ESR STUDIES OF INTER- AND INTRA-LAMELLAR CATION EXCHANGE PROCESSES IN Cd₂P₂S₆

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Abstract—The transition metal chalcogenides, M₂P₂S₆ (M = divalent transition metal cation), are lamellar materials that undergo an unusual cation exchange process when their crystals are placed in contact with a solution of a transition metal salt. EPR spectroscopy has been used to examine the exchange of paramagnetic Co²⁺ for Cd²⁺ in diamagnetic Cd₂P₂S₆. It has been possible to study the uptake of ions by the lamellar lattice and to examine the cation coordination environment through its effect on the parameters of the spin-Hamiltonian. In order to extract details of the Co²⁺ coordination from the observed EPR spectra, theoretical values of the g-value have been computed as a function of several crystal field parameters. The results for Co²⁺ are compared with previous results for the insertion of Mn²⁺ into Cd₂P₂S₆ by cation exchange. The relative occupancy of the inter- and intra-lamellar sites is a function of the cation type and the solvent employed.

Keywords: ESR, chalcogenide, intercalation, magnetic resonance.

INTRODUCTION

The transition metal phosphorus chalcogenides (M₂P₂S₆, M = transition metal) form a class of lamellar materials with crystal structures of the CdCl₂ type [1]. The layers are composed of arrays of [P₂S₆]₄⁻ units coordinated to the M²⁺ cations through M-S bonds, so that the transition metal ions are in a trigonally distorted octahedral environment. The transition metal layers are separated by two planes of chalcogenide atoms, bound together by relatively weak van der Waals interactions. Because of the weak nature of these interactions, various chemical species may be intercalated into the van der Waals gap (VWG) between the adjacent chalcogenide layers [2].

An unusual ion exchange process occurs in crystalline M₂P₂S₆ lattices in which the lattice cations are exchanged with intercalate cations [3]. The process is surprising in terms of the conventional view of transition metal chalcogenide lattices as covalently bonded, extended networks. In Mn₂P₂S₆, the Mn²⁺ can be exchanged at room temperature with mono- or dipositive guest cations (G⁺ or G²⁺) to form compounds with the composition Mn₂₋ₓGₓ⁺P₂S₆. EXAFS studies [4] have demonstrated that Ni²⁺ guest cations enter the intralamellar Mn²⁺ vacancies in Mn₂P₂S₆. Recently, Clement et al. [5] studied the exchange process by systematically exchanging the lattice cation in several of the M₂P₂S₆ lattices with K⁺. It was found that the ease of exchange followed the sequence Mn²⁺ > Cd²⁺ > Fe²⁺ and Ni²⁺ showed no exchange.

Previously [6], we have used EPR to study the exchange of diamagnetic Cd²⁺ cations in Cd₂P₂S₆ by paramagnetic Mn²⁺. The EPR analysis demonstrated that the guest cation is partitioned between interstitial sites in the VWG and host lattice Cd²⁺ vacancies. Heating the intercalated lattice is required to achieve insertion from the VWG into intralamellar vacancies. In the present work, EPR spectroscopy has been employed to examine the insertion of paramagnetic Co²⁺ into Cd₂P₂S₆. We have carried out the exchange reactions using different solvent media in order to qualitatively assess the importance of solvent coordination in the exchange process.

EXPERIMENTAL

Cd₂P₂S₆ was synthesized from the elements as described previously [7]. Crystals of Cd₂P₂S₆ containing about 1% by weight of Co²⁺ were obtained by the same method as for the undoped lattice, except that the required amount of dopant was also added to the reaction mixture. These crystals were used as standards for identification of intra-lamellar Co²⁺ residing on Cd²⁺ sites. In doped crystals prepared by sublimation at >650°C, the Co²⁺ cation resides exclusively on cadmium substitutional sites.

Co²⁺ cations were introduced into the Cd₂P₂S₆ lattice by a two step exchange procedure described by Clement and coworkers [5]. Large single crystals (typically 4 × 4 × 0.1 mm) of Cd₂P₂S₆ were immersed in a 1 M KCl solution at room temperature for 2–3 days. The solution was buffered to pH = 7 with
potassium bicarbonate and was approx. 0.1 M in EDTA. This procedure creates Cd\textsuperscript{2+} vacancies and inserts highly mobile solvated K\textsuperscript{+} cations into the interstitial sites in the VWG according to the reaction,

$$\text{Cd}_2\text{P}_2\text{S}_6(s) + 2xK^{+}(aq) \rightarrow H_{2}O\text{EDTA}$$

$$\text{Cd}_{2-x}K_x\text{P}_2\text{S}_6(s) + x\text{Cd}^{2+}(aq).$$

The EDTA assists in the extraction of Cd\textsuperscript{2+} by forming a strong complex with the metal cation. The product crystals were immersed in 1M aqueous solutions of CoCl\textsubscript{2} for 2 days in order to intercalate the host solid with Co\textsuperscript{2+} according to the following reaction

$$\text{Cd}_{2-x}K_x\text{P}_2\text{S}_6(s) + \text{Co}^{2+} \rightarrow H_{2}O$$

$$\text{Cd}_{2-x}\text{Co}_2\text{P}_2\text{S}_6(s) + 2xK^{+}(aq).$$

The solution was decanted and the crystals were washed a number of times with distilled water and then air-dried.

Crystals of Cd\textsubscript{2-x}Co\textsubscript{2}P\textsubscript{2}S\textsubscript{6} were also prepared by a single step procedure used previously for the exchange of Mn\textsuperscript{2+} with Cd\textsuperscript{2+} in Cd\textsubscript{2}P\textsubscript{2}S\textsubscript{6}. In this procedure, single crystals were immersed for 2-3 days at room temperature in a 1M CoCl\textsubscript{2} solution prepared with a solvent consisting of 80% by volume pyridine in water. The exchange may be represented as,

$$\text{Cd}_2\text{P}_2\text{S}_6(s) + x\text{Co}^{2+}(\text{pyridine}) \rightarrow \text{Cd}_{2-x}\text{Co}_2\text{P}_2\text{S}_6(s) + x\text{Cd}^{2+}(\text{pyridine}).$$

Because the product crystals may contain intercalated solvent, it was necessary to record EPR spectra after programmed heating cycles. Programmed heat treatment of the crystals was conveniently accomplished using a DuPont Instruments Model R90 thermal analyzer equipped with a Model 951 thermogravimetric analyzer. A nitrogen purge gas was used during heat treatment to avoid oxidation of the sample. EPR spectra were measured with a Bruker ER 200E-XRC X-band spectrometer with 100-kHz field modulation and a TM\textsubscript{10} cavity. The sample temperature was controlled with an Oxford Instruments Model ESR 900 continuous flow liquid helium cryostat.

**RESULTS**

An ESR spectrum obtained at 5 K from a Cd\textsubscript{2}P\textsubscript{2}S\textsubscript{6} crystal doped with 1% by weight of Co\textsuperscript{2+} during growth by vapor transport is reproduced in Fig. 1. This spectrum is observable below about 25 K and is substantially independent of temperature between 20 and 4K [8]. The spectrum is that of single Co\textsuperscript{2+} cations at Cd\textsuperscript{2+} vacancies. No EPR signal from pair spectra or interstitial ions was observed [9]. Experimental values of the spin-Hamiltonian parameters obtained from this spectrum are given in Table 1.

![Fig. 1. The EPR spectrum of Co\textsuperscript{2+} substitutional impurity centers in Cd\textsubscript{2}P\textsubscript{2}S\textsubscript{6} at T = 5 K. (a) H parallel to c* and (b) H perpendicular to c*.](image)

**Table 1.** ESR parameters for Co\textsuperscript{2+} in various coordination environments

<table>
<thead>
<tr>
<th>Compound (reference)</th>
<th>$g_1$</th>
<th>$g_2$</th>
<th>$\times 10^{-4}$ cm\textsuperscript{-1}</th>
<th>Site</th>
</tr>
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<tr>
<td>KFSt \textsuperscript{1}</td>
<td>5.80</td>
<td>3.44</td>
<td>184</td>
<td>47</td>
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<tr>
<td>Cd\textsubscript{2}P\textsubscript{2}S\textsubscript{6}</td>
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<td>3.99</td>
<td>183</td>
<td>97</td>
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<tr>
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<td>3.94</td>
<td>189</td>
<td>103</td>
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<tr>
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<td>3.94</td>
<td>199</td>
<td>118</td>
</tr>
<tr>
<td>MgO \textsuperscript{5}</td>
<td>4.28</td>
<td>4.28</td>
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<tr>
<td>CdO \textsuperscript{6}</td>
<td>4.37</td>
<td>4.37</td>
<td>Octahedral</td>
<td></td>
</tr>
<tr>
<td>Cd\textsubscript{2}P\textsubscript{2}S\textsubscript{6}</td>
<td>2.21</td>
<td>4.59</td>
<td>Trig. elong.</td>
<td></td>
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<tr>
<td>CdBr\textsubscript{2} \textsuperscript{7}</td>
<td>3.76</td>
<td>4.67</td>
<td>50</td>
<td>150</td>
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<tr>
<td>CdCl\textsubscript{2} \textsuperscript{8}</td>
<td>3.06</td>
<td>4.69</td>
<td>50</td>
<td>170</td>
</tr>
<tr>
<td>MgCl\textsubscript{2} \textsuperscript{8}</td>
<td>2.86</td>
<td>5.05</td>
<td>45</td>
<td>161</td>
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</tbody>
</table>

\textsuperscript{1} In potassium fluorosilicate [10].
\textsuperscript{2} Vapor doped.
\textsuperscript{3} Sharp spectrum.
\textsuperscript{4} Broad spectrum.
\textsuperscript{5} [17].
\textsuperscript{6} [18].
\textsuperscript{7} Calculated, this work.
\textsuperscript{8} [9].
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Fig. 2. The EPR spectrum of Co²⁺ intercalated cation in CdPzS₄ at T = 5 K.

Fig. 3. The EPR spectrum of Co²⁺ intercalated cation in CdPzS₄ at T = 10 K.

Fig. 4. Plot of the temperature dependence of the EPR site spectra obtained for Co²⁺ in CdPzS₄.

Fig. 5. The EPR spectrum of Co²⁺ intercalated cation CdPzS₄ at 5 K obtained after exchanging interstitial cations with K⁺.

temperature sensitivity of the EPR spectra of the ion exchanged lattice is markedly different from the behavior of the vapor doped lattice (Fig. 1).

The EPR spectra in Fig. 5 were obtained after briefly treating the sample crystals with a solution of 1 M KCl to replace interstitial Ca⁺⁺ cations with K⁺. After this treatment, the spectrum became independent of temperature between 4 and 20 K.

DISCUSSION

The EPR spectra (Figs 2 and 3) of Co²⁺ ions introduced into CdPzS₄ lattices by cation exchange are consistent with two, quite similar, cation coordination geometries. The magnitudes of the g-values and the g-value anisotropy are nearly identical and are very close to the g-value for substitutional Co⁺⁺ ions located at Cd⁺⁺ cation vacancies (Fig. 1). The greatest variation is observed in the value of g₁, suggesting that differences between the sites occur principally in the c* direction. The linewidth and temperature dependence of the site spectra are very different, indicative of different relaxation times for each site (see Fig. 4). Heating the crystalline samples to 600°C to remove intercalated solvent produced no significant changes in the EPR spectra.

The angle dependences of the EPR spectra of Co²⁺ observed after cation exchange indicate that the ions occupy lattice sites with their spin-axes aligned with respect to the crystallographic axes. Randomly oriented ions in the VWG would produce a powder EPR spectrum with peak absorption at g₁. There is no evidence of a powder EPR spectrum, therefore disordered ions are either not present or are EPR silent. The oriented ions may be located at either tetrahedral or octahedral interstitial sites within
the VWG or at octahedral cadmium ion vacancies. Further details concerning the coordination of the exchanged Co\textsuperscript{2+} ions can be extracted from the magnitude and anisotropy of the components of the g-tensor.

**g-Value anisotropy and coordination environment**

In tetrahedral coordination, the crystal field splitting of the \textsuperscript{4}F Co\textsuperscript{2+} free-ion ground term yields a \textsuperscript{4}A_2 ground state. The isotropic g-value is given approximately by,

\[
g = g_e - 8\lambda/10D_q. \tag{1}
\]

Setting the spin-orbit coupling parameter \(\lambda = -180\text{ cm}^{-1}\) and 10 \(D_q = 4300\text{ cm}^{-1}\) in eqn (1) gives \(g = 2.34\). No resonances are observed with \(g\)-values close to that expected for tetrahedral coordination. We conclude that the tetrahedral sites in the VWG or at octahedral cadmium ion vacancies are either not occupied or are EPR silent.

In the Cd\textsuperscript{2+} sites, the \textsuperscript{4}F ground state is split by a trigonally distorted octahedral crystal field and spin-orbit coupling. The octahedral crystal field component produces a splitting of approx. 20,000 \text{ cm}^{-1} into an orbital singlet (\textsuperscript{4}A_2) and two orbital triplets (\textsuperscript{4}T_{1g}, \textsuperscript{4}T_{2g}). The \textsuperscript{4}T_{1g} state lies lowest and is further split by spin-orbit coupling into six Kramers doublets. The isotropic g-value of the lowest Kramers doublet is given approximately by [10]

\[
g = (2/3)\alpha + 10/3 - 15\lambda/20D_q. \tag{2}
\]

For an F-state ion \(\alpha = 3/2\). If the value of \(D_q = 837\text{ cm}^{-1}\) obtained from the electronic spectrum of Co\textsubscript{6}P\textsubscript{6}S\textsubscript{6} is used in eqn (2), the isotropic g-value for octahedrally coordinated Co\textsuperscript{2+} is estimated to be \(g = 4.49\). The value estimated is in good agreement with the average g-value observed \(g_{\text{avg}} = 4.4\) (see Table 1) and we conclude that the EPR spectra arise from cations with predominately octahedral coordination. The anisotropy of the experimental g-values is indicative of a distorted octahedral Co\textsuperscript{2+} coordination. In distorted octahedral crystal fields, the g-tensor may become highly anisotropic and depend in a complex fashion on the details of the distortion. In the following, we employ a simple crystal field (point charge) model to describe the Co\textsuperscript{2+} coordination.

**Crystal-field model**

The EPR spectra of Co\textsuperscript{2+} are usually discussed in terms of the Abrahgam and Price model [10] in which the trigonally distorted octahedral coordination environment is described in terms of empirical values that are introduced as parameterizations of radial integrals arising from different terms in the spherical harmonic expansion of the crystal field (\(G \sim \langle r^6 \rangle\), \(H \sim \langle r^4 \rangle\), \(J \sim \langle r^2 \rangle\)). We have used an alternative parameterization of the trigonally distorted octahedral crystal field in terms of the three separable parameters: the trigonal angle (\(\theta\)), the familiar cubic field splitting parameter (\(D_q\)) and the trigonal field splitting parameter (\(C_p\)) [11].

In terms of these parameters, the crystal-field potential may be written as,

\[
V_{\text{cryst}} = 21\sqrt{\pi/5}C_p(3\cos^2 \theta - 1)Y_3^0 + 3\sqrt{\pi}D_q(35\cos^4 \theta - 30 + \cos^4 \theta + 3\lambda^2) + 3\sqrt{3\pi}(\sin^2 \theta - \cos \theta)(Y_2^2 - Y_4^0), \tag{3}
\]

where the \(Y_{\ell}^m\) are Legendre polynomials of order \(\ell\). The complete spin Hamiltonian is,

\[
H = V_{\text{cryst}} + H_{\text{Zeeman}}, \tag{4}
\]

where \(H_{\text{Zeeman}}\) is the magnetic field interaction given by:

\[
H_{\text{Zeeman}} = \beta H \cdot (\mathbf{L} + g_S \mathbf{S}) + i\lambda \mathbf{L} \cdot \mathbf{S}. \tag{5}
\]

Diagonalization of the matrix of the crystal-field Hamiltonian [eqn (4)] yields the energy levels of the ground state free-ion term split by the cubic and trigonal crystal-field components, spin–orbit coupling and the applied magnetic field. As the trigonal field splitting and the spin–orbit coupling are of the same order of magnitude, the Hamiltonian was diagonalized with both terms included.

A \(\textsuperscript{4}P\) state also arises from the \(d^7\) free ion and lies about 15,000 \text{ cm}^{-1} above the ground state. The \(\textsuperscript{4}P\) state must be included in calculations of the ground state splitting, because of its proximity to the ground state and the presence of matrix elements of the crystal field coupling the \(\textsuperscript{4}P\) and the \(\textsuperscript{4}F\) terms. Therefore, the crystal-Hamiltonian was diagonalized using the 26 functions of the decoupled basis \(L, M_L, S, M_S\) for the \(\textsuperscript{4}F\)-state \((L = 3, S = 3/2)\) and the 12 basis functions for the \(\textsuperscript{4}P\)-state \((L = 1, S = 3/2)\), constructed from one electron d-orbital functions. The values of other parameters used in the calculation were \(B\) (Racah) = 500 \text{ cm}^{-1} and \(\lambda\) (molecular spin–orbit coupling parameter) = −178 \text{ cm}^{-1}. Using the Hamiltonian [eqn (4)], the ground state g-value anisotropy was computed for trigonal angle values 50 < \(\theta\) < 58 and 2 < \(C_p/D_q\) < 12, corresponding to the full range of physically reasonable values of these parameters. The relationship between the g-value anisotropy \(\Delta g = (g_{||} - g_{\perp})\) and the parameters \(\theta\) and \(C_p/D_q\) is displayed as contour plots in Figs 6 and 7 for two values of the crystal field splitting.

The results of the calculation demonstrate that when \(C_p/D_q < 8\), the sign of the g-value anisotropy depends upon the sign of the trigonal distortion. When \(\theta < 54.7^\circ\) (trigonal elongation) \(g_{||} < g_{\perp}\) and when \(\theta > 54.7^\circ\) (trigonal compression) \(g_{||} > g_{\perp}\). The predicted behaviour is observed experimentally for Co\textsuperscript{2+} in a variety of trigonally distorted sites
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Fig. 6. g-Value anisotropy for Co²⁺ in a trigonally distorted octahedral coordination. The anisotropy \( \Delta g = g_\parallel - g_\perp \) is shown as a contour plot as a function of the parameters \( Cp/Dq \) and \( \theta \). Dq value held constant at 837 cm⁻¹.

When \( Cp/Dq \) takes larger values (>8) the dependence of the anisotropy upon the distortion angle becomes complex and the anisotropy alone cannot be used to determine the sign of the trigonal distortion.

The Co²⁺ sites in Cd₃P₂S₆ exhibit a g-value anisotropy \( \Delta g > +1 \). The results are unexpected, since in the pure M₃P₂S₆ lattices the metal coordination is trigonally elongated by about 2–3° [12]. Although antiferromagnetic alignment below \( T_N = 20 \) K prevents an experimental determination of the g-value anisotropy in Co₃P₂S₆, the calculated value is \(-2.33\), using parameter values obtained from infrared measurements of the trigonal splitting and X-ray diffraction [13] (\( Cp/Dq = 6\), \( \theta = 51.3^\circ \)).

Examination of the contour plots of the g-value anisotropy (Figs 6 and 7) reveals that \( \Delta g = +1 \) is consistent with either \( Cp/Dq < 2 \) and \( \theta < 54^\circ \), or a \( Cp/Dq > 5 \) and \( \theta > 55^\circ \). Thus, either \( Cp/Dq \) decreases dramatically from its value in Co₃P₂S₆ or the trigonal distortion changes sign. Insertion of a smaller cobalt ion into a cadmium vacancy (or into the VWG) results in a greater Co–S bond length than in Co₂P₂S₆. Both Dq and \( Cp \) decrease with increasing bond length, but they do not decrease at the same rate. The magnitude of the Dq parameter is proportional to the fourth-order radial one-electron integral and is given by,

\[
Dq = \frac{2e^2}{6a_0^5} \langle r^4 \rangle, \tag{6}
\]

where \( a_0 \) is the Bohr radius. \( Cp \) parametrizes second-order radial integrals of the form \( \langle r^2 \rangle \) and is given by,

\[
Cp = \frac{22e^2}{7a_0^3} \langle r^2 \rangle. \tag{7}
\]

Since \( Dq \) decreases more rapidly than \( Cp \) with increasing bond length, the behavior expected is for

Fig. 7. g-Value anisotropy for Co²⁺ in a trigonally distorted octahedral coordination. The anisotropy \( \Delta g = g_\parallel - g_\perp \) is shown as a contour plot as a function of the parameters \( Cp/Dq \) and \( \theta \). Dq value held constant at 700 cm⁻¹.
$C_p/D_q$ to increase as the bond length increases. We conclude that the experimental $g$-value anisotropy is consistent with Co$^{2+}$ located in trigonally compressed octahedral sites.

The $\Delta g = +1$ contour in Figs 6 and 7 requires a trigonal compression of at least 1° from octahedral coordination. The replacement of the larger Cd$^{2+}$ ion (ionic radius = 0.97 Å) by the smaller Co$^{2+}$ ion (ionic radius = 0.72 Å) without local lattice relaxation, leads to an increase in the trigonal angle (51.8°) by < 1°. In order to account for a 4° elongation from the original trigonal angle, considerable local lattice distortion about the Co$^{2+}$ must occur. It has been noted previously [14], that the P$_2$S$_6^{2-}$ groups exhibit a high degree of flexibility, allowing the first row M$_2$P$_2$S$_6$ materials to maintain essentially unchanged metal coordination geometry although the ionic radius varies substantially. This accommodation is achieved in the main part by variation of the P-P bond length and it is likely that this is the source of the local deformation of the Co$^{2+}$ coordination environment in Cd$_2$P$_2$S$_6$.

The magnitude of the atomic mean-square vibrational amplitudes in some M$_2$P$_2$S$_6$ phases suggests that there is a static disordering of the MS$_6$ octahedra. It has been suggested that this may arise through the existence of energetically equivalent minima for several MS$_6$ configurations [15]. The multi-site EPR spectrum of Co$^{2+}$ exchanged Cd$_2$P$_2$S$_6$ crystals indicates the presence of at least two nearly octahedral coordination environments. Since the site EPR spectra are similar to that of Co$^{2+}$ ions occupying Cd$^{2+}$ vacancies, it seems likely that the exchanged Co$^{2+}$ also occupy Cd$^{2+}$ vacancies. However, the low-temperature site interconversion indicates the presence of multiple potential minima for this site. The site responsible for the 4 K, broad EPR spectrum is evidently more labile since it is possible to effect its complete removal by exchanging the lattice with K$^+$ while leaving the second site spectrum undiminished (see Fig. 4).

Effect of solvent on the cation exchange process

We have studied the cation exchange reaction of Cd$_2$P$_2$S$_6$ using different paramagnetic ions (M$^{2+}$ = Co$^{2+}$, Mn$^{2+}$, Ni$^{2+}$) and two solvents (pyridine/H$_2$O and H$_2$O). The solvent system may play a significant role through the effect of solvation on the following schematic cation exchange equilibrium process,

$$\text{Cd}_2\text{P}_2\text{S}_6(\text{cryst.}) + x(M^{2+})_{\text{solv}} \rightleftharpoons \text{Cd}_{2-x}\text{M}_x\text{P}_2\text{S}_6(\text{cryst.}) + x(\text{Cd}^{2+})_{\text{solv}}$$

or equivalently, for the potassium exchanged host lattice,

$$\text{Cd}_{2-x}\text{K}_x\text{P}_2\text{S}_6(\text{cryst.}) + x(M^{2+})_{\text{solv}} \rightleftharpoons \text{Cd}_{2-x}\text{M}_x\text{P}_2\text{S}_6(\text{cryst.}) + 2x(K^+)_{\text{solv}}.$$  

The exchange of diamagnetic Cd$^{2+}$ for paramagnetic M$^{2+}$ located at the lattice cation vacancy site was followed by EPR. The results of these measurements are summarized in Table 2 and indicate that the stability constants of the solvated ions and the crystal field stabilization energies of the lattice ions exert comparable effects on the final composition of the ion exchanged lattice. The crystal field stabilization energies are:

$$\text{Mn}^{2+}(0) = \text{Cd}^{2+}(0) < \text{Co}^{2+}(8D_q) < \text{Ni}^{2+}(12D_q).$$

The M(py)$^{2+}$ stability constants at 25°C are [16]:

$$\text{Mn}^{2+}(1.38) < \text{Co}^{2+}(13.8) < \text{Cd}^{2+}(18.6) < \text{Ni}^{2+}(60.3).$$

On the basis of the crystal field stabilization, Co$^{2+}$ should enter the lattice vacancies more readily than Mn$^{2+}$; but, in the presence of pyridine the reverse is found, which we may understand in terms of the equilibria involving the solvent. The pyridine complex of Mn$^{2+}$ is much less stable than that of Co$^{2+}$, with the result that manganese occupies the lattice sites much more easily than cobalt when pyridine is used in the solvent medium. When, Mn$^{2+}$ is exchanged with Cd$_{2-x}$K$_x$P$_2$S$_6$ in aqueous medium, the solvated ion enters the VWG but does not enter lattice vacancies until the solvent is removed by

<table>
<thead>
<tr>
<th>Lattice</th>
<th>Ion</th>
<th>Solvent</th>
<th>EPR indication</th>
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<tr>
<td>Cd$_2$P$_2$S$_6$</td>
<td>Ni$^{2+}$</td>
<td>H$_2$O</td>
<td>No exchange</td>
</tr>
<tr>
<td></td>
<td>Co$^{2+}$</td>
<td>H$_2$O</td>
<td>No exchange</td>
</tr>
<tr>
<td></td>
<td>Mn$^{2+}$</td>
<td>H$_2$O</td>
<td>Rapid exchange at RT</td>
</tr>
<tr>
<td></td>
<td>Ni$^{2+}$</td>
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</tr>
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<td></td>
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<td>Pyridine</td>
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<td>Cd$_{2-x}$K$_x$P$_2$S$_6$</td>
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<td>H$_2$O</td>
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<tr>
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<tr>
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<td></td>
<td>Mn$^{2+}$</td>
<td>Pyridine</td>
<td>With heating to remove solvent</td>
</tr>
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</table>
heating. The large difference in the solvation energy may account for the ability to exchange \( \text{Cd}^{2+} \) for \( \text{Mn}^{2+} \) in pyridine solution without pre-exchange with \( \text{K}^+ \) [6]. This conclusion is supported by the observation that \( \text{K}^+ \) pre-exchange does not assist the insertion of \( \text{Mn}^{2+} \) into \( \text{Cd}^{2+} \) vacancies, but actually hinders it. \( \text{Ni}^{2+} \) is more strongly coordinated by pyridine than \( \text{Cd}^{2+} \) and will not exchange despite its large crystal field stabilization energy. Thus, solvation may reverse the effect of crystal field stabilization.

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