# NATIONAL BUREAU OF STANDARDS REPORT 

## Progress Report

on

## the crystal structure of the aragonite phase of $\mathrm{CaCO}_{3}$

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## U.S. DEPARTMENT OF COMMERCE

NATIONAL BUREAU OF STANDARDS
The Crystal Structure of the Aragonite Phase of $\mathrm{CaCO}_{3}$

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## Abstract

Aragonite $\left(\mathrm{CaCO}_{3}\right)$ crystallizes in the unit cell $a=$ $4.9598(5) \AA, \underline{b}=7.9641(9) \AA$, and $\underline{C}=5.7379(6) \AA$ at $25^{\circ} \mathrm{C}$ with four formula weights in space-group Pmen. The structure has been refined to $\underline{R_{W}}=0.024, \underline{R}=0.040$ using 765 X-ray reflections from a single crystal. The Ca ion is coordinated to nine oxygens with Ca...O distances in the range $2.414(1) \AA$ to 2.653(1) A. The two unique $\mathrm{C}-0$ distances in the $\mathrm{CO}_{3}$ group are $1.288(2) \AA$ (on the mirror plane) and $1.283(1) \AA$. The two unique $0-C-O$ angles are $119.62(4)^{\circ}$ (across the mirror plane) and $120.13(8)^{\circ}$.

## 1. INTRODUCTION

Aragonite $\left(\mathrm{CaCO}_{3}\right)$ is found in nature as a mineral and is an important biomineral because of its presence in coral, clam shells, gallstones and otoliths. It is isomorphous with the carbonates of large divalent cations such as $\mathrm{Ba}, \mathrm{Sr}$ and Pb .

Aragonite is less stable than the calcite phase of $\mathrm{CaCO}_{3}$ at room temperature, but is denser than calcite. This suggests that aragonite is more stable than calcite at low temperatures and/or high pressures. More details are available in reference l. Because of the importance of aragonite, and because of the possibility of performing calculations on the lattice energies of selected carbonates along the lines suggested by Busing[2], we have collected new X-ray data from a single crystal of aragonite and have refined the structure from the positions given in 1924 by Bragg [3].
2. STRUCTURE DETERMINATION

FORMULA (ideal): $\mathrm{CaCO}_{3}$ (aragonite phase); UNIT CELL: orthorhombic with $\underline{a}=4.9598(5) ~ \AA, \underline{b}=7.9641(9) \AA, \underline{c}=$ $5.7379(6) \AA$ at $25^{\circ} \mathrm{C}$ (calculated by least squares from $122 \theta$ values observed on a diffractometer); volume: $226.65 \AA^{3}$; radiation, $\mathrm{Mo}\left(\mathrm{K}_{\mathrm{Q1}}\right)$, $\lambda=0.70926 \AA$; monochromator: highly
oriented graphite crystal; space-group: Pmen; contents $4\left(\mathrm{CaCO}_{3}\right)$; reciprocal lattice extinctions, hkO: $h+k=2 n+1$, hol: $\ell=2 n+1$; observed density, $2.947(2) \mathrm{g} \cdot \mathrm{cm}^{-3}$ [4]; calculated density, $2.944 \mathrm{~g}^{\cdot \mathrm{cm}^{-3} \text {; CRYSTAL: material available }}$ heavily twinned; small wedge largest single crystal fragment found; this wedge was attached to thin borate glass fiber with clear household cement; fiber attached to insert in goniometer head with epoxy cement; origin of crystal, mineral sample \#75538 from National Museum of Natural History, Smithsonian Institution, Washington, D. C. Supplied by J. S. White, Jr.; linear absorption corrections made by 8 x 8 x 8 Gaussion quadrature using subroutines written by C. W. Burnham [5] and adapted by B. Dickens; maximum and minimum corrections for absorption $=0.963$ and 0.880 (transmission factors). INTENSITY DATA: number of reflections, 2356 collected from 2 octants and merged into a unique set of 765 , of which 619 are "observed" and 146 are "unobserved"; unobserved reflections are those less than $2 \sigma$ above background; maximum $\sin \theta / \lambda$ for data $0.907 \AA^{-1}$; method used to estimate data\& $\theta / 2 \theta \operatorname{scan}$, scintillation counter; diffractometer: Picker 4-circle single-crystal diffractometer automated by PDP 8/I computer through FACS-l interface and adapted to include least significant digit of counts; COMPUTATION: setting programs,
those of reference [6] as adapted by Picker Nuclear Corporation; scan range: $1.4^{\circ}+114.6 \frac{\Delta \lambda}{\lambda}, \Delta \lambda=0.692$, $\lambda=0.70926 \AA$; scan parameters: backgrounds counted at higher and lower $2 \theta$ for 100 sec . each; $\theta / 2 \theta \operatorname{scan}$ at $0.25^{\circ} / \mathrm{min}$ for $2 \theta$ from one background position to the other; attenuators: 0.025 mm thick layers of $\mathrm{Nb}, 1$ layer for first attenuator, 2 for second, 3 for third; scan range correction: table lookup method to obtain values recommended in reference [7]; paper tape processing program written by B. Dickens for Univac 1108 computer, contains adaptions of similar program by F. A. Mauer (NBS), standard reflection plotting routine and extince reflection editing routine from programs by J. M. this program Stewart, University of Maryland; Muses an intense standard reflection at low angle to correct for change in intensity of the primary X-ray beam. Counts in peak $=I=P-\left(T / 2 T_{B}\right)$ $\left(B_{L}+B_{H}\right), \sigma(I)=\left(P+\left(B_{L}+B_{H}\right)\left(T /\left(2 T_{B}\right)\right)^{2}\right)^{\frac{1}{2}}, F=((A F)(L P)$ $(I))^{\frac{1}{2}}, \sigma(F)=(\sigma((I) / 2)(L P / I))^{\frac{1}{2}}$ where $P=$ counts at the peak position, $B_{L}$ and $B_{H}=$ background counts at lower and higher $2 \theta$ respectively, $T=$ time spent counting peak, $T_{B}=$ time spent counting background, $A F=$ attenuator factor, $L P=$ Lorentz-polarization correction. Data merging program for equivalent reflections, written by B. Dickens for Univac in this program is 1108 computer; /each set of equivalent reflections/treated as
follows: Reflections which were all unobserved were averaged and given the largest individual standard deviation in the set. Unobserved reflections in the presence of at least one observed reflection were discarded. Observed reflections which, subsequent to this step, occurred only once in the list were copied unchanged but their standard deviations were increased by a factor of three. Observed reflections with magnitudes which agreed within the counting statistics and reflections with magnitudes whose ratios fell within the range 0.95 to 1.06 were averaged and given as standard deviation the maximum of the standard deviation from counting statistics and the standard deviation from the range estimate [8,9]. Under these circumstances, reflections whose magnitudes did not pass the criteria were discarded. If no reflections passed the criteria, the highest magnitude was taken and the associated standard deviation multiplied by five. The justification for these arbitrary increases of standard deviations is that without some corroboration, every reflection is suspect because of the possibilities of multiple reflection, including the "tail" of nearby intense reflectionsin its measurement, change in intensity of $X$ of crystal
ray beam, misalignment/etc. Scattering factors; those for
the neutral atoms in reference 10 , least-squares refinements: full-matrix, with $\Sigma\left(w\left\|F_{0}|-| F_{C}\right\|\right)^{2}$ minimized; refinements include unobserved reflections which calculate higher than $2 \sigma$ above background; least-squares weights: $1 / \sigma^{a}$; $\left.\underline{R}_{\underline{W}}=\left(\Sigma\left(w\left\|F_{0}|-| F_{C}\right\|\right)\right)^{2} / \Sigma\left(w\left|F_{0}\right|\right)^{z}\right)^{\frac{1}{2}}, \underline{R}=\Sigma\left\|F_{0}\left|-\left|F_{C} \| / \Sigma\right| F_{0}\right| ;\right.$ thermal parameters have the form $\exp \left(-1 / 4\left(\underline{a} *^{2} \underline{\underline{B}}_{11} \underline{h}^{2}+\right.\right.$ $\left.\underline{B}_{22} \underline{k}^{2}+\underline{C}^{* 2} \underline{B}_{33} \underline{l}^{2}+2 \underline{a}^{*} \underline{b}^{*} \underline{B}_{12} \underline{h k}+2 \underline{a}^{*} \underline{C}^{*} \underline{B}_{13} \underline{h} \underline{\ell}+2 \underline{b} \underline{c}^{*} \underline{B}_{2} \underline{B}_{2} \underline{k}\right)$. Least squares and electron density synthesis calculations carried out with X-ray 67 system [11] of computing programs. FINAL REFINEMENT: $R_{W}=0.024 ; R=0.040$, average shift/error for last cycle $=0.0017$; standard deviation of an observation of unit weight $=\left(\Sigma\left(w\left(F_{O}-F_{C}\right)\right)^{2} /(765-28)\right)^{\frac{1}{2}}=0.775$.

The highest peak in an electron density difference synthesis calculated after the final anisotropic refinement to $\underline{R}_{\underline{w}}=0.024$ corresponded to about $1 / 3$ of an electron and was about $0.95 \AA$ from $C$ towards $O(1)$. The largest correlation coefficients are $\quad 0.34-0.44$ between the scale factor and the $\underline{B}_{11}$. $\underline{B}_{29}$ and $\underline{B}_{33}$ temperature factors of Ca .

The atomic parameters are given in table l. The observed and calculated structure factors are given in table 2.
3. DESCRIPTION OF THE STRUCTURE

The structure of aragonite, the main points of which are well known, is shown in figure 1. The Ca ions lie in pseudohexagonal layers parallel to (001) and the layer sequence is
$A B A B$. The Ca layers are separated by $\mathrm{CO}_{3}$ groups which lie in two layers parallel to (001), and form columns parallel to [001].

### 3.1. The Ca ion environments

The Ca ion lies on the mirror plane at $x=1 / 4$. Its environment is shown in figure 2 and summarized in table 3. The coordination contains three $\mathrm{CO}_{3}$ edges, $0(1,2), 0\left(I^{I}, 2^{I}\right)$ and $O\left(2^{I V}, 2^{V}\right)$ and three apexes, $O\left(I^{I I}, 2^{I I}, 2^{I I I}\right)$. The apparent thermal motion of Ca is almost isotropic, (table 1, Fig. 2).

$$
\text { 3.2. The } \mathrm{CO}_{3} \text { group }
$$

The dimensions of the $\mathrm{CO}_{3}$ group are given in table 4 .
All $\mathrm{C} \rightarrow \mathrm{O}$ distances in the $\mathrm{CO}_{3}$ group are essentially equal with an average of $1.286 \AA$. This agrees well with the C-O distance of $1.283(2)$ reported [12] for calcite. There appears to be some significant deviation from trigonality in the angles; the reported angle of $119.62^{\circ}$ for $0(2)-C-0\left(2^{I}\right)$ is consistent with the $0\left(2,2^{I}\right)$ edge of the $\mathrm{CO}_{3}$ group being coordinated slightly more strongly to Ca as judged from the Ca...O distances. If the apparent thermal motions of the atoms in the $\mathrm{CO}_{3}$ group are attributed to thermal motion rather than to slight positional disorder, there seems to be oscillation within the $\mathrm{CO}_{3}$ layer, i.e.,
more or less perpendicular to the edge coordination to Ca . Similarly, $0(1)$ may be undergoing some additional wagging out of the $\mathrm{CO}_{3}$ layer.

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Table 3

The Ca environment in aragonite $\left(\mathrm{CaCO}_{3}\right)$

| Atoms | Distance, $\AA$ |
| :--- | :--- |
| Ca, $0\left(I^{I I}\right)$ | $2.414(1)$ |
| Ca, $0\left(2^{I I}, 2^{I I I}\right)$ | $2.445(1)$ |
| Ca, $0\left(2,2^{I}\right)$ | $2.520(1)$ |
| Ca, $0\left(2^{I V}, 2^{\mathrm{V}}\right)$ | $2.544(1)$ |
| Ca, $0\left(1,1^{I}\right)$ | $2.653(1)$ |

In all tables of interatomic distances and angles, the figures in parentheses are standard deviations in the last digit and were calculated from the standard deviations in the atomic positional parametersand the unit cell parameters.

## Table 4

The $\mathrm{CO}_{3}$ group and its environment in aragonite $\left(\mathrm{CaCO}_{3}\right)$

Atoms
C. $0(1)$

C, $0(2)$
$0(1), O(2)$
$0(2), O\left(2^{I}\right)$
$0(1), C, O(2)$
$0(2), C, O(2 I)$
$0(1), \mathrm{Ca}^{\mathrm{V}}$
$0(1),\left(\mathrm{Ca}^{I I}, \mathrm{Ca}^{\text {III }}\right)$

O(2), Ca
$\begin{array}{ll}0(2), & \mathrm{Ca} I I \\ 0(2), & \mathrm{Ca} I V\end{array}$

Distance, $\AA$, or angle, deg.

$$
\begin{aligned}
& 1.288(2) \AA \\
& 1.283(1)
\end{aligned}
$$

$$
2.229(1)
$$

$$
2.219(1)
$$

$$
120.13(8) \circ
$$

$$
119.62(4)
$$

$2.414(1) \AA$
2.653(1)
2.445(1)
2.520(1)
2. 544 (1)

## Figure legends

1. A stereoscopic illustration of the crystal structure of aragonite $\left(\mathrm{CaCO}_{3}\right)$. A unique set of atoms is labelled. The origin of the crystallographic coordinate system is marked by *.
2. The Ca environment in aragonite $\left(\mathrm{CaCO}_{3}\right)$. The labels refer to atoms in table 3.
3. The $\mathrm{CO}_{3}$ group environment in aragonite $\left(\mathrm{CaCO}_{3}\right)$. The labels refer to atoms in table 4.


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