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THE GYPSUM—ANHYDRITE EQUILIBRIUM AT ONE ATMOSPHERE PRESSURE

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ABSTRACT

The equilibrium temperature for the reaction $\text{CaSO}_4 \cdot 2\text{H}_2\text{O} = \text{CaSO}_4 + 2\text{H}_2\text{O}_{(\text{lq, soln})}$ has been determined as a function of activity of $\text{H}_2\text{O (a}_{\text{H}_2\text{O}}$) of the solution. Synthetic gypsum and anhydrite or 1:1 mixtures were stirred in solutions of known $a_{\text{H}_2\text{O}}$ (calculated from vapor pressure data for the $\text{Na}_2\text{SO}_4$ and $\text{H}_2\text{SO}_4$ solutions), at constant temperature for as much as 12 months. The reversible equilibrium was approached from both sides and is defined by: $a_{\text{H}_2\text{O}} = 0.960$ at 55°, 0.845 at 39°, 0.770 at 23°C. Provided the solids do not change in composition, the equilibrium at constant $P$ and $T$ is a function of $a_{\text{H}_2\text{O}}$ only and is independent of the constituents in solution. Extrapolation to the bounding system $\text{CaSO}_4 - \text{H}_2\text{O}$ ($a_{\text{H}_2\text{O}} = 1.000$) yields 58°± 2°C. This is within thermodynamic calculations (46°± 2°C) but higher than solubility measurements (38° to 42°C). The new data indicate that in seawater saturated with halite and gypsum should dehydrate above 18°C. The scarcity of anhydrite in modern evaporite deposits is predicted by the present results. The available data on the temperature-salinity conditions under which anhydrite and gypsum exist in the Recent supratidal flat sediments of the Trucial Coast, Persian Gulf, are compatible with the present experimental data.

INTRODUCTION

The stability relations of gypsum ($\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$) and anhydrite ($\text{CaSO}_4$) are of considerable interest because most natural marine evaporite deposits consist essentially of gypsum and/or anhydrite interbedded with dolomite, limestone and clastic sediments (e.g., Stewart 1963, Table 18). In the binary system $\text{CaSO}_4 - \text{H}_2\text{O}$, the reaction $\text{CaSO}_4 \cdot 2\text{H}_2\text{O} = \text{CaSO}_4 + 2\text{H}_2\text{O}_{(\text{lq, soln})}$ has been studied experimentally at one atmosphere pressure by van't Hoff et al (1903), Partridge and White (1929), Toriumi and Hara (1934), Hill (1934), Posnjak (1938) and D'Ans et al (1955). Kelley, Southard and Anderson (1941) measured the thermochemical properties of the solid phases of the system at atmospheric pressure, and from these data calculated an equilibrium temperature for the gypsum-anhydrite transition. Marsal (1952), MacDonald (1953), Zen (1965) and Hardie (1965, pp. 25-30) calculated the effect of pressure on the reaction.

The effect of salt solutions on the gypsum-anhydrite equilibrium at atmospheric pressure has been considered in some detail by several workers, all of whom have verified that the transition temperature is lowered with increasing salinity. The system $\text{CaSO}_4 - \text{NaCl - H}_2\text{O}$ has been investigated experimentally by van't Hoff et al (1903), D'Ans et al (1955), Madgin and Swales (1956), Bock (1961) and Zen (1965). MacDonald

1 This paper is taken from a Ph.D. dissertation submitted by L. A. Hardie to the Department of Geology, The Johns Hopkins University, Baltimore, Maryland.
(1953) calculated the effect of NaCl solutions on the equilibrium temperature. The transition in sea water has been studied by Toriumi et al (1938) and Posnjak (1940). Other pertinent laboratory studies are those of Hill and Wills (1938) and Conley and Bundy (1958) in the system CaSO$_4$-Na$_2$SO$_4$-H$_2$O, D'Ans and Hofer (1937) in the system CaSO$_4$-H$_3$PO$_4$-H$_2$O, and Ostroff (1964), who converted gypsum to anhydrite in NaCl-MgCl$_2$ solutions.

For the most part the results of the different workers are in poor agreement. For this reason, and because the methods used were largely indirect (e.g. solubility and dilatometer measurements, thermodynamic calculation), a re-examination of the problem, using a different laboratory approach, seemed desirable.

The conversion of gypsum to anhydrite, and anhydrite to gypsum, was studied at atmospheric pressure as a function of temperature and activity of H$_2$O ($a_{H_2O}$). For the reaction

$$\text{CaSO}_4\cdot2\text{H}_2\text{O}_{(s)} \rightleftharpoons \text{CaSO}_4_{(s)} + 2\text{H}_2\text{O}_{(\text{liquid}, \text{in solution})}$$

the equilibrium constant may be defined in terms of activities, as follows:

$$K_a(p, T) = \frac{a_{\text{CaSO}_4}}{a_{\text{CaSO}_4\cdot2\text{H}_2\text{O}}}$$

If the standard states of the components of the reaction are considered to be pure H$_2$O liquid water, pure crystalline CaSO$_4$ and pure crystalline CaSO$_4\cdot2$H$_2$O at one atmosphere total pressure and at the temperature of reaction, the equilibrium constant simplifies to

$$K_a(T, p=1) = a_{H_2O}^2$$

It follows that the dehydration of gypsum to anhydrite, at atmospheric pressure, is a function of temperature and activity of H$_2$O only. Therefore, provided the solids do not change in composition, the equilibrium is independent of the components in the co-existing solution.

The activity of H$_2$O of the solutions co-existing with gypsum or anhydrite was varied by adding Na$_2$SO$_4$ or H$_2$SO$_4$: these were chosen because the gypsum $\rightleftharpoons$ anhydrite conversion rates were found to be relatively rapid in sodium sulfate or sulfuric acid solutions. In the system CaSO$_4$-Na$_2$SO$_4$-H$_2$O neither gypsum nor anhydrite can co-exist with a solution of Na$_2$SO$_4$ concentration greater than that fixed by the one atmosphere isothermally invariant assemblage gypsum (or anhydrite) + glauberite (CaSO$_4$·Na$_2$SO$_4$) + solution + vapor. The $a_{H_2O}$ of the invariant solutions varies from about 0.90 at 25°C to about 0.96 at 70°C. This limits the
study to very dilute solutions only, a considerable disadvantage because many natural calcium sulfate deposits must have formed in brines with activities of $\text{H}_2\text{O}$ at least as low as 0.75, as defined by the assemblage gypsum (or anhydrite) + halite + solution + vapor in the "haplo-evaporite" (Zen, 1965, p. 125) system $\text{CaSO}_4$-$\text{NaCl}$-$\text{H}_2\text{O}$. In the system $\text{CaSO}_4$-$\text{H}_2\text{SO}_4$-$\text{H}_2\text{O}$, however, the stability fields of gypsum and anhydrite are not limited by double-salt formation. Thus, the reaction may be studied in $\text{H}_2\text{SO}_4$ solutions which have a range of activity of $\text{H}_2\text{O}$ comparable to that found in natural waters, that is, from near 1.00 to about 0.70.

**Experimental Methods**

**Starting Materials.** The solid starting materials were artificial $\text{CaSO}_4$·$2\text{H}_2\text{O}$, $\text{CaSO}_4$ (anhydrous) and $\text{Na}_2\text{SO}_4$ (anhydrous) of reagent grade.

The $\text{CaSO}_4$·$2\text{H}_2\text{O}$ (Baker Analyzed, Lots No. 25692 and 25286) was fine-grained but variable-sized material (0.1 mm to less than 0.01 mm), which showed the characteristic morphology of gypsum euhedra. The X-ray diffraction pattern was indistinguishable from that of natural gypsum, and the material was used as such with no further treatment. The $\text{CaSO}_4$ (Baker Analyzed anhydrous [sic] Lot No. 90128) yielded an X-ray pattern consistent with bassanite $\text{CaSO}_4$·$n\text{H}_2\text{O}$ ($n \leq 0.5$). When this material was heated at 450°C to 550°C for 2 to 5 days, a very fine-grained powder was produced which gave an excellent anhydrite X-ray pattern. In the experimental runs with anhydrite as a starting material, the heat-treated $\text{CaSO}_4$ was used. Zen (1965, p. 151) found that artificial anhydrite, prepared by dehydrating gypsum overnight at 300°C, readily reverted to gypsum when brought into contact with water; he therefore considered such anhydrite unsuitable as a starting material. In the present investigation no such rehydration of artificial anhydrite occurred—even in stirred runs of 6 months duration—under conditions where anhydrite was considered to be stable. Critical experiments however, were repeated using natural gypsum and/or anhydrite. The gypsum was large clear selenite plates from Montmartre, Paris, and the anhydrite was massive fine-grained material from Richmond Co., Nova Scotia (Williams Collection, The Johns Hopkins University).

The $\text{Na}_2\text{SO}_4$ (Baker Analyzed, anhydrous, Lots No. 25581 and 22088) gave a sharp thenardite X-ray pattern and was used without further refinement.

The sulfuric acid solutions were prepared by diluting Baker Analyzed 95 percent $\text{H}_2\text{SO}_4$ with double-distilled water to the required concentrations. The exact concentration in weight percent $\text{H}_2\text{SO}_4$ was then deter-
mained by titration of carefully weighed aliquots of each solution against 1N NaOH solution (CO₂ free) using methyl orange as an indicator (Welcher, 1962, p. 540). The results were checked against the H₂SO₄ concentration determined by specific gravity measurements, using the calibration curve of Hodgman (1953, p. 1894).

Experimental procedure

(a) Static Method. At the start of the study a technique similar to that used in hydrothermal work was employed. Finely ground mixtures of anhydrite, or gypsum, and thenardite were accurately weighed, with the required amount of distilled water, into pyrex glass tubes (7×60 mm) which were then sealed using an oxy-acetylene torch. Loss of distilled water was successfully avoided during the sealing process by wrapping the tubes in wet filter paper. The sealed tubes were then totally immersed in thermostatically controlled water-baths. At the end of the run periods, which varied from several days to many months, the tubes were broken open and the solid products separated from the solution on absorbent paper. Samples were immediately examined, both under the microscope and by X-ray diffraction. Although by this technique many runs can be carried out simultaneously, it has the obvious disadvantage that the solution volumes are too small for analysis. Unfortunately, the method proved to have an even greater disadvantage; in many runs equilibrium was not attained even after periods of many months. When it was apparent that some form of agitation was required to promote the reactions, the static method was abandoned. However, it was possible to salvage enough significant information to warrant discussion and comparison with the results of runs carried out with the dynamic method (continuous stirring of the charge), which was used through the rest of the study.

(b) Dynamic Method. Approximately 200g of starting materials were weighed (±0.1%) into a 250-ml Erlenmeyer flask, fitted with a mercury-in-glass air-tight seal through which the solution was stirred sufficiently to keep all the solid material in constant agitation. The charged reaction vessels were immersed in water baths of capacity 30 liters, thermostatically controlled to ±0.1°C.

A maximum variation of ±0.5°C was observed over a period of six months.

At intervals, one milliliter of the suspension-charged solution was withdrawn with a pipette and rapidly pressure-filtered through a Buchner funnel: this removed almost all the solution. The solid material was immediately washed several times with acetone and air-dried. A portion of the sample was examined under the petrographic microscope; the re-
remainder was hand-ground under acetone and a smear mount prepared for x-ray diffraction.

**Determinations of activity of H₂O in solutions.** The activity of any constituent of a solution is given by the ratio of the fugacities:

\[ a_i = \frac{f_i}{f_i^0} \]

At one atmosphere total pressure, water vapor may be regarded as an ideal gas so that the fugacities may be safely replaced by the partial pressures of H₂O:

\[ a_{\text{H}_2\text{O}} = \frac{p_{\text{H}_2\text{O}}}{p_{\text{H}_2\text{O}}^0} \]

The standard state is taken as pure liquid water, at one atmosphere pressure and at the temperature of reaction, for which the activity is unity.

The solubilities of gypsum and anhydrite in sodium sulfate and sulfuric acid solutions are very low (less than 0.25 percent by weight). Therefore, the vapor pressures of these solutions saturated with CaSO₄ are given, within experimental measurement, by the vapor pressures of the CaSO₄-free solutions.

For the sodium sulfate solutions the vapor pressure data given in International Critical Tables (III, p. 371) were used to calculate \( a_{\text{H}_2\text{O}} \). The \( a_{\text{H}_2\text{O}} \) of each solution was computed at the temperature of each experiment. At the end of a run the total dissolved solids content of the equilibrium solution was determined. This value was compared with the starting Na₂SO₄ content to provide a check on the assigned \( a_{\text{H}_2\text{O}} \) value. The activities of H₂O of the sulfuric acid solutions were taken from Harned and Owen (1958, p. 574). These data, reproduced in Table 1, show that within the range of H₂SO₄ concentration used in the study, \( a_{\text{H}_2\text{O}} \) values determined by EMF measurements are in excellent agreement with those calculated from vapor pressure measurements. The initial gypsum and (or) anhydrite constituted only about 3 percent of the total charge. Consequently the H₂SO₄ concentration of the solution, and hence the activity of H₂O, was not significantly changed by the H₂O released or absorbed by the gypsum-anhydrite conversion. The H₂SO₄ content of the solution was checked by titration at the completion of each run.

The \( a_{\text{H}_2\text{O}} \) of the final solution in each of two runs (one in H₂SO₄ and one in Na₂SO₄) was measured directly using an H₂O-sensing apparatus (Hardie, 1965a, p. 252): the values did not differ measurably from the activities of H₂O determined in the CaSO₄-free solutions.
TABLE 1. ACTIVITY OF H₂O IN AQUEOUS SULFURIC ACID SOLUTIONS
(After Harned and Owen, 1958, p. 574)

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
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</tr>
</thead>
<tbody>
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<td>1</td>
<td>8.93</td>
<td>0.9620</td>
<td>0.9620</td>
<td>0.9624</td>
<td>0.9624</td>
<td>0.9630</td>
<td>0.9630</td>
</tr>
<tr>
<td>1.5</td>
<td>12.82</td>
<td>0.9391</td>
<td>0.9389</td>
<td>0.9402</td>
<td>0.9402</td>
<td>0.9415</td>
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<tr>
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<td>16.40</td>
<td>0.9136</td>
<td>0.9129</td>
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<td>0.9155</td>
<td>0.9180</td>
<td>0.9180</td>
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<tr>
<td>3</td>
<td>22.73</td>
<td>0.8506</td>
<td>0.8514</td>
<td>0.8548</td>
<td>0.8548</td>
<td>0.8602</td>
<td>0.8602</td>
</tr>
<tr>
<td>4</td>
<td>28.18</td>
<td>0.7775</td>
<td>0.7795</td>
<td>0.7850</td>
<td>0.7850</td>
<td>0.7950</td>
<td>0.7950</td>
</tr>
<tr>
<td>5</td>
<td>32.90</td>
<td>0.6980</td>
<td>0.7030</td>
<td>0.7086</td>
<td>0.7086</td>
<td>0.7229</td>
<td>0.7229</td>
</tr>
<tr>
<td>6</td>
<td>37.05</td>
<td>0.6200</td>
<td>0.6252</td>
<td>0.6288</td>
<td>0.6288</td>
<td>0.6505</td>
<td>0.6505</td>
</tr>
<tr>
<td>7</td>
<td>40.71</td>
<td>0.5453</td>
<td>0.5497</td>
<td>0.5608</td>
<td>0.5608</td>
<td>0.5815</td>
<td>0.5815</td>
</tr>
</tbody>
</table>

DESCRIPTION AND PROPERTIES OF THE SOLID PHASES

The gypsum synthesized by hydration of anhydrite (hereinafter referred to as "synthetic gypsum") commonly consists of thin plates flattened parallel to (010), with the characteristic monoclinic outline (Fig. 1). Between crossed nicols the thin plates show first-order white or grey interference colors and oblique extinction. In runs where gypsum was converted to anhydrite, the first stage of the process was recrystallization of the fine-grained artificial gypsum used as a starting material to coarser bladed crystals. This recrystallized gypsum, seemingly of a more

FIG. 1. Photomicrograph of synthetic gypsum, prepared from anhydrite at 40°C and 1 atmosphere in 22% H₂SO₄ solution, with seeding, in 50 days (Run LS 23). Average length of crystals 0.2 to 0.3 mm.
stable habit than the original smaller crystals, was morphologically and optically indistinguishable from the synthetic gypsum.

The optical properties of the synthetic gypsum were not significantly different from those of the natural material.

Wooster (1936) has determined the crystal structure of gypsum. In the ac plane two sheets of \( \text{SO}_4 \) tetrahedra are bound by Ca atoms within them. Between these sheets lie layers of water molecules. The Ca atoms are linked to 6 oxygens of \( \text{SO}_4 \) tetrahedra and to 2 water molecules. Water molecules thus occupy important structural positions and even partial dehydration must result in the destruction of the gypsum structure.

Unit cell parameters of the synthetic gypsum were determined by X-ray powder diffraction methods with either silicon or quartz as internal standard, and found to be in excellent agreement with those given by Deer et al. (1962) for natural gypsum.

Anhydrite synthesized from gypsum appeared under the microscope as a mass of minute birefringent grains; individual crystal outlines were barely distinguishable under high power, and refractive index measurements were unreliable.

The presence of synthetic anhydrite could be readily detected in the reaction flask by inspection: a fine white mass stayed in suspension long after the stirrer was stopped. In contrast, the well-crystallized synthetic and seed gypsum settled very rapidly, leaving, in the absence of anhydrite, a remarkably clear solution.

In one run (LS 51, Table 2), anhydrite grains as large as 0.3 mm across were synthesized from gypsum in 22 percent sulfuric acid solution at 50°C. They showed a stubby prismatic to equant shape with very high interference colors and parallel extinction.

The cell parameters of synthetic anhydrite were in excellent agreement with those given by Swanson et al. (1955).

Bassanite (\( \text{CaSO}_4 \cdot n\text{H}_2\text{O} \), \( n \leq 0.5 \)) was encountered only in static runs in which NaCl had been added to the charges. It was identified by X-ray diffraction techniques only, using the data of Posnjak (1938, p. 253). No distinction was made between calcium sulfate hemidydrate (\( \text{CaSO}_4 \cdot 1/2\text{H}_2\text{O} \)) and soluble anhydrite (\( \text{CaSO}_4 \)) because there is considerable uncertainty as to the relationship between these two phases (Deer, Howie and Zussman, 1962, v. 5, pp. 207–208).

**Experimental Results**

*Results of dynamic runs.* The data used to define the position of the \( a_\text{H}_2\text{O} \)-temperature equilibrium curve for the reaction
gypsum $\rightarrow$ anhydrite + $2\text{H}_2\text{O}$ (liquid, in solution)

are given in Table 2 and plotted in Figure 2.

Determination of the curve rested on the ability to convert gypsum to anhydrite, and vice versa, and on the ability to reverse the conversion when either one of the parameters was varied.

A run in which no change occurred in the starting phase, even after a reaction time of many months, was not considered evidence of stability of that phase, although in many cases such data provided confirmation of reversed runs. Taken by itself, a run in which no reaction occurs is at best inconclusive since metastable persistence of starting phases is commonly encountered in experimental studies of mineral equilibria. In the present study this was particularly true of the rehydration of anhydrite in sulfuric acid solutions, and induced nucleation by seeding proved necessary. Anhydrite remained unchanged in most unseeded runs for up to eight months; the addition of seeds of gypsum promoted relatively rapid conversion of the anhydrite (compare runs LS 5 and LS 14 at 35°C, Table 2). In the system CaSO$_4$-$\text{H}_2\text{O}$, the dehydration of gypsum to anhydrite has been shown to be incredibly slow (e.g. run AG 1, 70°C, Table 2) but, according to Posnjak (1938, p. 262) seeds of anhydrite do initiate the reaction. Considerable doubt has been thrown on the determination of stability by experiments in which seeds have been added to the charge because metastable growth of a phase from solution on seeds of its own kind is known to occur (Fyfe et al., 1958, p. 83). However, it has never been demonstrated that seeding would promote the disappearance of a stable phase and growth of a less stable one. In the present study, charges containing equal parts by weight of gypsum and anhydrite were used. The proof of stability in these seeded runs was growth of one phase and disappearance of the other. When the extent of reaction exceeded about 7 to 10 percent, metastable precipitation of either phase on seeds could be ruled out. This follows from a consideration of the solubilities of gypsum and anhydrite in Na$_2$SO$_4$ and H$_2$SO$_4$ solutions (maximum about 0.3 percent CaSO$_4$ by weight), the mass of solution in the reaction vessel (about 200–250 g) and the mass of excess starting solids (about 10 g). The extent of reaction was gauged by microscope and X-ray examination of a series of samples taken from the reaction vessel at intervals. From the X-ray diffraction patterns, the relative intensities of the 020 peak of anhydrite (3.499 Å) and the 140 peak of gypsum (3.065 Å) were measured. The amount of gypsum in each sample was read from a calibration

1 This was not substantiated in the present work, perhaps because the runs were not of sufficient duration. However, it was found that the presence of lime-water ($\text{pH} = 12.4$), with no seeding, markedly increased the dehydration rate where a seeded run in distilled H$_2$O showed no reaction (compare AG 1 and AG 5, 70°C, Table 2).
### Table 2. Experimental Data for Gypsum and Anhydrite Stability as a Function of Activity of H$_2$O and Temperature at Atmospheric Pressure. Results of Dynamic Runs Only

<table>
<thead>
<tr>
<th>Run no.</th>
<th>Starting materials</th>
<th>Solution (wt. %)</th>
<th>$\phi_{H_2O}$ soln</th>
<th>Temp. °C</th>
<th>Time days</th>
<th>Solid products</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>AG1</td>
<td>95g, 5a</td>
<td>100% H$_2$O</td>
<td>1.000</td>
<td>70°</td>
<td>359</td>
<td>95g, 5a</td>
<td>n.c.</td>
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<tr>
<td>AG5</td>
<td>100g</td>
<td>limewater</td>
<td>1.000</td>
<td>70°</td>
<td>193</td>
<td>93g, 7a</td>
<td>g-a</td>
</tr>
<tr>
<td>AG23</td>
<td>100g</td>
<td></td>
<td>0.965</td>
<td>70°</td>
<td></td>
<td>100g</td>
<td>g-a</td>
</tr>
<tr>
<td>AG25</td>
<td>100g</td>
<td></td>
<td>0.961</td>
<td>60°</td>
<td>46</td>
<td>42g, 58a</td>
<td>g-a</td>
</tr>
<tr>
<td>LS30-2</td>
<td>95g, 5a</td>
<td>limewater</td>
<td>0.955</td>
<td>55°</td>
<td>51</td>
<td>94g, 6a</td>
<td>g-a</td>
</tr>
<tr>
<td>LX49-2</td>
<td>100g</td>
<td></td>
<td>0.951</td>
<td>55°</td>
<td>35</td>
<td>55g, 45a</td>
<td>g-a</td>
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<tr>
<td>LS20</td>
<td>50g, 30a</td>
<td>9.43</td>
<td>0.941</td>
<td>50°</td>
<td>50</td>
<td>100g</td>
<td>n.c.</td>
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<tr>
<td>LS22</td>
<td>50g, 30a</td>
<td>4.03</td>
<td>0.960</td>
<td>50°</td>
<td>25</td>
<td>100g</td>
<td>n.c.</td>
</tr>
<tr>
<td>LS30-1</td>
<td>50g, 50a*</td>
<td>9.43</td>
<td>0.954</td>
<td>50°</td>
<td>47</td>
<td>100g</td>
<td>g-a</td>
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<tr>
<td>LS38</td>
<td>100g</td>
<td>15.0</td>
<td>0.953</td>
<td>50°</td>
<td>155</td>
<td>95g, 5a</td>
<td>349</td>
</tr>
<tr>
<td>LS49-1</td>
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<td>15.0</td>
<td>0.953</td>
<td>50°</td>
<td>47</td>
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<tr>
<td>LS52</td>
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<td>22.24</td>
<td>0.946</td>
<td>50°</td>
<td>99</td>
<td>100a</td>
<td>g-a</td>
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<tr>
<td>LS21</td>
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<tr>
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<td>9.56</td>
<td>0.945</td>
<td>50°</td>
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<td>100g</td>
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<tr>
<td>AG3</td>
<td>50g, 30a</td>
<td>9.56</td>
<td>0.956</td>
<td>50°</td>
<td>25</td>
<td>100g</td>
<td>n.c.</td>
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<tr>
<td>LS1</td>
<td>100g</td>
<td>15.0</td>
<td>0.953</td>
<td>50°</td>
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<td>95g, 5a</td>
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<td>LS16</td>
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<td>0.953</td>
<td>50°</td>
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<td>100g</td>
<td>n.c.</td>
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<td>0.953</td>
<td>50°</td>
<td>47</td>
<td>100g</td>
<td>g-a</td>
</tr>
<tr>
<td>LX11</td>
<td>100g</td>
<td>20.6</td>
<td>0.930</td>
<td>50°</td>
<td>23</td>
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</tr>
<tr>
<td>LS32-2</td>
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<tr>
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<tr>
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<tr>
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<tr>
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<tr>
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<td>g-a</td>
</tr>
<tr>
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<td>49</td>
<td>75g, 25a</td>
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</tbody>
</table>

**g**—gypsum  
**a**—anhydrite  
**gl**—glauberite  
**th**—thenardite  
**mr**—mirabilite  
**n.c.**—no detectable change  
**g-a**—natural gypsum and anhydrite  
**(*)**—trace
Starting materials
Solution (wt. %)
Na$_2$SO$_4$ H$_2$SO$_4$

<table>
<thead>
<tr>
<th>Run no.</th>
<th>Solid phases</th>
<th>Solution (wt. %)</th>
<th>Temp.</th>
<th>Time</th>
<th>Solid products</th>
<th>Result</th>
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<td>35°</td>
<td>80g, 20a</td>
<td>a-g</td>
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</tr>
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</tr>
<tr>
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<td>0.842</td>
<td>30°</td>
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<td>n.c.</td>
</tr>
<tr>
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<td>100a</td>
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<td>0.842</td>
<td>30°</td>
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</tr>
<tr>
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<td>0.773</td>
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<td>90g, 10a</td>
<td>a-a</td>
</tr>
<tr>
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</tr>
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</tr>
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<td>20°</td>
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<td>a, g, mc</td>
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<td>0.768</td>
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</table>

Fig. 2. The stability of gypsum and anhydrite determined experimentally as a function of temperature and activity of H$_2$O at atmospheric pressure. Only runs in which a conversion was achieved are plotted.
curve of I(anhdydrite)/I(gypsum) against weight percent gypsum with a reproducibility of better than 2 percent.

Because of the time limit, most of the runs were stopped before complete conversion of one phase to the other had occurred. Those runs which were allowed to react completely provided material for optical and X-ray studies.

Figure 2 shows that the experimental data, in general, are consistent. However, some exceptions must be noted. At 50°C, in 15 percent sodium sulfate solution ($a_{H_2O} = 0.953$), anhydrite started growing at the expense of gypsum after about 90 days reaction time (run LX 11, Table 2).

However, the reaction apparently stopped (!) as no further growth of anhydrite occurred in 10 months. Under the same conditions, anhydrite as a starting phase remained unaltered after 5 months (run LX 17). With sulfuric acid of about the same activity of $H_2O$, anhydrite was converted to gypsum at 50°C in a seeded run within a month (run LS 16), while gypsum was unchanged in an unseeded run of 9 months (run LS 1). The anomaly remains unexplained. An inconsistent result was also obtained at 55°C and $a_{H_2O} = 0.96$. In an unseeded run, gypsum was converted to anhydrite in sulfuric acid solution (run LS 8, about 50 percent reaction); in seeded runs both synthetic and natural anhydrite were transformed into gypsum (runs LS 22 and LS 30-1). These results are taken to indicate that the runs are very close to the equilibrium curve.

With the exception of run LX 11, then, the results obtained using sulfuric acid solutions are perfectly consistent with those obtained using sodium sulfate solutions.

The $a_{H_2O}-T$ values which define the equilibrium curve are given in Table 3.
Results of static runs. The results of experiments on the gypsum-anhydrite reaction carried out under static conditions are not included in Figure 2 or Table 2 because the method was soon abandoned in favor of agitated runs. A discussion of the data, however, is warranted. All the static runs were made with sufficient Na₂SO₄ to produce glauberite (CaSO₄·Na₂SO₄) as an additional phase. Synthetic anhydrite was the starting solid, although in a few runs natural anhydrite or synthetic gypsum were used instead. Seeds were not added to any of the charges. Reaction times varied from a few days to over 12 months. The results were most unexpected: anhydrite was found to rehydrate to gypsum at temperatures as high as 75°C. The reverse reaction, the dehydration of gypsum to anhydrite, was never achieved. Several possible explanations come to mind. First, the properties of the synthetic anhydrous CaSO₄ may be different from those of natural anhydrite. However, the results of the dynamic runs indicate that the differences, if any, are not significant. Another possibility is solid solution between gypsum and sodium sulfate. Unfortunately, this could not be checked by chemical analysis due to separation difficulties. However, no significant change in cell dimensions of the gypsum was observed, which suggests little or no substitution of Na⁺ for Ca⁡⁺⁺ in the gypsum structure. Indeed, a direct substitution is impossible since it would create a charge imbalance. The substitution of Na⁺ for Ca⁡⁺⁺ perhaps could be achieved if accompanied by an HSO₄⁻ for SO₄²⁻ substitution. Chemical analyses of natural gypsum and anhydrite (Stewart, 1963, p.33; Deer, Howie and Zussman, 1962, p. 206 and p. 221) show no evidence of this: sodium only occurs in trace amounts, if at all.

A third possibility is related to the experimental method. In the preparation of the charges, water was added to a solid mix of anhydrite + thernardite. Local high concentrations of sodium sulfate solution certainly existed, and probably persisted, in the initial stages of the runs. Conley and Bundy (1958) and Hardie (1965, p. 126) have shown that anhydrite reacts very rapidly with concentrated Na₂SO₄ solutions to form Ca-Na double sulfates. These double-salts are unstable in dilute Na₂SO₄ solutions (Hardie, 1965, p. 136; Hill and Wills, 1938, p. 1652) and decompose to gypsum and/or glauberite. In distilled water they immediately decompose to gypsum+Na₂SO₄ solution at all temperatures up to 100°C. It is possible, therefore, that in the static runs early formation of double-salts occurred in the regions of local high Na₂SO₄ concentration. This reaction removed anhydrite from the system. With time, diffusion led to a uniformly concentrated solution too low in Na₂SO₄ content for double-salt stability. Decomposition followed, giving gypsum+glauberite as products. Then, with prolonged time the gypsum should convert
to anhydrite. This is a plausible explanation because in all runs (static or stirred) in which the starting anhydrite was added to pre-mixed, uniformly concentrated sodium sulfate solution, no anomalous formation of gypsum was observed.

The work of Conley and Bundy (1958) is pertinent here since they proposed essentially this mechanism for the conversion of anhydrite to gypsum in salt solutions. They suggested that the reaction for the conversion with activator solutions such as sodium or potassium sulfate is:

$$\text{CaSO}_4 \rightarrow^{\text{act.}}_{\text{H}_2\text{O}} \text{Ca}^{++} + \text{SO}_4 = \text{H}_2\text{O} \rightarrow \text{CaSO}_4 \cdot 2\text{H}_2\text{O}$$

and is primarily dependent upon temperature and concentration. However, they achieved the conversion only by washing the reaction products with water. This in fact really only demonstrates that anhydrite will react rapidly at low temperatures and high alkali sulfate concentrations (see Conley and Bundy, 1958, Figs. 1 and 2) to form double-salts which decompose in water to give gypsum and salt solution. It certainly does not prove, as they maintained in the abstract of their paper (p. 57), that “contrary to recent hypothesis of gypsum dehydration by concentrated salt solutions, double salts and/or gypsum are stable phases below a temperature of 42°C.”

Comments on the mechanism of dehydration of gypsum to anhydrite. Three different mechanisms by which gypsum in contact with an aqueous medium could dehydrate to anhydrite appear possible:

1. a solution-precipitation process.
2. direct dehydration to anhydrite (loss of structural water).
3. step-wise dehydration through the intermediate hydrate, bassanite.

The present experimental results throw some light, albeit very diffuse, on the problem.

In a few of the runs in which anhydrite was produced from gypsum, a rind, presumed from X rays to be anhydrite, was observed on the surface of, and along cleavage cracks in, gypsum crystals (Fig. 3). It is conceivable that the rind material is bassanite, formed as an intermediate dehydration product, but no trace of this phase was identified in the charges by X rays even though perhaps as many as 10 to 15 percent of the gypsum grains showed some alteration. If the rind is anhydrite, the process would appear one of direct dehydration to anhydrite beginning at the crystal surfaces where H$_2$O may be transferred to the solution phase. The effect of seeding on the dehydration rate could not be gauged with any certainty because too few duplicate runs were made. However, in
those runs which were strictly comparable (e.g. LS 17 and LS 12 at
25°C; LS 51 and LS 32-2 at 50°C, Table 2), no significant rate increase,
which would have suggested a solution-precipitation mechanism, was
observed. In this respect the results of Zen's (1965) precipitation exper-
miments loom large: direct precipitation of anhydrite from super-saturated
solutions could not be achieved at temperatures up to 70°C, even with
seeding. Similar attempts by the present author also failed.

![Photomicrograph of a dark rind of anhydrite (?) on a prismatic grain of gypsum](image)

Fig. 3. Photomicrograph of a dark rind of anhydrite (?) on a prismatic grain of gypsum (colorless). Note incipient development of rind material in cleavage crack of large, colorless (recrystallized) gypsum plate. Data: Run LS 51, gypsum stirred in 22% H₂SO₄ solution at
50°C and 1 atm for 211 days. About 50% conversion to anhydrite at end of run. No reac-
tion was observed until 146 days. Magnification about 250X.

always precipitated under conditions where anhydrite was presumed
stable. This inability to precipitate anhydrite must remain the most
telling evidence against a solution-precipitation mechanism for the de-
hydration of gypsum to anhydrite in aqueous media. On the other hand,
the reverse reaction, the hydration of anhydrite, may well be accom-
plished through a solution-precipitation process because the hydration
rate is measurably increased (in sulfuric acid solutions, at least) by the
addition of seeds (compare runs LS 7 and LS 15 at 45°C; LS 5 and LS 14
at 35°C; LS 6 and LS 4-3 at 30°C; Table 2).

Ostroff (1964) observed the formation of calcium sulfate hemihydrate
as an intermediate step during the conversion of gypsum to anhydrite in
sodium-magnesium chloride solutions at 90.5°C. That this is not the
GYPSUM—ANHYDRITE EQUILIBRIUM

invariable mechanism is proved by the present experiments: bassanite was not formed in any of the sodium sulfate or sulfuric acid runs. With NaCl solutions, however, bassanite rather than anhydrite was the dehydration product of gypsum (Hardie, 1965, Table 30, p. 185). Zen (1965) also found bassanite instead of anhydrite in concentrated NaCl solutions. Apparently sodium chloride solutions promote a step-wise dehydration process. The picture remains unclear and an exhaustive study of the kinetics of the reaction using more sensitive methods is obviously needed.

Comparison of the Results with Previous Work

Extrapolation of the measurements of the present study to solutions in the system CaSO₄·H₂O (a₂H₂O = 1.00) gives a transition temperature of 58° ± 2°C, slightly lower than that of van’t Hoff et al. (1903) but significantly higher than the oft-quoted temperature of 42° ± 2°C given by Posnjak (1938) from solubility measurements. This discrepancy was puzzling, at first, because the solubility data seemed well supported by the thermodynamic data of Kelley et al. (1941), who calculated an equilibrium temperature of 40°C. Zen (1962), however, pointed out that these calculations employed internally inconsistent data. Recalculation put the transition temperature at 46° ± 21°C, and Zen considered the agreement with the solubility measurements as fortuitous, a conclusion supported by the present calculations (Appendix). Indeed, re-examination of the solubility data indicates that the value of 42°C is by no means securely established. In Figure 4 are compiled all the available solubility measurements on gypsum and anhydrite in the system CaSO₄·H₂O; in large part the older data were taken from the tables of D’Ans (1933, pp. 203–205). If curves are drawn to enclose the maximum density of each set of points, a transition temperature of anywhere between 38° and 50° is indicated (see shaded area in Fig. 4). An uncertainty may well be real, since, as Zen (1965, p. 126) pointed out, all the available measurements were made by approaching the equilibrium solubility curves from the side of undersaturation only. This is, of course, a serious drawback in any solubility study but is particularly crucial in the case of poorly soluble substances. For example, Backström (1921) measured the solubility of calcite and aragonite by approaching the equilibrium curves from both sides: he found that after a few days the rate of change of the solubility had reached zero but that the supersaturation and undersaturation values differed by as much as 5 percent. Alexander et al. (1954) and Krauskopf (1956) obtained similar results for amorphous silica: even greater deviations were the case here. There is no guarantee, then, that the equilibrium saturation value of either gypsum or anhydrite had been reached in any except Zen’s (1965) gypsum determinations. The available
solubility data therefore must be regarded as yielding minimum values only. The argument is crucial, for if the points plotted in Figure 4 for anhydrite are indeed minimum values then the transition point must lie at some temperature above at least 44°C, Zen’s (1965) data being taken as the upper limit of gypsum solubility. Hill (1934, 1937) did recognize the necessity for approaching a solubility curve from both sides, and, indeed, reported his anhydrite values as obtained from both undersaturated and supersaturated solutions. The data, however, were extrapolated from solubilities measured in potassium sulfate solutions because he was unable to achieve supersaturation with respect to anhydrite using pure water at temperatures below 65°C. At this and higher temperatures he apparently was successful, but, unfortunately, he neither described the procedure nor reported direct precipitation of anhydrite from solution.
A similar criticism applies to the available gypsum and anhydrite solubility measurements made in salt or acid solutions, and probably explains why the results of different workers are in such poor agreement. Comparison of the results of these workers is most easily made by computing the activity of H$_2$O of the solutions reported to be in equilibrium with gypsum±anhydrite. Because the CaSO$_4$ content of these solutions is very low, activity of H$_2$O may be obtained with considerable accuracy from the water vapor pressures of the CaSO$_4$-free salt or acid solutions. The vapor pressure data for the NaCl and Na$_2$SO$_4$ equilibrium solutions were taken from the International Critical Tables (1928, III, pp. 370–371) and those of sea water from Arons and Kientzler (1954). For the phosphoric acid solutions the vapor pressure values of Kablukov and Zagwosdkin (1935) were used. The results are summarized in Table 4 and shown graphically in Figure 5.

The values of van't Hoff et al. (1903) alone are higher than those of the

<table>
<thead>
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<th>Investigator</th>
<th>Trans. temp. °C</th>
<th>Co-existing solution</th>
<th>$a_{H_2O}$</th>
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<td>50°</td>
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<td>45°¹</td>
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<tr>
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<td>31.0% H$_3$PO$_4$</td>
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<tr>
<td>Bock (1961)</td>
<td>35°</td>
<td>11.85% NaCl</td>
<td>.920</td>
</tr>
<tr>
<td>Zen (1965)</td>
<td>35°</td>
<td>15.25% NaCl</td>
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</tr>
<tr>
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<td>6.09% NaCl</td>
<td>.963</td>
</tr>
<tr>
<td>Bock (1961)</td>
<td>30°</td>
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</tr>
<tr>
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<tr>
<td></td>
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<td>0.82% MgSO$_4$</td>
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<tr>
<td>Bock (1961)</td>
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<td>18°</td>
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<td>.751</td>
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¹ Extrapolated by the present author.
² The results of these two studies were taken from Seidell (1958, pp. 665–667).
present study. The approach of van't Hoff and his co-workers to the problem was brilliantly conceived. They first partially converted gypsum to anhydrite in saturated sodium chloride solution in a dilatometer at 70°C. The rate of conversion of gypsum to anhydrite and vice versa at different temperatures was then measured by the rate of change of volume of the contents of the dilatometer. At the equilibrium temperature there should be no volume change. They observed a volume decrease at 25°C and an increase at 35°C, so the transition temperature was taken as 30°C. The partial pressure of H₂O of the solution co-existing with gypsum, anhydrite and halite at 30°C was then measured, giving a value of 24 mm. The method was repeated using saturated sodium bromate solution and a transition temperature of 50°C was obtained; the water vapor pressure of the equilibrium solution was measured as 83.3 mm. Using these two points in the relationship

\[ \log p = \log p^\circ + \frac{A}{T} - B/T \]

they were able to extrapolate to solutions in the system CaSO₄·H₂O. Implicit in this approach is the principle that the gypsum-anhydrite transition is independent of the constituents in solution and that the equilibrium temperature is a function of the ratio \( p/p^\circ \). This is, of course, the principle on which the present study is based!

Figure 5 also shows that although the results of different workers are in very poor agreement, they all fall within the limits predicted by the thermodynamic calculations (Appendix)!
It has been suggested by some workers that most, if not all, calcium sulfate of natural evaporites was originally deposited as gypsum (Posnjak, 1940; Bundy, 1956; Conley and Bundy, 1958; Murray, 1964; Zen, 1965). The argument is based on (1) petrographic observations that much anhydrite is pseudomorphous after twinned gypsum, (2) the scarcity of anhydrite in modern evaporites deposits, and (3) experimental evidence that anhydrite cannot be synthesized under pressure-temperature conditions consistent with natural evaporite environments.

It is clear that the present relationship between gypsum and anhydrite in the pre-Recent marine evaporites of the world, to a great extent, is secondary, due to the effects of post-depositional burial. Gypsum at surface may be traced downward into anhydrite so that at depths below about 2000–3000 feet gypsum is practically absent (MacDonald, 1953; Stewart, 1963). Evidence of replacement is abundant. Anhydrite pseudomorphous after twinned gypsum has been reported by Schaller and Henderson (1932) in the Salado formation of Texas and New Mexico, by Stewart (1953) in the Permian evaporites of Yorkshire, England, and by Borchert and Baier (1953) in the German Zechstein. At shallower depths gypsum has been shown to have replaced anhydrite (Stewart, 1953; Goldman, 1952; Ogniben, 1955; Sund, 1959); such replacement has been recorded recently at a depth as great as 3500 feet (Murray, 1964). Therefore, both gypsum and anhydrite in sedimentary deposits may be metamorphic but this evidence does not prove that the replaced gypsum, or anhydrite, was primary in origin, a point emphasized by Zen (1965, p. 147).

More significant evidence is provided by the distribution of gypsum and anhydrite in Recent marine and nonmarine evaporites where effects due to burial are not involved. In these deposits gypsum is ubiquitous and, in all certainty, primary (Bramkamp and Powers, 1955; Morris and Dickey, 1957; Masson, 1955; Phleger and Ewing, 1962; Wells, 1962; and others) whereas anhydrite has been reported from only one locality, in supratidal flat sediments on the Trucial Coast, Persian Gulf (Curtis et al., 1963; Kinsman, 1964).¹ This single occurrence of Recent sedimentary gypsum—Anhydrite Equilibrium

Geological Implications

It has been suggested by some workers that most, if not all, calcium sulfate of natural evaporites was originally deposited as gypsum (Posnjak, 1940; Bundy, 1956; Conley and Bundy, 1958; Murray, 1964; Zen, 1965). The argument is based on (1) petrographic observations that much anhydrite is pseudomorphous after twinned gypsum, (2) the scarcity of anhydrite in modern evaporites deposits, and (3) experimental evidence that anhydrite cannot be synthesized under pressure-temperature conditions consistent with natural evaporite environments.

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¹ Other occurrences of Recent anhydrite have been reported recently by Hunt et al. (1966, p. 59) as a surface layer in Death Valley, California, and by Moiola and Glover (1965) from a sediment dump on Clayton Playa, Nevada. In both cases the anhydrite has formed from gypsum, via bassanite, in the absence of a liquid phase. This dehydration process

\[
\begin{align*}
\text{CaSO}_4 \cdot 2\text{H}_2\text{O} & \rightleftharpoons \text{CaSO}_4 \cdot \frac{3}{2}\text{H}_2\text{O} + \frac{3}{2}\text{H}_2\text{O} (g) \\
\text{CaSO}_4 \cdot \frac{3}{2}\text{H}_2\text{O} & \rightleftharpoons \text{CaSO}_4 + \frac{3}{2}\text{H}_2\text{O} (g)
\end{align*}
\]

involves a set of thermochemical conditions very different from those encountered in "wet" evaporite deposits.
Anhydrite is of great importance because, if not evidence of primary precipitation of anhydrite, it is, at least, proof that metamorphism of gypsum on burial is not essential to anhydrite formation. Even accepting that anhydrite can be primary, an inconsistency between the field evidence from modern evaporites and available experimental evidence exists. Temperature-salinity conditions necessary for anhydrite stability, as predicted from solubility experiments and thermodynamic calculations, are commonly found in modern evaporite environments, yet gypsum is the common phase of such deposits. This observation, coupled with the inability of experimenters to synthesize anhydrite at low temperatures, has led workers such as Murray (1964) to conclude that gypsum is always the primary precipitate. Further, under conditions theoretically favoring anhydrite, this gypsum will persist metastably (except where temperatures at surface are very high) until burial causes dehydration to anhydrite.

The present experimental results have a two-fold bearing on the problem. First, it is demonstrated that anhydrite can be synthesized at one atmosphere pressure under geologically reasonable conditions of temperature and activity of H$_2$O in a geologically reasonable time, reckoned in months. Primary precipitation of anhydrite, however, could not be achieved, indeed, has not been achieved by previous workers. This would suggest, but, of course, not prove, that gypsum is always the first formed CaSO$_4$ phase on evaporation of natural waters. Be that as it may, the experiments do show that gypsum, maintained in the stability field of anhydrite, would be dehydrated to anhydrite soon after deposition. Second, it is demonstrated that higher temperatures than previously were entertained are required for anhydrite formation. This, qualitatively, is more in keeping with the observation that gypsum is the common phase in Recent evaporites.

Quantitative application of the experimental results to natural deposits is valid and possible but is hindered by the paucity of precise information on the temperature and $a_{H_2O}$ of natural solutions co-existing with gypsum and anhydrite. Fortunately, the Persian Gulf deposit is an important exception. Quantitative data have been collected by D. J. J. Kinsman (personal communication, 1964). Brine temperatures range from 24$^\circ$ to 39$^\circ$C and anhydrite is found in carbonate muds of the sabkha, or supratidal salt flat, where ground water chlorinities exceed about 130$^\circ$/oo (about 24 percent salinity). Part of the anhydrite deposit

---

1 It is possible, of course, that natural waters differ from experimentally tested solutions in that they contain additional components which would induce direct precipitation of anhydrite, as, for example, trace elements ("impurities") have been found to influence the nucleation of aragonite and calcite (Wray and Daniels, 1957).
GYPSUM—ANHYDRITE EQUILIBRIUM

Table 5. Data on Solutions Co-existing with Gypsum or Anhydrite in Natural Evaporite Deposits

<table>
<thead>
<tr>
<th>Locality</th>
<th>Mineral Assemblage</th>
<th>Solution</th>
<th>Investigator</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trucial Coast, Persian Gulf</td>
<td>gypsum, carbonate</td>
<td>24°C</td>
<td>64-91</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>96</td>
</tr>
<tr>
<td></td>
<td>anhydrite, carbonate</td>
<td></td>
<td>134&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>152&lt;sup&gt;1&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>159</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>166</td>
</tr>
<tr>
<td>Boccana de Virrila, Peru</td>
<td>gypsum</td>
<td>27°C</td>
<td>108&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saline Valley, Calif.</td>
<td>gypsum</td>
<td>9-39°C</td>
<td>16-50</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.99-0.95</td>
</tr>
<tr>
<td>Salina fm., Mich. (Silurian)</td>
<td>anhydrite, halite&lt;sup&gt;3&lt;/sup&gt;</td>
<td>32-48°C</td>
<td>satd. NaCl &lt;0.75</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<sup>1</sup> Anhydrite in the zone of capillary draw. Chlorinities as given are for the underlying groundwaters. Actual solutions in which anhydrite formed presumably were more concentrated.

<sup>2</sup> Precise location in Boccana where gypsum is precipitating is difficult to read from Morris and Dickey's descriptions. Value given here is taken from their data for location C, which may be incorrectly interpreted by the present author as the gypsum site.

<sup>3</sup> Delicately preserved “hopper” crystals which have clearly not suffered alteration since their formation. The same argument must apply to the intimately associated anhydrite. Temperatures of formation of the hopper halite was determined by fluid inclusion studies.

occurs in the zone of capillary draw above the present water table but, more important, anhydrite is found in direct contact with brine (Table 5). Where ground waters are less concentrated (up to about 96<sup>°</sup>/oo chlorinity) gypsum is precipitated within the carbonate muds. These data, together with the limited information from a few other deposits, are summarized in Table 5. To compare these data, chlorinity values
have been recalculated to activities of H₂O using the vapor pressure data for seawater of different chlorinities given by Arons and Kientzler (1954). The natural brine data and the equilibrium curve determined in the present study are plotted in Figure 6. Included in this figure is a gypsum-anhydrite transition curve computed from the solubility measurements of Bock (1961) in the haplo-evaporite system CaSO₄-NaCl-H₂O. His results are considered representative of the stability range for gypsum and anhydrite predicted by most existing solubility studies (see Fig. 5). If the natural deposits are to be interpreted in terms of this transition curve, then it is clear from Figure 6 that metastable persistence of gypsum in natural brines is the rule. On the other hand, the equilibrium curve of the present study is remarkably compatible quantitatively with the data from the natural deposits, particularly that of the Persian Gulf (Fig. 6). This would strongly suggest that chemical equilibrium prevails in each of these deposits. Taken one step further, this could mean that the scarcity of anhydrite in modern evaporite deposits is simply a reflection that the conditions for its formation are seldom reached, or, at least, maintained for any length of time. Metastable persistence of gypsum would not be a necessity.

While this is most plausible it is surely an oversimplification because gypsum is found under nonequilibrium conditions in some modern evaporitic environments. For example, in Laguna Ojo de Liebre, Baja California, gypsum co-exists with halite at temperatures up to 27°C (Phleger and Ewing, 1962) whereas the present experimental data predict that in a seawater brine saturated with halite (aH₂O ≤ 0.75) gypsum should dehydrate to anhydrite at temperatures above about 18°C (see Fig. 2).

The questions this discussion raises are intriguing. Does the Persian Gulf, where anhydrite is forming, combine a freakish set of chemical and/or physical circumstances not found in other modern evaporite environments? Or, are the conditions under which gypsum is found in Baja California (and perhaps other areas) not maintained for long enough periods of time each year to produce anhydrite? It is clear that the problem is one of kinetics which must, therefore, become a most important consideration in interpreting gypsum-anhydrite deposits, modern or ancient.

**Summary**

1. The present results show that anhydrite can be synthesized experimentally from gypsum under p, t and aH₂O conditions reasonable for natural evaporite environments.
2. The gypsum-anhydrite equilibrium temperatures determined in the
Fig. 6. Temperature-activity of H₂O relations of brines co-existing with gypsum and anhydrite in natural evaporite deposits. Experimentally determined equilibrium relations are shown as curves: solid curve: this study; dashed curve: solubility data of Bock (1961). The open symbols represent solutions in contact with gypsum and the solid symbols are those in contact with anhydrite.

- **Open circle:** Morris & Dickey (1957)
- **Squares:** Bramkamp & Powers (1955)
- **Triangles:** Hardie (1965)
- **Solid circles:** Dellwig (1955)
- **Bars:** Kinsman (pers. comm., 1964)

Present study are considerably higher than those based on solubility measurements and on thermodynamic calculations. The new data are considered more reliable than the existing data because (a) the present results are based on reversible reactions whereas in available solubility studies the saturation curves for gypsum and anhydrite were approached only from the side of undersaturation and, therefore, do not necessarily represent equilibrium curves; (b) significant uncertainties exist in the available thermochemical values for gypsum and anhydrite.
3. The new equilibrium values of this study are more compatible with the field observations that gypsum is the common phase, and anhydrite rare, in unmetamorphosed evaporite deposits. Further, these values are quantitatively consistent with the $a_{H_2O} - T$ conditions under which gypsum and anhydrite are found in the Recent evaporite deposit of the Trucial Coast, Persian Gulf.

**Acknowledgments**

Professor Hans P. Eugster suggested the project, guided the research and critically reviewed the manuscript. To him, I am deeply indebted. The manuscript was also reviewed by Owen P. Bricker, James L. Munoz and George B. Skippen and their comments led to a considerable improvement in the manuscript. Mr. Arthur Everhart designed and built the relays used to control the water-bath units. A special tribute must be paid to the late Thomas Banks (The Johns Hopkins University and Harvard University) for helpful discussions and stimulation during the latter part of the study.

The research was supported, in part, by Grant 680-A from the Petroleum Research Fund to The Johns Hopkins University.

**Appendix. Thermodynamic Considerations**

_Calculation of the gypsum-anhydrite transition temperature in the system CaSO₄·H₂O at one atmosphere pressure._ Kelley, Southard and Anderson (1941) measured the thermochemical properties of the solid phases of the system CaSO₄·H₂O (Table 6).

For the reaction

$$\text{CaSO}_4 \cdot 2\text{H}_2\text{O} = \text{CaSO}_4 + 2\text{H}_2\text{O}$$

Kelley et al. (1941, p. 44) obtained

$$
\Delta G_T^\circ = -2495 - 65.17 T \log T + 0.0215 T^3 + 163.89 T
$$

(1)

This equation gives 313 K (40°C) as the temperature at which gypsum, anhydrite and liquid water are in equilibrium at one atmosphere total pressure, in surprisingly good agreement with the value of 42°C derived from solubility data (Hill, 1937; Posnjak, 1938).

Zen (1962; 1965) has pointed out that equation (1) was obtained from inconsistent data. The present calculations confirm Zen’s criticism. Differentiation of equation (1) with respect to temperature, $-d\Delta G^\circ/dT = \Delta S^\circ$, yields

$$
\Delta S_T^\circ = -135.59 + 65.17 \log T - 0.043 T
$$

(2)

and a value of 12.85 cal/deg for 298 K. This is inconsistent with the sum of the individual entropies at 298 K as given by Kelley et al. (see Table 6), that is,

$$\Delta S_{298}^\circ = 25.5 + (2 \times 16.8) - 46.4 = 12.7 \text{ cal/deg}$$

The discrepancy arises from the use of an integration constant of -33.18 which is the mean of their own consistent value of -33.03 and the value of -33.34 obtained from the indirect vapor pressures of Toriumi and Hara (1934). The resulting small error in entropy has a con-


Gypsum—Anhydrite Equilibrium

Table 6. Thermochemical Properties of the Phases of the System CaSO₄·H₂O

<table>
<thead>
<tr>
<th>Phase</th>
<th>Cᵥ, cal/deg mole (298°-450°K)</th>
<th>S° (298°K) cal/deg mole</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaSO₄·2H₂O</td>
<td>21.84+0.0767T</td>
<td>46.4 ± 0.4</td>
<td>I (p. 36 &amp; p. 19)</td>
</tr>
<tr>
<td>CaSO₄</td>
<td>14.10+0.0337T</td>
<td>25.5 ± 0.4</td>
<td>I (p. 36 &amp; p. 19)</td>
</tr>
<tr>
<td>H₂O (liquid)</td>
<td>18.02</td>
<td>16.8</td>
<td>II (p. 105)</td>
</tr>
<tr>
<td>H₂O (gas)</td>
<td>7.45+0.0027T</td>
<td>45.13</td>
<td>II (p. 105)</td>
</tr>
</tbody>
</table>

Referencel

I = Kelley et al. (1941)
II = Kelley (1950)
III = Kelley (1960)

Siderable effect on the free energy values since it is incorporated into the aT term of equation (1).

Recalculation of the entropy for the reaction using the accepted value of S°₂₉₈ = 16.75 cal/deg-mole for liquid water (Giauque and Stout, 1936; Kelley, 1960) yields ΔS°₂₉₈ = 25.5 + (2×16.75) = 12.6 cal/deg.

Hence, equation (2) becomes

\[ ΔS_T^o = -135.84 + 65.17 \log T - 0.043 T \]  

Using (Kelley et al, 1941, p. 44),

\[ ΔH_T^o = -2495 + 28.30 T - 0.0215 T^2 \]  

the free energy expression becomes

\[ ΔG_T^o = -2495 + 164.14 T - 65.17 T \log T + 0.0215 T^2 \]  

This equation gives 319°K (46°C) as the gypsum-anhydrite equilibrium temperature, an increase of 6°C over the value obtained from the expression of Kelley et al (1941) (Equation (1)).

The free energy expression can be further modified by employing the revised heat capacity of anhydrite given by Kelley (1960, p. 46). The following relationships for the reaction are then obtained:

\[ C_v = 31.02 - 0.0524 \ T \]

and

\[ ΔH_T^o = ΔH_2^o + 31.02 T - 0.0262 T^2 \]  

---

1. I = Kelley et al. (1941)
   II = Kelley (1950)
   III = Kelley (1960)
Substituting the mean value, $\Delta H_{298}^0 = 4030$ cal$^1$ (Kelley et al, 1941), of the heat of solution measurements in equation (7):

$$\Delta H_{298}^0 = -2890 \text{ cals}$$

so that

$$\Delta H_{298}^0 = -2890 + 31.02T - 0.0262T^2$$  \hspace{1cm} (7)

Using

$$\Delta S_{298}^0 = 25.5 + (2 \times 16.75) - 46.4 = 12.6 \text{ cal/deg. in}$$

$$\Delta S_{298}^0 = \Delta S_{298}^0 + 31.02 \ln T - 0.0524T$$

we obtain

$$\Delta S_{298}^0 = -148.55 + 31.02 \ln T - 0.0524T$$  \hspace{1cm} (8)

From equations (7) and (8) it follows that

$$\Delta G_{298}^0 = -2890 + 179.57T + 0.0262T^2 - 71.44T \log T$$  \hspace{1cm} (9)

This equation gives $T(\text{equil.}) = 46^\circ C$, demonstrating that the equilibrium temperature is insensitive to small variations in the heat capacity of anhydrite.

If the uncertainties of measurement assigned to each one of the thermodynamic values (Table 6) used in the derivation of equation (9) are assembled, then the confidence to be placed in this equation can be assessed. From the maximum and minimum possible values of each property, except the heat capacity, we obtain two limiting free energy equations:

$$\Delta G_{298}^0 = -2870 + 180.43T + 0.0262T^2 - 71.44T \log T$$  \hspace{1cm} (10)

and

$$\Delta G_{298}^0 = -2910 + 178.71T + 0.0262T^2 - 71.44T \log T$$  \hspace{1cm} (11)

Equation (10), designed to give a maximum temperature, was derived using $\Delta H_{298}^0 = 4030 + 20 = 4050$ cal and $\Delta S_{298}^0 = 25.1 + (2 \times 16.72) - 46.8 = 11.74$ cal/deg. The equivalent values for equation (11), which gives a minimum temperature, are $\Delta H_{298}^0 = 4030 - 20 = 4010$ cal and $\Delta S_{298}^0 = 25.9 + (2 \times 16.78) - 46.0 = 13.46$ cal/deg.

The equilibrium temperatures given by equations (10) and (11) are $68^\circ C$ and $25^\circ C$ respectively. It is clear that the available thermochemical data can fix the gypsum-anhydrite transition point at no better than $46 \pm 22^\circ C$!

Calculation of the effect of dissolved salts on the gypsum-anhydrite transition temperature at atmospheric pressure. For the conversion of gypsum to anhydrite in the presence of any aqueous solution containing dissolved salts, the reaction may be written

$$\text{CaSO}_4 \cdot 2\text{H}_2\text{O} = \text{CaSO}_4 + 2\text{H}_2\text{O}_{(\text{liq.}, \text{soln.})}$$

A simplified equilibrium constant for this reaction can be applied if the compositions of the solid phases remain unchanged (pure liquid H$_2$O and the pure solids at 1 atmosphere being taken as the standard states):

$$(K_a)_{p=1,T} = a_{2\text{H}_2\text{O}}^2$$

$^1$ Kelley et al (1941, p. 15) give $\pm 20$ cals. as the uncertainty in this value.
Now, for any chemical reaction

$$\Delta G_T = \Delta G_T^0 + RT \ln K_a$$

Therefore, for the dehydration of gypsum to anhydrite

$$\Delta G_T = \Delta G_T^0 + 2RT \ln a_{H_2O}$$  \hspace{1cm} (12)

It follows from this equation that a lowering of the activity of $H_2O$ of the solution (e.g., by increasing the salinity) would decrease the free energy of reaction. The effect would be to lower the dehydration temperature. To evaluate this quantitatively it is necessary to know $\Delta G_T^0$ (reaction) and the $a_{H_2O}$ of the solutions in which the reaction occurs.

An expression for $\Delta G_T^0$ (reaction) as a function of temperature has been derived (equation 9 above):

$$\Delta G_T^0 = -2890 + 179.57 T + 0.0262 T^2 - 71.44 T \log T$$  \hspace{1cm} (9)

Therefore, the change in equilibrium temperature with change in $a_{H_2O}$ of the co-existing solution can be determined from

$$\Delta G_T = -2890 + 179.57 T + 0.0262 T^2 - 71.44 T \log T + 2RT \ln a_{H_2O}$$  \hspace{1cm} (13)

An expression similar to (13) but based on the free energy equation of Kelley et al (1941) (equation 1), was derived by MacDonald (1953, p. 889) using a slightly different thermodynamic treatment:

$$\Delta G_T = -2495 + 163.89 T + 0.0215 T^2 - 65.17 T \log T + 2RT \frac{2.303 \log p/p^o}{(14)}$$

From this, MacDonald determined the transition temperature as a function of concentration of sodium chloride.

Kelley et al (1941, Fig. 8, p. 41) also had considered the effect of activity of $H_2O$ on the gypsum-anhydrite transition temperature. They presented the results in diagrammatic form only and did not give the equation used in the calculation. This, however, is most certain to be the equation given by MacDonald, who used their data and produced exactly equivalent results.

Equation (13) gives 20°C and equation (14) 15°C for the transition temperature in the presence of halite in the system CaSO$_4$-NaCl-H$_2$O ($a_{H_2O}$=0.75). The uncertainties in these temperatures will remain in the order of ±22°C, the uncertainty range for the solution of equation (9).

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