, in relation with their high ionic mobility. The electrochemical intercalation of different cations (Mg²⁺, Ni²⁺, Cu²⁺, Zn²⁺, Cd²⁺...) has already been demonstrated in the Mo₆X₈ host matrix, and presents an important potential for numerous applications. For instance, Mo₆S₈ thin films have been proven as an accurate sensor for the detection and quantification of hazardous elements, and Mo₆S₈ is a promising candidate as cathode material for the development of high performance rechargeable batteries, with particularly interesting results for Mg batteries.

Among the several potential applications, the Electrochemical Transfer Junction (ETJ) method for cadmium removal, proposed C. Boulanger et al., seems very promising. Cadmium (Cd) is a rare metallic element and its natural occurrence in the environment is always low (< 1 mg kg⁻¹), higher concentration levels result mainly from anthropogenic activities such as mining, smelting, metal plating, battery, waste disposal, or the application of phosphate fertilizers. The accumulation in soil and in the food chain generate a critical world-wide problem, being Cd highly toxic for plants and animals, and contaminated products consumption results in serious human diseases. For such reasons, several methods for the treatment of industrial wastewater have been developed, among them ETJ presents as advantages a high selectivity in cations removal but also the possibility to easily recuperate the removed cations with clear economic benefit for high valuable elements, as Co. The electrochemical process is based on the use of a Mo₆X₈ matrix as mineral junction between two half-cells, one containing the contaminated solution and the other the accumulation solution.

However, such technique has not been industrially applied to the removal of Cd²⁺ atoms yet and one of the possible explanations of such retard is the incertitude in the chemistry of Cd.
intercalation in Mo$_6$X$_8$ and its selectivity. Indeed, although the reaction has been already studied by Boulanger et al.\textsuperscript{8,22} and by Gocke et al.\textsuperscript{23} the results are still not clear and remain subject to reinterpretation\textsuperscript{7}. Effectively, Gocke et al.\textsuperscript{23} observed two potential plateau regions for Mo$_6$Se$_8$ and only one for MoS$_8$ during Cd$^{2+}$ galvanostatic intercalation in an aqueous electrolyte solutions. Such observation, with the support of \textit{ex situ} crystallographic cell parameters evaluation have been interpreted by the existence of two line-phases for the Mo$_6$Se$_8$ (CdMo$_6$Se$_8$ and Cd$_2$Mo$_6$Se$_8$) and only one-line phase for Mo$_6$S$_8$ ascribed to CdMo$_6$S$_8$. Boulanger et al.\textsuperscript{8} by cyclic voltammetry, confirmed the interpretation for Mo$_6$Se$_8$ but underlined some discrepancies in the case of Mo$_6$S$_8$. Although the experiment was in agreement with the existence of a single step intercalation process, the authors noticed that the initial stoichiometry CdMo$_6$S$_8$ could be enriched till Cd$_2$Mo$_6$S$_8$ by imposing the potential of the cathodic peak. Finally, such interpretation for S has been questioned by Levi on the basis of similitudes in respect to other cation intercalation for Mo$_6$X$_8$ phases\textsuperscript{7}.

Clearly a better knowledge of thermodynamic, kinetic and selectivity of intercalation electrochemistry would improve the development and industrialization of the ETJ technology. The understanding of the evolution of the phases with the potential, apart to represent a further step toward a more general comprehension of the interaction chemistry of Mo$_6$X$_8$ phases and a significant stage for future investigations, will allow to understand the evolution of materials during applications and prevent heterogeneous transformation, as proven by the several studies coupling structural techniques with electrochemistry\textsuperscript{3,24–27}. However, on a more fundamental point of view, the \textit{condicio sine qua non} to determine the atomistic origin of the intercalation chemistry and high mobility of Chevrel phases is a complete structural characterization of the intercalation compounds as proven by Levi et al.\textsuperscript{7,28}.

In this study, we report an \textit{in situ} coupled electrochemistry and synchrotron powder diffraction study of the intercalation and de-intercalation of Cd$^{2+}$ in Chevrel phases Mo$_6$X$_8$ (X=S, Se) in order to elucidate the intercalation processes (and possible pitfall in the galvanostatic intercalation experiment for these materials) and the structural determination of the phases obtained during the electrochemistry processes.

\textbf{Experimental}

Mo$_6$X$_8$ phases were synthesized by solid state reaction. Ni$_{1.7}$Mo$_6$S$_8$ was first prepared via solid-state synthesis from a stoichiometric mixture of nickel (99.7%, Sigma-Aldrich), molybdenum (99.9%, Sigma-Aldrich), and molybdenum sulfide (99.98%, Sigma-Aldrich) at
1150 °C for 48 h in a vacuum-sealed fused-silica tube. Then, Mo6S8 was obtained via nickel-ion extraction (oxidation) from Ni1.7Mo6S8 by stirring the powder overnight in a HCl·H2O (50:50) solution, followed by suction filtration, distilled water washing, and finally drying at 100°C in an oven for 24 h. Energy-dispersive X-ray spectroscopy (EDS) probe during SEM characterization and Cyclic Voltammetry show no residual trace of Nickel in the powder. Mo6Se8 is directly obtained by mixing the molybdenum (99.9%, Sigma-Aldrich), and molybdenum selenide (99.98%, Sigma-Aldrich) in stoichiometric quantities. The reagents were sealed in a Mo crucible and heated at 1300°C for 48 h.

Electrochemical measurements has been performed in a three-cell electrode at ambient temperature using as reference a commercial Ag/AgCl electrode (Bioblock) in aqueous electrolytes CdSO4 0.1 M (CdSO4·8/3H2O, Roth, purity >99%). Electrochemical experiments were conducted on an Autolab PGSTAT204 controlled by Nova software with calibrated electrode with a 10 mV precision. For the current densities the precision is equal to 0.2 %. Cyclic voltammetry (CV) sweep started from the open-circuit voltage (OCV) to negative potentials, sweep rate used being of 0.01 and 1 mV/s. The potential limits were set from -1 V (to avoid metal deposition and water reduction) to 0.3 V the limit of oxidation stability of the Mo6X8 phases.

Combined synchrotron X-ray powder diffraction and electrochemistry measurements have been performed in an ad hoc conceived electrochemical cell equipped by a working electrode aligned with the center of the two circle diffractometer, a reference electrode composed by Pt wire enrolled in spiral, and an Ag/AgCl reference electrode (all the potentials will be expressed in this potential scale). In order to avoid gradient concentrations of Cd during experiment, the electrolyte solution has been continuously refreshed in the cell by the use of a peristaltic pump. A scheme of the setup is drawn in figure S1 of the supplementary materials.

In order to avoid strong X-ray absorption effects, the working electrode containing Mo6X8 has been designed with the constraint of a maximum thickness of 100 µm. The working electrode was built using a Pt-wire enrolled in a spiral with an hole in the middle, coated with a slurry composed by a mix of Mo6X8 (X=S; Se) powder, and a polyvinylidene difluoride, (PVDF) binder (W#1300, Kureha Co.) (9:1 weight ratio), dispersed in carbonate propylene at 80°C. The electrode was finally dried at 60 °C for 72 hours, and at this stage the hole in the center was covered by a thin film of Mo6X8-PVDF in electronic contact with the wire, allowing the X-ray beam to pass without the interference with Pt. Each electrode was loaded with 15 mg of
Mo\textsubscript{6}X\textsubscript{8} over an area of about 1 cm\textsuperscript{2}, however such loading should be considered as nominal due to the possibility of a residual fraction of PVDF (< 10%) still present in the sample. The experiments were performed at the high resolution powder diffraction beamline ID22 at ESRF. The source of the beamline is a double undulator, with variable gap, and the emission is monochromatized by a Si (111) channel cut monochromator. The photons scattered have been collected by 9 scintillation detectors preceded by 9 Si (111) analyzer \textsuperscript{29} crystals. A wavelength of 0.354170 Å has been selected in order to minimize sample environment parasitic signals still maintaining a good resolution. During the \textit{in situ} experiments, in order to obtain a more general description, for each acquisition the electrochemical cell has been translated, sampling three different positions of the working electrode. Consequently, each acquisition is the average of three individual scans of 5.5 minutes going from 0.5° to 33° 2θ (d\textsubscript{min} ≈ 0.63 Å). The use of very thin electrodes combined with the high resolution configuration and the absence of sample movement (i.e. rotation) resulted in a poor grain orientation statistics strongly affecting the relative intensity of the different reflections. The existence of such inconvenient, that prevent a proper Rietveld analysis of \textit{in situ} data, is proven by the strong difference between the signal collected by each of the nine detectors, the impossibility to perform a physically sensible fit and the large difference between patterns collected for the same sample in \textit{in situ} configuration in respect to optimal rotating capillary configuration (i.e. \textit{ex situ}).

For such reasons, the \textit{in situ} data have been treated in the approximation of the crystallites composing the electrode don't change orientation significantly, the fraction of the different phases has been evaluated by the integrated intensity of an isolated specific reflection for the three phases scaled by the square of structure factor calculated from the corresponding crystal structure. Such procedure has been performed by the use of sequential Rietveld refinements, as coded in the FullProf code \textsuperscript{30} but limiting the refinement to the three ranges in which the selected reflection (using hexagonal setting (211) and (101) reflections for Cd\textsubscript{x}Mo\textsubscript{6}Se\textsubscript{8} and Cd\textsubscript{x}Mo\textsubscript{6}S\textsubscript{8}, respectively) for all Chevrel phases fall and to constrain the structural parameters to the values obtained by ex situ characterization.

In view of performing accurate Rietveld refinements, complementary experiments were carried out \textit{ex situ} on the binary Mo\textsubscript{6}X\textsubscript{8} and on powders obtained from laboratory electrochemistry experiments, using a rotating borosilicate glass capillary of 0.3 mm in diameter and higher counting statistics (around 1 hour for each sample). Rietveld refinements for \textit{ex situ} investigations have been performed by the use of Jana \textsuperscript{31} software for the structure
refinement and the localization of Cd ions, applying cylindrical absorption correction as implemented in the code using a packing fraction of 0.5 and calculated density. The regions with the few low intensity spurious peaks, ascribed to impurities, were excluded from the fit, in order to obtain more reliable refinements convergence and representative agreement factors. During refinement background has been modelled by a twenty five terms Chebyshev polynomial, and peaks profiles have been modelled by pseudo Voigt functions, further details could be retrieved in the CIF files. Maximum entropy method (MEM) calculations using observed intensity and phasing obtained by Rietveld refinement have been performed by the means of the Dysnomia code.

The structures for Cd,Mo6X8 have been determined from two samples that were biphasic, for X=S Cd1Mo6S8: 75.0(7) wt% Cd2Mo6S8: 25.0(15) wt% and for X= Se Cd1Mo6Se8: 40(1) wt%; Cd2Mo6Se8: 60(1) wt%. The results obtained by ex-situ refinements have been graphically represented with the help of the Vesta software.

Scanning Electron Microscopy (SEM) images of Mo6Se8 and Mo6S8 were collected at 30 kV with a Tescan Vega 3 microscope, in secondary electron imaging mode.

Results and discussion

Pristine materials characterization

Synthesized Mo6Se8 and Mo6S8 phases have been characterized by ex situ synchrotron radiation Rietveld refinement, confirming results obtained from laboratory diffractometers and absence of Nickel residual trace in Mo6S8. The crystal structure of both phases is trigonal with space group $R\bar{3}$ (148), and lattice parameters for hexagonal setting ($hP$) $a = 9.565247(15)$ Å and $c = 11.17121(2)$ Å for the Se phase and $a = 9.193645(14)$ Å and $c = 10.89027(2)$ Å for the S phase corresponding in the rhombohedral setting ($hR$) to $a=6.6606(1)$ Å, $\alpha=91.79(1)^\circ$ and $a=6.4305(1)$ Å, $\alpha= 91.26(1)^\circ$ for Se and S respectively. As expected, Mo6Se8 presents a larger cell volume than Mo6S8 (885.163(3) versus 797.158(2) Å³) due to a larger atomic radius of Se with respect to S, 1.20 Å versus 1.05 Å, respectively. The refined atomic parameters and selected interatomic distances for both phases are summarized in supplementary materials and are in good agreement with previous reports. However, due to the high counting statistics and resolution of synchrotron powder diffraction,
few spurious peaks have been detected ascribed to minor impurity phases (lower than 1 %) such as MoSe$_2$ and MoS$_2$, and probably few cluster condensed phases not detected during preliminary powder characterization. Analysis of peaks broadening using profile coefficients from the Rietveld refinement shows a small isotropic broadening due to internal strain (average max-strain $7.43(3) \times 10^{-4}$ and $6.02(3) \times 10^{-4}$ for Mo$_6$Se$_8$ and Mo$_6$S$_8$, respectively) while broadening due to crystallite size has been only detected for Mo$_6$S$_8$ with an average size of 3.06(1) μm. SEM images support XRD evaluation, both materials showing grains with irregular parallelepiped shape with a larger dimension for Mo$_6$Se$_8$ than for Mo$_6$S$_8$, from 10 to 30 μm for the former and few μm for the latter (Figure S2 part c and d).
Cyclic voltammetry

*Mo₆Se₈*

In figure 1 part a) is shown the cyclic voltammetry of Mo₆Se₈ at 0.01 mV/s in an aqueous solution of CdSO₄ 0.1 M. The voltamperogram is characterized by two intercalation peaks  E_{int1} and E_{int2} (around 0.35 V and -0.52 V) and two deintercalation peaks  E_{deint1} and E_{deint2} (around -0.42 V and -0.25 V), in agreement with the literature. In accordance with the works of Boulanger et al the first peak is ascribed to the intercalation of a single Cd²⁺:

\[
Mo₆X₈ + Cd^{2+} + 2e^- \rightarrow CdMo₆X₈ \quad (1),
\]

while the second peak corresponds for the selenide phase to the intercalation of the second Cd²⁺:

\[
CdMo₆X₈ + Cd^{2+} + 2e^- \rightarrow Cd₂Mo₆X₈ \quad (2).
\]

Both reactions are reversible and the ionic transfer is mainly limited by cation diffusion in the solid, with the first insertion showing a cation diffusion slower than the second due to the different broadening of the peaks when potential sweep rate is changed. However, differently from previous experiments when the cyclic voltammetry is performed with a sweeping rate of 1 mV/s or higher (Figure 1 part b), a new peak (E_{inta}) appears at intermediate values between the first and second intercalation peak. The nature of this intermediate peak will be discussed later with the support of *in situ* structural investigations as cyclic voltammetry alone is not sufficient to ascribe this peak.

Figure 1: cyclic voltammetry of part a) and b Mo₆Se₈ at two different cycling rate and part c) of Mo₆S₈ with the rate 0.01 mV/s for 0.1 M solutions of Cd²⁺.
Mo₆S₈ cyclic voltammetry is clearly different from MoSe₈, the voltamperogram (Figure 1 part c)) exhibits only one intercalation peak and one de-intercalation signal (at -0.62 V and -0.49 V) independently from the sweep rate. According to the literature⁸,²³,³⁷, the assignment of such peaks is debated among the reaction (1) or reaction (2).

Galvanostatic intercalation

The intercalation reactions observed by cyclic voltammetry have been investigated *in situ* by coupled diffraction/electrochemistry experiments in galvanostatic mode *i.e.* imposing a fixed current and monitoring the potential variation necessary to counterbalance the change in resistance of the system, following intercalation/deintercalation processes. Such approach, focused on the thermodynamics of the reaction, is very widespread for combined XRD/electrochemical experiment²,³,²³–²⁷ because it has the advantage to promote reactions in the bulk and not limited to the surface of the electrode (as CV). Moreover, although this technique doesn’t allow a kinetic modeling of the reactions, it allows to directly relate the type of reaction with the shape of potential evolution with a clear complementarity with structural techniques³⁸.

**Selenium phases:**

A typical series of diffraction patterns taken during an intercalation cycle is shown in Figure 2 a), displaying in the inset a zoom at low angles around reflection (100) in *hR* setting of binary Mo₆Se₈. The patterns could be deconvoluted as the sum of up to three contributions, the initial binary Mo₆Se₈ phase (space group *R̅3*, *a*=9.543, *c*= 11.17 Å), a first rhombohedral intercalated phase (space group *R̅3*, *a*= 9.76, *c*=11.20 Å) that is ascribed to CdMo₆Se₈²⁸, and a second rhombohedral ternary phase (space group *R̅3*, *a*=10.2489, *c*=10.8789 Å) that has been previously ascribed to Cd₂Mo₆Se₈³⁶, but whose structure is still unknown. A more clear correlation between stoichiometry and cells metric could be noticed adopting the rhombohedral setting, insertion of cations generate an elongation of the packing and an
increase of the trigonal distortion $(\text{Mo}_6\text{Se}_8 \ a = 6.65 \ \text{Å}, \ \alpha= 91.7^\circ$, $\text{CdMo}_6\text{Se}_8 \ a = 6.76 \ \text{Å}, \ \alpha= 92.44^\circ$, $\text{Cd}_2\text{Mo}_6\text{Se}_8 \ a = 6.94 \ \text{Å}, \ \alpha= 95.19^\circ$).

Figure 2: part a) In situ diffraction pattern during galvanostatic intercalation in $\text{Mo}_6\text{Se}_8$: part a) typical pattern (inset: zoom around the reflection of the different phases during the first intercalation stage); part b) from to top to bottom: estimated phase fractions and corresponding voltage evolution as a function of time.

In Figure 2 b) the evolution of the weight fractions of the phases associated with the potential evolution during a galvanostatic cycle is shown. During the intercalation process, in the initial stage of reduction ($t < 50 \ \text{s}$), the potential decreases rapidly in the range between 0 and -0.35 V with negligible change in XRD patterns, such abrupt variation being typical for galvanostatic electrochemistry experiment and generally associated to small deterioration of the electrode or other parasitic currents. As a consequence, the real first stage of reaction is between 50 and 1350 s with an apparent single phase reaction region, i.e. a region with continuous and monotone potential variation, where the potential pass from -0.35 to -0.50 V which corresponds in accordance with CV to the potential of the first insertion of Cd in the structure. However, the inspection of the diffraction data shows clearly that it is rather a two-phase region with the appearance and increase of the peaks associated to $\text{Cd}_1\text{Mo}_6\text{Se}_8$. 
A second intercalation stage is present between 1350 and 2820 s (0.14 e-/mol) where the potential remains almost constant at around -0.50 V. This region, from an electrochemistry point of view, due to the constant potential, presents a clearer two-phase character and could be ascribed to the second intercalation peak $E_{\text{int}_2}$ presents on the CV in fast sweeping condition. From a structural point of view, the intercalation mechanism is not clear, due to the insufficient time resolution of XRD acquisition. In the interval of just one data acquisition of 16.5 min, several processes mixed together, inducing a small increase of Cd$_1$Mo$_6$Se$_8$, a clear increase of Cd$_2$Mo$_6$Se$_8$, and a strong decrease of the original Mo$_6$Se$_8$ phase.

The final stage of intercalation occurs between 3000 and 13350 s where a second zone with a constant potential (-0.56 V) is present along with a clear decrease of Mo$_6$Se$_8$, specular to the increase of Cd$_2$Mo$_6$Se$_8$, as visible in Figure 2b, meanwhile the Cd$_1$Mo$_6$Se$_8$ XRD peak remains rather unchanged. On the basis of the structural results and cyclic voltammetry, this process should be related to the direct intercalation of 2 Cd$^{2+}$ atoms within the binary grains, as described in the reaction (3)

$$\text{Mo}_6\text{X}_8 + 2\text{Cd}^{2+} + 4e^- \rightarrow \text{Cd}_2\text{Mo}_6\text{X}_8 \quad (3)$$

The de-intercalation process has been studied on the same sample just by inverting the current flow (red curve in Figure 2b). De-intercalation is limited to only a minority part of the sample before the potential rises to high values. The de-intercalation process could be divided in two double phase transformation regions: a first region from 14370 to 17670 s with a potential at 0.45 V and a second from 18500 to 21100 s at -0.28 V corresponding to the two peaks present in the deintercalation electrochemical signal of Figure 1a. In the first region of de-intercalation, the decrease in intensity of the peaks ascribed to Cd$_2$Mo$_6$Se$_8$ is concomitant with the increase of the ones of Cd$_1$Mo$_6$Se$_8$ (the binary XRD peaks remaining almost constant in intensity), leading to the almost complete disappearance of the Cd$_2$Mo$_6$Se$_8$ phase, so this de-intercalation step could be ascribed to the inverse of reaction 2; during the second step of de-intercalation, peaks ascribed to Cd$_1$Mo$_6$Se$_8$ decreases accompanied by an increase of the intensity of peaks associated to the binary phase following the inverse of reaction (1). It should be underlined that the binary phase obtained presents a peculiar peak profile with a pronounced asymmetry on the left that could be caused by experimental inconvenience such as a gas bubble standing in front of the beam or by the existence of very strained additional phase parent of the initial Mo$_6$Se$_8$ (Figure S3).
In order to better investigate the intercalation steps after the de-intercalation process, a further intercalation process has been repeated on the same electrode, but with a lower imposed current of -0.075 mA (dashed black curve on figure 2 b). The lower current has two effects: (i) it avoids the polarization of the electrode due to the low ionic conductivity; (ii) it allows a better reaction sampling. Such second intercalation process presents a simpler electrochemical profile with respect to the precedent, with two double phase regions. The first one at a potential of -0.32 V that could be associated by XRD to reaction (1) while the second at around -0.50 V that should be linked to a mechanism involving the transformation of Cd$_1$MoSe$_8$ into Cd$_2$MoSe$_8$ as described by reaction (2).

It is worth underlining, that for Mo$_6$Se$_8$, the limited rate of intercalation and deintercalation reaction affects chronovoltametry with large imposed current as well as CV for rapid sweeping, revealing interesting kinetic effects. In the hypothesis that solid state diffusion is the limiting step for the intercalation and that the diffusion of Cd$^{2+}$ in Chevrel phases depends on the concentration of Cd$^{2+}$ within the phase, the higher the concentration, the faster the diffusion $^7$, we could propose the following interpretation to explain the results during intercalation reaction in Mo$_6$Se$_8$:

For slow intercalation (present experiment 0.075 mA for chronopotentiometry and 0.01 mV/s for CV). During galvanostatic intercalation, if the current imposed is lower than the maximal possible diffusion of Cd$^{2+}$ in Mo$_6$Se$_8$, the reaction proceeds passing from reaction 1, and when all the Mo$_6$Se$_8$ is transformed in Cd$_1$Mo$_6$Se$_8$, a second Cd per formula unit starts to be intercalated as described in reaction (2), therefore only two steps in the potential are present as shown for dashed curve in Figure 2 b). Also CV could be interpreted in the same approach, the first peak corresponds to reaction 1 and the second to reaction 2, both peaks being related to a single intercalation and present similar integrated area.

For fast intercalation (present experiment 0.15 mA for chronopotentiometry and 1mV/s for CV). During galvanostatic intercalation, reaction 1 will happen in the initial period of intercalation but will be rapidly limited to the surface of the Mo$_6$Se$_8$ grains, due to the insufficient diffusion of Cd$^{2+}$ in Mo$_6$Se$_8$, at such point the potential decreases till it is sufficient to activate reaction 2, but such reaction will cease rapidly once all the limited amount of Cd$_1$Mo$_6$Se$_8$ has been consumed. This statement is supported by similar duration for the first two steps of the intercalation at -0.15 mA. Once reaction 2 is terminated, the system is forced to decrease further the potential to activate reaction 3, the direct conversion of
Mo$_6$Se$_8$ into Cd$_2$Mo$_6$Se$_8$. Owing to the fact that CV with rapid sweeping time presents three anodic peaks, the first could be ascribed to reaction 1 and it is strongly deformed by the rapid sweep of voltage, the second one, noted as $E_{\text{inta}}$ in figure 1 b) in accordance with galvanostatic intercalation should correspond to the second peak for slow sweeping condition (reaction 2), and finally the third peak should correspond to reaction 3.

For de-intercalation. From a chemistry point of view, for Cd$_x$Mo$_6$Se$_8$ de-intercalation reactions consist in inverting the reactions discussed in the previous paragraphs. However, based on the physics of transport, it is worth underlining that for intercalation reaction, the front of reaction move from the surface to the center of the crystallites while for de-intercalation the front of the reaction is on the liquid-solid interface. This difference implies that if during galvanostatic de-intercalation the imposed current is higher than the maximum diffusion in the electrode, the surface region will be significantly depleted of Cd compared to the bulk. As the diffusion is depending from the Cd loading in Mo$_6$X$_8$, most of the surface results in Cd being depleted, lowering the diffusion, with the consequence to arrive to a kind of trapping mechanism for Cd in the bulk of the electrode. In the de-intercalation experiment at 0.15 mA (red curve Figure 2 b)), in the first two phase region, the Cd$_2$Mo$_6$Se$_8$ is oxidized to Cd$_1$Mo$_6$Se$_8$ with a rate slightly higher than the Cd diffusion, after the potential increases for the inverse of reaction 1, but the diffusion from the bulk will be also lower and consequently forming a surface layer of Mo$_6$Se$_8$ which hinders further oxidation and consequently a significant fraction of Cd$_x$Mo$_6$Se$_8$ is still present at the end of the de-intercalation. For CV at low sweeping rate, two cathodic peaks are present corresponding to the inverse of reaction 2 and 1 with similar intensity and width whereas for fast sweeping rate the peak corresponding to the inverse of reaction 1 decreases in intensity and increases in width due to the lower diffusion.

Mo$_6$S$_8$

Figure 3 shows, in analogy with the work done for selenium phase, a set of diffraction patterns collected in situ in the part a and the evolution as function of time of the integrated intensity of a few selected reflections for the different phases as well as the potential. The XRD patterns are also a convolution of up to three phases: the initial binary Mo$_6$S$_8$ phase (space group $R\bar{3}$, $a=9.19$ Å, $c=10.89$ Å), a first rhombohedral intercalated phase (space group $R\bar{3}$, $a=9.41750$ Å, $c=10.74200$ Å that is ascribed to Cd$_1$Mo$_6$S$_8$, and a second rhombohedral
ternary phase (space group $R\bar{3}$, $a=9.82$ Å, $c=10.57$ Å) refined as to Cd$_2$Mo$_6$S$_8$ (see below). As in the case of Mo$_6$Se$_8$ the rhombohedral setting shows a clearer correlation with Cd stoichiometry with an increase of the cell parameter and the rhombohedral angle: Mo$_6$S$_8$ $a = 6.43$ Å, $\alpha = 91.26^\circ$, Cd$_1$Mo$_6$S$_8$ $a = 6.51$ Å, $\alpha =92.63^\circ$, Cd$_2$Mo$_6$S$_8$ $a = 6.67$ Å, $\alpha =94.74^\circ$.

Figure 3 a) Left: typical pattern (inset: zoom around the hP (101) reflection of the different phases during the first intercalation stage); part b) from to top to bottom: estimated phase fractions and corresponding voltage evolution as a function of time.

Interestingly although three phases are potentially present as for the Se case, the electrochemistry appears at first sight very different, with only one “plateau” present for both intercalation and de-intercalation in accordance with CV where only one cathodic and one anodic peaks are present. Looking in details to the electrochemistry data and to the structural ones, we notice that in the first step of intercalation $t < 10000$ s (in Figure 3 b) the potential stabilizes rapidly around -0.57 V, suggesting a two phase region, this range could be clearly ascribed to reaction (3) with the direct conversion of Mo$_6$S$_8$ to Cd$_2$Mo$_6$S$_8$.

The de-intercalation process (red curve Figure 3) behaves electrochemically similarly, with another plateau at -0.5 V that lasts for a much shorter time (around 5000 s). However, when looking at the diffraction data, it is clear that this two phase region is not the inverse of the previous intercalation reaction (3), but the inverse of reaction (2), i.e., the conversion of Cd$_2$Mo$_6$S$_8$ into Cd$_1$Mo$_6$S$_8$. That implies that a Cd trapping effect is present, with the
irreversible accumulation of Cd in the structure of CdMo₆S₈ as proposed by Levi et al.⁷. As a consequence, the anodic peak of CV should be ascribed to reaction 2 (or 3 if the sample has never been intercalated) while the cationic peak should be related to the inverse of reaction 2 and not to the reaction 1. This is thus clarifying the Boulanger et al.⁸ and Gocke et al.²³ experiments.

In order to clarify this process, a second intercalation/de-intercalation cycle has been performed. During this second cycle, the main difference is that at the beginning of intercalation, half of Mo₆S₈ has been converted to Cd₁Mo₆S₈ from the previous cycle. During intercalation the main reaction is the conversion of Cd₁Mo₆S₈ into Cd₂Mo₆S₈, during a plateau that, although characterized by a strong curvature, could be tentatively defined as -0.55 V. Such value is very close to the intercalation potential of reaction (3) at least in the present experimental configuration, explaining the difficulty to discriminate between the two reactions in the CV. In order to support such attribution, it is possible to use the first cycle, where a small shoulder is present before the plateau (time <2000 s) in which the potential is slightly higher and it is possible to notice that the small amount of Cd₁Mo₆S₈, created during CV test before interaction, is converted into Cd₂Mo₆S₈. For the second de-intercalation cycle, the condition are the same as for the first cycle, although Cd₁Mo₆S₈ has a higher concentration and consequently the reaction follows the scheme of the previous one.

![Graphical representation of kinetic effect during intercalation and de-intercalation of Cd.](image-url)

Figure 4: graphical representation of kinetic effect during intercalation and de-intercalation of Cd.
In a more general approach, the electrochemistry of Mo₆S₈ should be interpreted as an asymptote of an extremely fast experiments for Mo₆Se₈. For instance, in the CV case, the peaks $E_{\text{int1}}$, $E_{\text{inta}}$, $E_{\text{deint1}}$ decrease in intensity and broaden as sweep rate is increased. Asymptotically for very fast rate, these peaks will practically disappear and only the peaks $E_{\text{int2}}$ and $E_{\text{deint2}}$ remain visible. Considering that electrochemical parameters are similar, the current interpretation is that for reaction (1), Cd diffusion in Mo₆S₈ should be several order of magnitude lower than in the case of Mo₆Se₈.

The present model has been graphically depicted in figure 4, it is worth underlining that in the case of fast intercalation the rate determining step will be the diffusion of Cd²⁺ from Mo₆X₈ to Cd₂Mo₆X₈ (process (3)) with the later phase continuously replenished by the intercalation. For the fast de-intercalation, the elucidation of the rate determining step is more complex. After a first period in which the surface is completely depleted and an interface of Cd₄Mo₆X₈ has been formed between the bulk Cd₄Mo₆X₈ phase and the surface, the prosecution of the de-intercalation is limited by the diffusion of Cd²⁺ from Cd₄Mo₆Se₈ or Cd₂Mo₆Se₈. However, more and more the de-intercalation proceed, more the phase are depleted in Cd and slower is the diffusion. As a consequence, if the diffusion of Cd in Mo₆X₈ is very low, the cation is practically trapped.

**Structural analysis**

Independently from the chemical composition, the crystal structure of the Chevrel phases is basically the same: a trigonal distorted simple cubic stacking of Mo₆X₈ blocks (Mo₆-octahedral clusters inside an anion cube) forming an open three-dimensional channel system in which cations are hosted. The channels are formed by the association of two types of face-sharing pseudocubic cavities: cavity 1 (blue polyhedra in Figure 5) and cavity 2 (green polyhedra in Figure 5), corresponding to the body centered and the face centered sites of the Mo₆X₈ blocks packing, respectively.

According to the literature⁷,³⁹ cation positions are strongly correlated with ionic mobility. on the base of this parameter, the ternary Chevrel phases could be divided in two families: type I were cations are located only near or exactly in the origin of cavity 1, that is the case of large cations with no mobility(like Pb²⁺ for instance) and type II for which cations could be present
in both cavities and shifted from the center of the cavities as observed for small and mobile cations (like Li$^+$ for instance). Cd$^{2+}$ has an ionic radius near 1 Å (0.95 Å for coordination number = VI according to R.D. Shannon) as a consequence it is quite difficult to classify it into one of the two classes \emph{a priori}, and structural analysis is mandatory for the understanding of the materials. For such reasons the following sections will be devoted to describe the structures of intercalated phases resulting from Rietveld refinement with particular interest for the positions of Cd$^{2+}$ that have been determined by Fourier difference and successively refined (positions, occupancy and isotropic atomic displacement parameters). In particular, the Cd position will be discussed with the help of electronic density maps obtained by the Maximum Entropy Method (MEM)$^{40}$. In this context, the main advantages of MEM are the possibility to use fully phased data (the phase contribution of Cd is included) and the quality of the density distribution concerning the background, which is usually close to zero and allows the visualization of details, which are inconclusive with standard Fourier techniques.

Figure 5: Part a) General structural model for Chevrel phases in hexagonal setting ($R\bar{3}:hP$) with cavity 1 and 2, blue and green polyhedra. Part b) Mo$_6$X$_4$ blocks simple cubic packing in
rhombohedral setting ($R3\overline{3}:hR$) with cavity location.

$Cd_1Mo_6S_8$. As clear from the previous paragraph, it is reasonable to suppose two different models for $Cd_1Mo_6S_8$ crystal structure: the first for which $Cd^{2+}$ is located in the center of cavity 1 in Wyckoff position $3a$ (0.666, 0.333, 0.333) ($R3\overline{3}:hP$), and a second model, still inside the cavity 1 but with Cd at general position $18f$. Both models have been used as input for Rietveld refinement of the ex situ synchrotron powder diffraction data. However the first model leads to abnormally too huge value for the $B_{iso}$ of Cd. Analysis of Fourier difference map obtained after setting Cd occupancy to zero (Figure S4) shown a residual the electron density within cavity 1 rather centered on the $3a$ site but very diffuse in the $xy$ plane. In order to discriminate between the two models, MEM analysis has been applied, as visible in Figure 6, the map obtained shows a clear shift of Cd density from the center of cavity 1, supporting model 2 with Cd at $18f$ position (0.644(9), 0.265(2), 0.333(3) $R3\overline{3}:hP$) slightly shifted from the $3a$ site (0.57(2) Å).

As a consequence, we propose for describing the current data the second model with Cd on the $18f$ general site. Such model is supported by a previous combined neutron and laboratory X-ray powder diffraction study$^{24}$ in which Cd has been refined at a very similar position, with a shift of 0.60 Å from the origin. It is worth noticing, that Cd positions generated by symmetry form a six-membered ring in the cavity 1, with a maximum distance between the sites of 1.14 Å, value too small to allow the presence of two Cd in the same cavity 1. Occupancy of Cd was freely refined (its $U_{iso}$ parameter being made identical to the $U_{iso}$ value of Mo) and converged near 1/6 value (0.1598(17)); it implies that each cavity 1 in the structure is fully occupied.
Figure 6: Cd\textsubscript{1}Mo\textsubscript{6}S\textsubscript{8}. Part a) MEM electronic density map within the a-b plane focused on cavity 1, z=1/3 (hP setting) Max./min. saturation levels at 80/0 e\textsuperscript{-}Å\textsuperscript{3}. Part b) Graphical representation of the Cd\textsubscript{1}Mo\textsubscript{6}S\textsubscript{8} structure in the (hR setting).

\textit{Cd}\textsubscript{2}Mo\textsubscript{6}S\textsubscript{8}. Once Cd\textsubscript{1}Mo\textsubscript{6}S\textsubscript{8} has been classified as a type 2 ternary Chevrel Phase, and as the ionic radius of Cd\textsuperscript{2+} prevents a further occupation of cavity 1, the most reasonable hypothesis is that cavity 2 has to be filled. Indeed, Fourier difference (Figure S5) and MEM (Figure 7 a)) maps obtained after refinement using as starting model the previous crystal structure of Cd\textsubscript{1}Mo\textsubscript{6}S\textsubscript{8} show an additional electron density peak located at position 0.503(5) 0.045(5) 0.464(5) (R\textsubscript{3}:hP), generating two equivalent sites (around the 9d special site at ½ 0 ½) distant from each other by 1.14(11) Å, located in the cavity 2. It is worth underlining that the Cd site in cavity 1 is also strongly modified by the insertion of the second Cd, moving to position 0.606(8) 0.198(5) 0.353(5) and drastically shifted further from the center of the cavity (from 0.57 Å to 1.17 Å). MEM and Fourier difference map clearly show electron density maxima more shifted from the site 3a at the center of cavity 1 (see Figure S6 and Figure 7 b)), in respect of the case of Cd\textsubscript{1}Mo\textsubscript{6}S\textsubscript{8}. The refinement of the occupancy for Cd in cavity 1 and 2 converges freely to 0.167(8) and 0.187(12), respectively, corresponding to a stoichiometry of Cd\textsubscript{2.12}Mo\textsubscript{6}S\textsubscript{8} and an average occupancy of 1.12 and 0.37 for cavity 1 and 2, respectively.
Figure 7: Cd$_2$Mo$_6$S$_8$. Part a) Cavity 2: MEM electronic density map within the $a$-$b$ plane at $z=0.5347$ showing three maxima corresponding to the three in-plane Cd$_2$ atoms, shifted from the ideal 9$d$ Wyckoff site at $\frac{1}{2} 0 \frac{1}{2}$ (max./min. saturation levels at 8/0 e/Å$^{-3}$). Part b) Cavity 1: MEM electronic density map within the $a$-$b$ plane focused on cavity 1, $z=0.3472$ ($hP$ setting). Max./min. saturation levels at 5/0 e·Å$^{-3}$. Part b) Graphical representation of the Cd$_2$Mo$_6$S$_8$ structure in the rhombohedral cell.

$Cd_1Mo_6Se_8$. $Cd_1Mo_6Se_8$ has been refined starting from the structural model (site 18$f$) obtained for $Cd_1Mo_6S_8$ achieving satisfactory agreement factors and acceptable $U_{iso}$ values ($R=3.06\%$, $\chi^2=1.63$). The position of Cd also turns out to be shifted from the center of cavity 1 to the position 0.65(3) 0.261(8) 0.331(11), which is slightly more shifted from the center of the cavity in respect of $CdMo_6S_8$ (0.64 versus 0.57 Å), with an occupancy of 0.143(11) corresponding to $Cd_{0.86}Mo_6Se_8$. Also in such case the observed density from Fourier difference maps (Figure S7) presents a hugely delocalized apparent maximum on the 3$a$ site, but MEM map shows a clear preference for Cd in site 18$f$. The data fitted in accordance with MEM analysis shows a very similar Cd disorder in cavity 1 in both $Cd_1Mo_6S_8$ and $Cd_1Mo_6Se_8$ phases, despite the different anionic lattices (including very close Cd-X distances distribution, ≈0.08 Å longer for Se as expected according to the ionic radii considerations).
Figure 8: \( \text{Cd}_1\text{Mo}_6\text{Se}_8 \) Part a) phase, cavity 1: MEM electronic density map within the \( a-b \) plane centered at \( 2/3 \ 1/3 \ 0.3194 \) showing three maxima corresponding to the three in-plane Cd1 atoms (max./min. saturation levels at 30/0 e/Å\(^3\)). Part b) graphical representation of the \( \text{Cd}_1\text{Mo}_6\text{Se}_8 \) structure in the rhombohedral cell.

\( \text{Cd}_2\text{Mo}_6\text{Se}_8 \). By analogy with the other Chevrel phases, the starting model used for such phase has been the crystal structure of \( \text{Cd}_1\text{Mo}_6\text{Se}_8 \) plus an additional site for Cd in a general Wyckoff positon \( 18f \) inside the cavity 2. However, both Cd sites converged into cavity 2. Fourier difference maps (Figure S8 a)) confirmed that cavity 1 is essentially empty and all the Cd electron density lies within cavity 2 (see Figure 9). For such reasons the fit has been performed finally using a single site in cavity 2 that converged at position 0.518(3) 0.100(2) 0.445(2) with occupancy 0.325(10) corresponding to a stoichiometry of \( \text{Cd}_{1.95}\text{Mo}_6\text{Se}_8 \). Symmetrically equivalent Cd atoms are separated by 2.24(3) Å within cavity 2, twice the value encountered in cavity 2 of \( \text{Cd}_2\text{Mo}_6\text{S}_8 \). MEM maps confirmed this cavity occupancy scheme (Fig. 9a) and b)).
Figure 9: Cd$_2$Mo$_6$Se$_8$. Part a) MEM electronic density map within the $a$-$b$ plane at $z=0.4444$ ($hP$) showing three maxima corresponding to the three in-plane Cd2 atoms, shifted from the ideal 9$d$ Wyckoff site at ½ 0 ½ (max./min. saturation levels at 10/0 $e^{-}/Å^{-3}$). Part b) MEM electronic density map within the $a$-$b$ plane centered at 2/3 1/3 0.3194 cavity 1, same zoom, orientation and scale of Figure 8 a). Part c) Graphical representation of the Cd$_2$Mo$_6$Se$_8$ structure in the $hR$ setting.

Crystal structure of Cd$_2$Mo$_6$Se$_8$ is not following the expected scheme of successive occupation of the two cavities, implying also the migration of Cd from cavity 1 to cavity 2 during intercalation. Although unexpected, a similar behavior has been reported for the intercalation of Mg$^{2+}$ in Mo$_6$Se$_8$\textsuperscript{41}, with the two line phases MgMo$_6$Se$_8$ and Mg$_2$Mo$_6$Se$_8$. Although both triclinic, those phases present the same cavity filling scheme as observed in the trigonal Cd$_2$Mo$_6$Se$_8$ phase. In this context, it is worth noticing that such similarity between the two cations is maintained also for Sulphur phases. MgMo$_6$S$_8$ has a trigonal structure with Mg hosted in cavity 1 at 0.90(3) Å from its center, that presents also some trapping character, further intercalation form the trigonal Mg$_2$Mo$_6$S$_8$ that, as for Cd$_2$Mo$_6$S$_8$, hosts the additional Mg in the cavity 2 (18f site) and the site in the cavity 1 moves further away from the center (1.117(18) Å from site 3a).

In order to better understand the intercalation chemistry of Cd into the Chevrel phases, Bond valence sum (BVS) and Bond valence site energy (BVSE) maps have been calculated for the four intercalated ternary structures. BVSE is a method derived from bond valence sum \textsuperscript{42} that
combines the bond valence sum mismatch map in a form of a Morse potential with a cation-cation electrostatic repulsion term. It presents as main advantage, compared to bond valence sum mismatch map, an energy scale that could be compared with atomistic calculations and a more accurate treatment of the sites near other cations. However, it is worth noticing that in the specific case of transition metal cluster compounds the results must be considered only due to the difficulties to treat in such simple approximation the cluster charges and lattice strain effects. The obtained map for selenium are visible in Figure 10, and the method is able to reproduce the main features of the experiments such as the lowest conductivity and preferential occupation of cavity 1 for Cd₁Mo₆X₈ or the stabilization of the site in cavity 2 accompanied by an increase of shift from the center of the cavity 1 for Cd₂Mo₆X₈ although the BVSE doesn’t show a clear inversion of stability between the two sites for Cd₂Mo₆Se₈.

Figure 10: Part a) Chevrel phases in hexagonal setting (R̅3:mP), BVSE map for Cd₁Mo₆Se₈ and Cd₂Mo₆Se₈ part b) and c), respectively. Red and yellow isosurfaces corresponds to 0.3 and 0.5 eV above the minimum and the percolation energy, respectively.

Due to the relatively good agreement of such method, it is of interest to consider separately the two main components of the BVSE: valence mismatch and coulomb interaction. Effectively the BVS clearly shows that the site on cavity 2 presents a higher valence mismatch also for Cd₂Mo₆X₈ phases (estimated valence around 2.5 and 2.0 for cavity 2 and 1, respectively) although in the BVSE map the two sites result with a similar stability. As a consequence, the stabilization of the cavity 2 for BVSE method in Cd₂Mo₆X₈ should be ascribed to Mo-Cd repulsion. Actually, the use of a higher oxidation state for Mo in BVSE calculation results in a further stabilization of cavity 1. In this approximation, excluding a direct covalent interaction between the cluster and Cd²⁺ due to the relatively long distance between Mo and Cd, the key parameter for the stabilization of cavity 2 is the actual charge on the cluster that is in turn dependent from the electron count but also to steric effects.
variation of the cluster geometry during the intercalation could be used as indirect indicator of the cluster charge, and effectively it could be noticed that cluster apical distance is almost independent from the anion for Mo$_6$X$_8$ (X=Se: 3.9087(12) Å, X=S: 3.9346(11) Å) and CdMo$_6$X$_8$ (X=Se: 3.842(7) Å, X=S: 3.844(7) Å but not for Cd$_2$MoX$_8$ (X=Se: 3.790(14) Å, X=S: 3.73(2) Å) for which the cavity occupation scheme is different.

It is worth underscoring that the different cavities occupations for Cd$_2$Mo$_6$X$_8$ have a fundamental consequence in ionic transport mechanism. Effectively, although the topology of the percolation paths remain the same, the availability of unoccupied nodes during cationic transport changes drastically. For instance, in the case of Cd$_2$Mo$_6$S$_8$, Cd hopping is formally hindered by the full average occupancy of cavity 1, implying locally a vacancy mechanism of ionic conduction, while in the case of Cd$_2$Mo$_6$Se$_8$, two empty cavities 2 over the six connected to the closest empty cavity 1 are always available for Cd hopping.

**Conclusion**

In the present work the electrochemical intercalation/de-intercalation of Cd into the Chevrel phases Mo$_6$X$_8$ (X=S, Se) has been studied in situ, and the crystal structure of the intercalated phases so obtained has been determined. The processes of the electrochemical Cd insertion into the host material Mo$_6$X$_8$ occurred stepwise, involving only two separate phases Cd$_1$Mo$_6$X$_8$ and Cd$_2$Mo$_6$X$_8$, although the electrochemical characterization shows a more complex picture than debated in the past. Effectively, the coupling of in situ X-ray powder diffraction and electrochemical measurements has allowed to reveal a strong dependency of the process from the imposed electrochemistry kinetics, mainly dependent on the less common increase of ionic conductivity during intercalation, in particular the competition for the formation of Cd$_2$Mo$_6$X$_8$ between the two step (Mo$_6$X$_8$ → CdMo$_6$X$_8$ → Cd$_2$Mo$_6$X$_8$) and the single step (Mo$_6$X$_8$ → Cd$_2$Mo$_6$X$_8$) intercalation processes or the trapping mechanism for CdMo$_6$S$_8$.

Ex situ synchrotron X-ray Rietveld refinement of samples in various stage of intercalation has allowed to determine the structure of Cd$_2$Mo$_6$X$_8$ phases previously unknown. The Cd$_2$Mo$_6$Se$_8$ shows an unexpected structure, in which Cd is hosted integrally in the cavity 2, in opposition to the classical scheme and to the structure of the isostoichiometric Cd$_2$Mo$_6$S$_8$. Such variation in cavity stability cannot be rationalized only with valence misfit calculated by BVS; BVSE calculations suggest that also coulomb interaction should be taken in account and the
stabilization of cavity 2 should be related with the actual charge on the Mo cluster. The different cavity filling scheme for Cd$_2$Mo$_6$Se$_8$ implies a different transport mechanism that could have strong effect on conductivity and applications.

The results shown in this work should be an important basis for a better understanding of the electrochemical intercalation reaction of divalent ions into Mo$_6$X$_8$ phases and an useful indication for device development that are founded on such processes.

ASSOCIATED CONTENT

Supplementary figures (PDF), including experimental setup, Rietveld refinements and Fourier differences.

Detailed crystallographic information (CIF files), copy of the CIF files are available on ICS database codes:

1864057  Cd1Mo6S8
1864019  Cd2Mo6S8
1864058  Cd1Mo6Se8
1864021  Cd2Mo6Se8

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References


Synopsis:

The chemistry of electrochemical intercalation of Cd\(^{2+}\) in Chevrel phase has been investigated by combined synchrotron X-ray powder diffraction and electrochemistry. The crystal structure of Cd\(_2\)Mo\(_6\)X\(_8\) (X=S, Se) phases, only synthesizable by a topotactic intercalation, has been determined, showing for Cd\(_2\)Mo\(_6\)Se\(_8\) a novel trigonal structure with only cavity 2 occupied.