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Synthesis and Thermal Properties of Strontium and Calcium Peroxides

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SYNTHESIS AND THERMAL PROPERTIES OF STRONTIUM AND CALCIUM PEROXIDES

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SUMMARY

This report describes the synthesis and discusses some thermal properties of strontium and calcium peroxides (SrO₂ and CaO₂). Our interest in these peroxides is founded on their use in the preparation of the new high-temperature superconductors. In other reports we have demonstrated several advantages in substituting BaO₂ for BaCO₃ in the preparation of the YBa₂Cu₃O_{7-x} bulk superconductor. The required, high-purity BaO_2 is easily made by heating BaO in dry O_2 , and it is commercially available. However, SrO_2 and CaO_2 , which would be used in the newer superconductors, are best synthesized by first precipitating the peroxide octahydrate from a cold, aqueous ammoniacal Sr or Ca salt solution with dilute H_2O_2 . Anhydrous SrO_2 is conveniently prepared by heating the octahydrate at 115 °C for 24 hr, and the less stable anhydrous CaO₂ is best synthesized by drying the octahydrate at room temperature in a dry box containing an effective desiccant such as P_2O_5 . The yields of anhydrous SrO_2 and CaO_2 by this method were about 66 percent based on the amount of Sr or Ca salt used. The yield of both peroxides may be increased to about 95 percent by adding a strong base, such as NaOH, which neutralizes the ammonium salt by-product. SrO₂ is considerably more thermally stable and less susceptible to hydrolysis and CO_2 pickup than CaO_2 . In addition, we give a new x-ray diffraction pattern for $CaO_2 \cdot 8H_2O$ from which we calculated the lattice parameters a = 6.212830and c = 11.0090 on the basis of the tetragonal crystal system.

INTRODUCTION

In preparing the high-temperature superconductor $YBa_2Cu_3O_{7-X}$, we found that substituting barium peroxide, BaO_2 , for the commonly used barium carbonate, $BaCO_3$, produced a better bulk superconductor in terms of greater sample homogeneity and density and improved reproducibility (ref. 1). Our success in using BaO_2 as the barium source in the $YBa_2Cu_2O_{7-X}$ formulation prompted us to investigate the synthesis and properties of strontium and calcium peroxides, SrO_2 and CaO_2 , for the preparation of the newer high-temperature superconductors involving these alkaline earth oxides.

The advantages of using alkaline earth metal peroxides $(BaO_2, SrO_2, and CaO_2)$ in place of the corresponding carbonates are briefly outlined as follows:

(1) Less rigorous reaction conditions are required in preparing superconductors because the carbonates are considerably more stable than the corresponding peroxides. (2) There is no carbon in the starting materials; thus, it should be possible to produce a carbonate-free superconductor. Deposits of nonconducting carbonate in the grain boundaries of bulk superconductors are believed to decrease the critical current.

(3) Peroxides provide an oxidizing media that favors the formation of higher oxidation states of metals, especially copper. In the $YBa_2Cu_3O_{7-X}$ superconductor, the oxygen stoichiometry must be greater than that required for the normal valency if superconductivity is to take place (ref. 2).

(4) Decomposition of peroxides liberates oxygen instead of the carbon dioxide liberated during the decomposition of carbonates. With peroxides, processing becomes simpler: a muffle furnace can be used to fire compositions containing peroxides because it is not necessary to use flowing oxygen during firing to remove liberated CO_2 . Liberated CO_2 must not remain in the sample environment because it will be taken up by the material during cooling, resulting in carbonate formation. As stated before, carbonate deposites have a detrimental effect on superconducting properties.

(5) One desirable attribute of BaO_2 is that it melts before it decomposes. This affords a liquid-solid phase reaction that is more efficient than the totally solid-solid phase reaction obtained with $BaCO_3$. The more efficient liquid-solid phase reaction with BaO_2 may explain the greater homogeneity that we observed in the bulk $YBa_2Cu_3O_{7-X}$ superconductor when BaO_2 was used instead of $BaCO_3$ (ref. 1).

(6) The anhydrous peroxides CaO_2 , SrO_2 , and BaO_2 are less reactive and have better storage capabilities than their respective normal oxides CaO, SrO, and BaO. The alkaline earth oxides are strong bases and, as such, tend to pick up acid gases in the atmosphere, such as CO_2 , to form salts (e.g., carbonates). In addition, the oxides react vigorously with water to form hydroxides. On the other hand, the anhydrous peroxides, being less basic than the oxides, are practically inert toward CO_2 pickup under ambient conditions. They also do not appreciably react with water at room temperature to produce hydroxides.

Alkaline earth peroxides undergo three general reactions as shown. With calcium peroxide used to illustrate these reactions, they are as follows:

Decomposition	$CaO_2 \rightarrow CaO + \frac{1}{2}O_2$	(1)
Dehydration	$CaO_2 \cdot 8H_2O \rightarrow CaO_2 + 8H_2O$	(2)

Hydrolysis $CaO_2 \cdot 8H_2O \rightarrow Ca(OH)_2 + \frac{1}{2}O_2 + 7H_2O$ (3)

Table I gives the calculated standard enthalpy of reaction ΔHF° from the National Bureau of Standards Tables (ref. 3). Unfortunately, the free energy of formation ΔGF° data for the alkaline earth peroxides and their octahydrates are not available; thus ΔHF° values are used instead to explain reaction trends. For all three of the peroxides and their octahydrates, the decomposition, dehydration, and hydrolysis reactions are endothermic - unlike some peroxides, which can decompose exothermically to form the oxide, for example, hydrogen peroxide

 $H_2 O_2 \rightarrow H_2 O_1 + \frac{1}{2} O_2 \qquad \Delta HF^\circ = -98.0 \text{ kJ mol}^{-1}$ (4)

The thermodynamic data for the anhydrous peroxides indicate that they are more stable than the corresponding oxides. Thus these peroxides may be stored safely without danger of violent decomposition.

There are two general methods for synthesizing peroxides (ref. 4). The first involves heating the oxide in a stream of pure, CO_2 -free dry oxygen. This method is favored for making peroxides that are considerably more stable than the corresponding oxide - for example, for the preparation of BaO₂. In fact, BaO₂ is commercially made by heating BaO at 500 °C in flowing, pure oxygen. The synthesis of the less stable SrO_2 requires more drastic conditions, namely heating the oxide at a temperature of 350 °C in a bomb containing high-pressure oxygen (250 atm). Finally, CaO₂, the least stable of the three, cannot be conveniently synthesized by direct combination of oxide with oxygen.

The second technique, which is more suitable for the preparation of less stable peroxides involves precipitating the insoluble peroxide from aqueous solution by adding H_2O_2 to a basic solution of the metal salt. Again using CaO₂ as the example,

 $CaCl_2 + H_2O_2 \rightarrow CaO_2$ (hydrate) + 2HCl

 $2HC1 + 2NH_3 \rightarrow 2NH_4C1$

Addition of aqueous ammonia to neutralize the HCl, forces the reaction to favor the precipitation of the peroxide hydrate.

This report presents various aspects of the preparation of both the hydrated and anhydrous calcium and strontium peroxides using the second synthesis technique. Since the chemistry of CaO_2 is more complicated because it is less stable than SrO_2 , it was studied in greater detail.

The thermal decomposition properties of SrO_2 and CaO_2 were also investigated. A new x-ray diffraction powder pattern was determined for $CaO_2 \cdot 8H_2O$, and lattice parameters were calculated from it. Because BaO_2 is commercially available as a pure anhydrous powder, no effort was made to investigate the synthesis and chemical properties of this compound.

EXPERIMENTAL PROCEDURE

The synthesis of strontium and calcium peroxides was accomplished by a modification of the method outlined in Brauer's Handbook of Preparative Chemistry (ref. 4). Calcium chloride dihydrate (CaCl₂·2H₂O) or strontium nitrate (Sr(NO₃)₂ (0.4 mol)) was dissolved in 1.5 liters of deionized water. An excess of aqueous ammonia (ammonium hydroxide, NH₄OH), 2 mol in 600 ml water, was added to the stirred salt solution. No precipitated Ca or Sr salts (carbonates) were observed. In all of our cases, the solution remained clear. The solution was adjusted to the desired temperature by either heating it in a water bath or cooling it in an ice bath. Because of the ease with which the CaO₂·8H₂O loses water even under water, the octahydrate was precipitated in ice

cold solution. To maintain the solution temperature at close to 0 °C, the ammoniacal CaCl₂ solution was kept in the freezer until some ice formed. To the well-stirred, cold solution, ice cold 3 percent H_2O_2 containing a slight excess of H_2O_2 (0.45 mol) was added at a rate of about two drops per second. The pearly white, crystalline hydrated peroxide was allowed to settle, and excess solution was removed via decantation. The crystals were filtered using a suction flask, sucked as dry as possible, washed with about 200 mQ water containing several milliliters of NH₄OH, and then, again, sucked as dry as possible. After the peroxide was removed from the filter, it was either air dried by allowing it to stand at ambient conditions in the laboratory or dried under specified conditions (see tables II and IV). The SrO₂ and the other CaO₂ preparations were synthesized in the same way except at different temperatures. For these preparations, the Sr(NO₃)₂ or CaCl₂ solution was maintained at the indicated temperature while the H_2O_2 solution was added at room temperature.

We took special care to prevent the $CaO_2 \cdot 8H_2O$ from dehydrating so that we could generate a new x-ray powder diffraction pattern of this compound. It was important that $CaO_2 \cdot 8H_2O$ remain as the pure octahydrate throughout the manipulation to ensure that the pattern was a true representation of the octahydrate. After filtration, the $CaO_2 \cdot 8H_2O$ crystals were kept moist. The moist peroxide was stored in a tightly capped plastic bottle that was kept cold in a freezer. The diffraction pattern of the wet crystals was recorded immediately after the octahydrate was removed from the freezer.

In other experiments involving $CaO_2 \cdot 8H_2O$, the moist crystals were removed from the filter and treated as indicated in the Results section. In most cases, further treatment involved heating the crystals at various temperatures in a drying oven or drying them in a flowing-nitrogen dry box containing P_2O_5 drying agent in a dish.

The peroxide yield was considerably less than stoichiometric based on the limiting reagent $CaCl_2 \cdot 2H_2O$, thus indicating some reverse reaction with the NH₄Cl by-product. We found that adding a strong base (NaOH solution) to increase pH, increased the yield to near stoichiometric. This was accomplished by the dropwise addition of 500 mL of a solution containing 0.75 mol of NaOH to the well-stirred precipitated peroxide suspension just following the addition of H₂O₂. Slightly less than the stoichiometric amount of NaOH was added so as to prevent the precipitation of calcium compounds other than $CaO_2 \cdot 8H_2O$, namely $Ca(OH)_2$ and $CaCO_3$. If sodium contamination presents a problem, an organic strong base such as tetramethylammonium hydroxide ((CH₃)₄NOH) may be substituted for NaOH.

The anhydrous CaO_2 was also directly precipitated from aqueous solution. The previously described method was employed except that it was done at higher solution temperature (e.g., 70 °C).

 $SrO_2 \cdot 8H_2O$ was synthesized in a fashion similar to that for $CaO_2 \cdot 8H_2O$. However, because of its lesser tendency to dehydrate, it was unnecessary to keep the $SrO_2 \cdot 8H_2O$ moist and cold. The diffraction pattern was taken on the dried and heat treated samples and compared with the known patterns for the octahydrate and anhydrous SrO_2 . We determined the purity and degree of hydration of both peroxides by firing them for 12 hr at 1000 °C in air, thereby converting the strontium and calcium compounds to their normal oxides, SrO and CaO. X-ray diffraction was used to identify various components resulting from different runs. These components include peroxides, hydroxides, carbonates, and oxides. Particle morphology was investigated with scanning electron microscope (SEM) and optical photomicrographs. Thermogravimetric analysis (TGA) runs in helium (30 cm³/min) at a heating rate of 10 °C/min established the decomposition temperatures of anhydrous SrO₂ and CaO₂.

RESULTS

Synthesis of SrO_2

Addition of H_2O_2 to ammoniacal $Sr(NO_3)_2$ solution (as described in the experimental section) produced pearly white, crystalline $SrO_2 \cdot 8H_2O$. The crystals settled rapidly and were readily isolated by filtration. The theoretical yield was about 66 percent based on the amount of $Sr(NO_3)_2$ starting material. Table II gives the effects of various treatments on the degree of hydration of $SrO_2 \cdot XH_2O$. The value of X in the table corresponds to the degree of hydration, X = moles $H_2O/moles SrO_2$. This value is calculated from the weight loss when $SrO_2 \cdot XH_2O$ is fired to SrO. Values of X less than 8 probably represent a mixture of $SrO_2 \cdot 8H_2O$ and anhydrous SrO_2 rather than the presence of a definite hydrated peroxide with less than eight waters of hydration.

Freshly precipitated $SrO_2 \cdot 8H_2O$ loses water of hydration gradually when it sits in a crystallizing dish at ambient laboratory conditions (about 21 °C). After 2 days the value of X drops from 8.00 to the range 7.05 to 7.70. The x-ray diffraction pattern, however, showed only the presence of the octahydrate. When freshly precipitated $SrO_2 \cdot 8H_2O$ is placed in a dry box at ambient temperature with flowing dry nitrogen and with P_2O_5 drying agent, water of crystallization is gradually lost. After 5 days the degree of hydration decreases to 0.29, and, according to x-ray diffraction, anhydrous SrO₂ becomes the major component. On drying in air at 100 °C for 24 hr, only a small amount of water (X = 0.14) remains, and at 115 °C for 24 hr, pure anhydrous SrO_2 is produced. Up to this point, no decomposition products $(Sr(OH)_2 \text{ or } SrO)$ were observed. And, at this point, x-ray diffraction showed anhydrous SrO2 as the sole component of the material. Chemical analysis gave 86.94 percent SrO (86.62 percent is the theoretical value), and the results of atomic adsorption analysis were also in good agreement: 73 percent Sr (whereas the theoretical value is 73.25 percent Sr). For superconductor formulations where a pure SrO₂ of known composition is required, we recommend drying the freshly precipitated hydrated SrO_2 in air at 115 °C for 24 hr. Raising the drying temperature to 125 °C resulted in a calculated negative value of X = -0.15. This denotes that in addition to complete dehydration, possibly a small amount of the normal oxide SrO was present probably because of a slight decomposition of the peroxide. TGA analysis indicated that anhydrous SrO₂ begins rapid decomposition at about 400 °C.

The theoretical yield of SrO_2 was increased from 66 percent to about 95 percent by neutralizing most (about 94 percent) of the NH_4NO_3 reaction byproduct with the strong base, NaOH, thereby increasing the pH of the solution:

$NH_4NO_3 + NaOH = NaNO_3 + NH_3 + H_2O$

We expect an increase in pH to drive the overall SrO_2 -producing reaction to the right, thereby favoring the SrO_2 yield:

 $Sr(NO_3)_2 + H_2O_2 + 2NH_3 + 8H_2O = SrO_2 \cdot 8H_2O + 2NH_4NO_3$

Neutralization with NaOH was not investigated further.

Synthesis of CaO_2

The relative instability of CaO_2 , especially when compared with SrO_2 , warranted a more extensive investigation into the chemistry of CaO_2 . It is the least stable of the alkaline earth peroxides, especially with regard to hydrol-ysis, decomposition, and loss of water of hydration.

Table III lists the degree of hydration (value of X in the formula $CaO_2 \cdot XH_2O$) when the peroxide is precipitated at various solution temperatures. It is evident that the average degree of hydration of the precipitated CaO₂ depends on the solution temperature. When the precipitation is carried out at 0 °C, pearllike crystals of the octahydrate form. Like the SrO₂·8H₂O precipitate, the insoluble, hydrated CaO₂ crystals settle rapidly and are easily filtered. At this time, we attribute the unknown x-ray diffraction pattern to $CaO_2 \cdot 8H_2O$. In a later part of this report, this supposition is further substantiated. As the precipitation temperature is increased, especially above 50 °C, the beige-colored anhydrous peroxide precipitates. The insoluble anhydrous peroxide is less crystalline and is more difficult to filter. For precipitation above 50 °C, the x-ray diffraction pattern shows only the presence of the anhydrous peroxide. On the basis of the value of X, practically pure anhydrous CaO2 is precipitated directly at the maximum temperature of our experiments, 70 °C. An important aspect to be considered in the synthesis of CaO₂ is that the peroxide yield decreases with increasing precipitation temperature; this yield is calculated from the amount of CaCl₂.8H₂O used as the limiting reagent. The yield ranges from 67.3 percent at 0°C to 51.0 percent at 70 °C. As was the case with the deposition of SrO₂.8H₂O described earlier, increasing the pH of the solution by neutralizing 90 percent of the NH_4Cl by-product with aqueous NaOH increased the calcium peroxide yield in cold solution to about 95 percent of theoretical based on the amount of $CaCl_2 \cdot 2H_2O$ starting material. Again, the results of neutralization experiments were not studied further.

Figure 1 presents micrographs of the octahydrate and anhydrous calcium peroxide. Parts (a) and (b) show the octahedral crystal plates of freshly precipitated $CaO_2 \cdot 8H_2O$ from cold solution (0 °C). Figure 1(a) shows transparent octahedral crystals; figure 1(b) shows the same material at higher magnification. More detail can be seen of the morphology of the eight-sided plates of $CaO_2 \cdot 8H_2O$. Parts (c) and (d) show the CaO_2 that was precipitated from hot solution (70 °C). It was surprising that the CaO_2 precipitated in the form of spheres instead of well-defined crystals. Such a phenomenon suggests that when the H_2O_2 solution was added to the hot ammoniacal $CaCl_2$ solution, the octahydrate $CaO_2 \cdot 8H_2O$ formed first, then melted in its own water of crystallization with the formation of small suspended spherical droplets. As the hydrated water was lost, the liquid droplets solidified into spheres of anhydrous CaO_2 . A liquid phase was observed when the octahydrate was heated to 90 °C in the oven prior to thermal dehydration to the anhydrous peroxide. It is apparent that this liquid phase is molten hydrated CaO_2 .

The final investigation into calcium peroxide chemistry concerned its thermal decomposition. The results (table IV) illustrate the ease with which hydrated CaO_2 loses water of hydration. It is difficult to obtain the pure octahydrate because attempts to remove excess absorbed water from wet, freshly prepared $CaO_2 \cdot 8H_2O$ resulted in some dehydration. The original sample used in the thermal decomposition studies was octahydrate freshly precipitated from cold solution at 0 °C and dried at room temperature overnight. The composition of this material was CaO₂·7.4H₂O. To illustrate the ease with which this material loses water, we left a sample of $CaO_2 \cdot 7.4H_2O$ standing for 5 days on a lab bench. It lost most of its water of hydration, and the x-ray diffraction pattern showed only the presence of anhydrous CaO_2 . The same material when placed in a flowing-nitrogen dry box containing P2O5 drying agent for 48 hr at room temperature lost practically all of its hydrated water (table IV, X = 0.11). Thus, it is not surprising that thermal dehydration takes place at elevated temperatures (table IV). Above 100 °C some decomposition (eq. (3)) takes place as indicated by $Ca(OH)_2$ showing as a minor component. At the highest drying temperature in the table, 120 °C, decomposition of the hydrated peroxide to $Ca(OH)_2$ followed by CO_2 pickup from the atmosphere becomes important. These reactions are indicated by $CaCO_3$ showing as a minor phase in the x-ray diffraction pattern. From these results, it is apparent that $CaO_2 \cdot 8H_2O$ should be dehydrated in a dry, CO_2 -free atmosphere at a relatively low temperature to obtain pure, anhydrous CaO₂.

Thermal decomposition data for CaO_2 and SrO_2 are summarized in table V. However, these decomposition temperatures are for the pure compounds. When mixed with copper oxide (CuO), the decomposition temperatures are probably considerably lower.

The last aspect of this report concerns the new x-ray diffraction pattern attributed to calcium peroxide octahydrate, $CaO_2 \cdot 8H_2O$. We were unable to verify an exact stoichiometry for the octahydrate because of the ease with which dehydration took place during attempts to remove excess water from the freshly precipitated compound. Because of the isostructural similarity of the unknown $CaO_2 \cdot XH_2O$ with tetragonal octahydrates of other alkaline earth peroxides, $SrO_2 \cdot 8H_2O$ and $BaO_2 \cdot 8H_2O$, we attributed our new x-ray powder pattern (table VI) to $CaO_2 \cdot 8H_2O$. The isostructuralism was confirmed through a lattice parameter program with a diffractometer extrapolation function. Photomicrographs of the $CaO_2 \cdot 8H_2O$ (JCPDS 25-905). This platelet morphology supports the conclusion of preferred orientation occurring and explains the well-defined, integrated-intensity results for the low index (0.0.1) planes. As expected, the lattice parameters of $CaO_2 \cdot 8H_2O$ would be smaller than those for $SrO_2 \cdot 8H_2O$ and $BaO_2 \cdot 8H_2O$ would be smaller than the Sr and Ba atoms.

CONCLUSIONS

The most important practical aspect of our experimental results is the development of a simple synthesis for pure CaO_2 and SrO_2 . The peroxides are of

particular interest as replacements for the corresponding alkaline earth carbonates for the synthesis of the new high-temperature superconductors. Their use in BiCaSrCuO and the most recent TlBaCaCuO systems should offer the same advantages as we found when BaO₂ was substituted for BaCO₃ in the preparation of the YBa₂Cu₃O_{7-x} superconductor. Furthermore, CaO₂ and SrO₂ may be generally useful when an oxidizing alkaline fusion media is required to synthesize compounds other than superconductors.

In view of our experimental results, we recommend the following procedure for the practical synthesis of pure SrO_2 and CaO_2 . For both peroxides, the first step involves precipitation of the octahydrate by dropwise addition of H_2O_2 to a cold solution (about 0 °C) of ammoniacal $Sr(NO_3)_2$ or $CaCl_2$. The best way to prepare anhydrous SrO_2 is to dry the freshly prepared octahydrate for 24 hr at 115 °C, preferably in a CO_2 -free atmosphere. In the case of the less stable CaO_2 , the freshly precipitated octahydrate is best dehydrated at room temperature for at least 24 hr in a CO_2 -free drybox containing an efficient desiccant such as P_2O_5 . The octahydrates of both peroxides are crystalline precipitates that tend to settle out rapidly, thereby allowing most of the mother liquor to be removed by decantation. The crystalline octahydrates are easy to recover by filtration and have little tendency to peptize during filtration and washing. If peroxide yield is important, it can be increased by neutralizing most of the ammonium salt byproduct with a strong base such as NaOH. If sodium contamination is to be avoided, a strong organic base such as tetramethylammonium hydroxide $(CH_3)_4NOH$ may be used instead.

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TABLE I. - CALCULATED STANDARD ENTHALPY OF REACTION (Δ HF°) FOR DECOMPOSITION, DEHYDRATION, AND HYDROL-YSIS OF THE ALKALINE EARTH PEROXIDES

Reaction	Equation	ΔHF° , $kJ-mol^{-1}$		1-1
		Ca0 ₂	Sr0 ₂	BaO ₂
Decomposition Dehydration Hydrolysis	(1) (2) (3)	17.6 65.6 18.0	41.5 103.3 63.6	80.8 85.7 61.1

[From National Bureau of Standards Tables (ref. 3).]

TABLE II. - EFFECT OF VARIOUS TREATMENTS ON Sr02 · XH20

Treatment	$\frac{\text{moles } H_20}{\text{moles } SrO_2}, \\ X$	Composition (from x-ray diffraction analysis)
Air dried, room temperature, 2 days	7.05 to 7.70	$\mathrm{SrO}_2\cdot\mathrm{8H}_2\mathrm{O}$
Dried in dry box over P ₂ O ₅ , room temperature, 5 days	0.29	Major: SrO ₂ Minor: SrO ₂ ·8H ₂ O
Air dried, 100 °C, 24 hr Air dried, 115 °C, 24 hr ^a	0.14 0	SrO ₂ SrO ₂
Air dried, 125 °C, 24 hr	-0.15	Sr0 ²

 $^{\rm a}X-{\rm ray}$ diffraction analysis: ${\rm SrO}_2$ is the only phase. Chemical analysis: 86.94 percent SrO (86.62 percent theoretical). Atomic adsorption analysis: 73 percent Sr (73.25 percent theoretical).

Precipitation temperature, °C	$\frac{\text{moles H}_20}{\text{moles CaO}_2},$	Composition (from x-ray diffraction analysis)	Yield, percent ^a
0 5 10 50	7.4 6.3 5.1 .42	(b) (b) (b) CaO ₂ CaO	67.32 64.23
65 70	.13 .10 .09	CaO_2 CaO_2 CaO_2	54.09 51.03

TABLE III. - SYNTHESIS OF $\text{CaO}_2\cdot\text{XH}_2\text{O}$

^aBased on amount of CaCl·2H₂O (limiting reagent) used. $b_{Attributed}$ to $CaO_2 \cdot 8H_2O$.

TABLE IV. - EFFECT OF HEAT TREATMENT ON MOLES H20/MOLES CaO2 [Starting material: 7.4 mol $H_20/mol \ CaO_2$. Hydrated CaO_2 tends to dehydrate; reacts with ambient CO_2 to form $CaCO_3$.]

Temperature, °C	Duration, hr	<u>moles H20</u> moles CaO2	Composition (from x-ray diffraction analysis)
90	24	0.15	Ca0 ₂
100		.13	Major: CaO ₂
107		.098	Minor: Ca(OH) ₂ Major: CaO ₂ Minor: Ca(OH) ₂
120	\downarrow	.18	Major: CaO_2
Room temperature	48	.11	Minor: CaCO ₃ CaO ₂

TABLE V. - DECOMPOSITION TEMPERATURES TO FORM OXIDES (ref. 5)

$2MO_2 \rightarrow 2MO + O_2$		$2MCO_3 \rightarrow MO + CO_2$		
Compound	Decomposition temperature, °C	Compound	Decomposition temperature, °C	
Ba02	^a 800	BaCO ₃	1450	
Sr0 ₂	b ₄₀₀	SrC0 ₃	1340	
Ca0 ₂	b ₃₃₀	CaC03	899	

^aMelting point, 450 °C. ^bDetermined from our own TGA data.

TABLE VI. - X-RAY POWDER DATA FOR

$CaO_2 \cdot 8H_2O$

[Crystal system: tetragonal lattice parameters (Å); a = $6.212830\pm7.628441\times10^{-3}$; c = $11.00990\pm1.083599\times10^{-2}$; all peaks attributed to CaO₂·8H₂O; no known impurity peaks for CaO₂, CaCO₃, and Ca(OH)₂ were observed.]

2-theta (obs)	d(Å)	hkl	I/I ₀
14.287	6.1993	100	1
16.182	5.4774	002	100
20.295	4.3758	110	1
21.656	4.1037	102	2
26.003	3.4268	112	13
28.759	3.1043	200	1
32.601	2.7467	004	63
33.188	2.6994	211	11
35.724	2.5134	104	7
36.250	2.4782	212	1
38.695	2.3270	114	23
40.765	2.2135	213	11
41.118	2.1953	220	1
44.017	2.0572	204	11
44.407	2.0400	222	2
46.182	1.9657	310	5
46.496	1.9532	214	10
47.003	1.9333	302	1
49.248	1.8503	312	3
52.029	1.7577	106	18
53.111	1.7244	215	5
54.232	1.6914	116	2
55.532	1.6549	304	1
58.495	1.5779	206	1
60.502	1.5290	216	8
60.715	1.5279	216	4
62.120	1.4942	410	1
63.528	1.4645	324	<1
66.032	1.4149	332	<1
67.323	1.3909	420	1
68.215	1.3737	306	6
68.602	1.3669	217	6
68.831	1.3663	217	3
69.467	1.3531	404	2
70.083	1.3416	108	10
70.333	1.3407	108	5
71.328	1.3223	118	<1
73.186	1.2922	334	1
/5./08	1.2563	326	1
77.422	1.2317	218	2
//./50	1.2304	218	1



(a) CaO $_2\cdot$ 8H_2O precipitated from O °C solution.



(b) Enlarged view of CaO $_2\cdot$ 8H $_2$ O precipitated from O °C solution.



(c) Anhydrous CaO_2 precipitated from 70 $^\circ\text{C}$ solution.



(d) Enlarged view of CaO $_{\rm 2}$ precipitated from 70 °C solution.

Figure 1.—Optical photomicrographs of precipitated anhydrous and octahydrate calcium peroxide.

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16. Abstract						
A practical synthesis and a discussion of presented. The general synthesis of these aqueous ammoniacal Sr(NO ₃) ₂ or CaCl ₂ tion techniques. A new x-ray diffraction a = 6.212830 and c = 11.0090 were calcu	some chemical prop peroxides involves p . The octahydrates ar powder pattern for C lated on the basis of	erties of pure stronti precipitation of their e converted to the a $aO_2 \cdot 8H_2O$ is given the tetragonal crysta	um peroxide and calc octahydrates by addi nhydrous peroxides b from which lattice pa al system.	eium peroxide are ition of H ₂ O ₂ to oy various dehydra- arameters		
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