Communication

Single-Crystalline Fibers of Deuterated Potassium Dihydrogen Phosphate

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Abstract: Single-crystalline fibers have distinct structures and optical properties comparing with the bulk crystals. In this article, two types of single-crystalline fibers of deuterated potassium dihydrogen phosphate (K(\(\text{H}_{1-x}\text{D}_x\))\(_2\)PO\(_4\), DKDP) are obtained by rapid growth in room-temperature supersaturated solution. X-ray diffraction analysis reveals that these DKDP single-crystalline fibers belong to tetragonal (I-42d) and monoclinic (P2\(_1\)/c) phases, respectively. The crystal structure of the tetragonal DKDP single-crystalline fiber is identical to that of the bulk DKDP tetragonal crystal reported. The lattice parameters of the monoclinic DKDP fiber (with the deuterium content of 55\%) are a = 14.6571 Å, b = 4.5187 Å, c = 18.6962 Å, and \(\beta = 108.030^\circ\), which is a new crystal phase of DKDP. The monoclinic DKDP single-crystalline fiber is metastable at the present experimental condition and readily transit to the corresponding DKDP tetragonal phase in solution and in solid by grinding. The optical experiment shows that the highly deuterated tetragonal DKDP single-crystalline fiber possesses excellent optical guided-wave and effective second-harmonic generation properties. DKDP single-crystalline fibers are expected to be the suitable candidates for fabrication of the miniaturized nonlinear optical devices.

Keywords: single-crystalline fiber; deuterated potassium dihydrogen phosphate; crystal phase; phase transition; second harmonic generation

1. Introduction

Potassium dihydrogen phosphate (KH\(_2\)PO\(_4\), KDP) and its isotope compound, deuterated potassium dihydrogen phosphate (K(\(\text{H}_{1-x}\text{D}_x\))\(_2\)PO\(_4\), DKDP), are irreplaceable electro-optical and nonlinear optical materials in the inertial confinement fusion (ICF) system because of their outstanding capability of growing into large size crystals with high optical quality as well as large nonlinear and electro-optic coefficients [1]. For the past few decades, therefore, the crystal growth and fabrication techniques are the main research fields for KDP-type crystals in order to satisfy the urgent demand of ICF for these two crystals [2].

Recently, we reported the crystal growth of one-dimensional KDP microstructures with a large length-to-diameter ratio grown under the ambient conditions. They exhibit unique crystal growth mechanism, which is quite different from that of the bulk KDP crystals. Moreover, a highly efficient second-harmonic generation was observed in these KDP single-crystalline fibers [3]. The discovery of these one-dimensional KDP materials opens a door to the unknown world of low-dimensional micro-/nano- KDP crystals. It indicates that except for the traditional bulk KDP/DKDP crystals, low-dimensions KDP-type crystals may become another group of robust nonlinear materials, especially for flexible and miniaturized device application in optical communications, high-energy laser, and so on.
Crystal growth is the first problem before considering their device applications. The crystallization of specific crystals is mainly controlled by the growth condition and molecular structure. For the isotopic crystals, on one hand, they generally have similar crystallization habits as well as phase species or phase structures under the same growth conditions. For example, DKDP crystallizes in the tetragonal crystal phase (I-42d) [4], which is isomorphic to the KDP tetragonal phase formed at ambient temperature. On the other hand, the isotope deuterium plays a role in crystallization habit and formation of the crystal structure. The growth process of DKDP tetragonal phase usually accompanies the crystallization of the DKDP monoclinic phase (P2₁) [5,6]. However, P2₁ phase of DKDP has no isomorphic KDP crystal phase crystallized in the aqueous solution. The structure of DKDP P2₁ phase is similar to a monoclinic phase of KDP crystal formed at a relatively high temperature [7]. In this article, we study the crystallization habits of DKDP during the fast growth in high supersaturated aqueous solution at room temperature. Two types of single-crystalline fibers of DKDP with tetragonal (I-42D) and monoclinic (P2₁/c) phases are obtained. The phase transition property between these two phases and the nonlinear frequency conversion property of the tetragonal DKDP single-crystalline fibers are also studied.

2. Materials and Methods

Hot solution was prepared using high purity KDP, D₂O (99.9 atom% deuterium, Aladdin, Shanghai, China) and ultrapure water (Millipore, America). All glasswares were washed before use in hydrochloric acid and then thoroughly washed with ultrapure water in an ultrasonic cleaner in order to remove the insoluble particles and possible metal ions (see Supplementary Materials) [8–10]. The hot DKDP solution is dropped on a cleaned glass slide, and evaporatively cooled at room temperature in a vacuum drier (Shuniu, Sichuan, China), which rapidly crystallizes and creates fiber-like DKDP single-crystals. The crystals are kept in vacuum dryer waiting for use. The final humidity of the vacuum drier is less than 10%.

3. Results and Discussion

SEM (Scanning electron microscope) shows that the fibers obtained exhibit two types of shapes with polyhedral or tetragonal cross-section, respectively (see Figure 1). Moreover, the solid-core/tubular structures can be found in both types of fibers. The length-to-diameter ratio for both types of fiber is up to 500:1. The smooth prismatic surfaces of these fibers indicate an excellent crystal quality. Obviously, the shape of these single-crystalline fibers is very different to that of the bulk tetragonal or monoclinic phases of DKDP crystals previously reported [2,11].

The X-ray diffraction analysis is performed in order to determine the crystal structure of these single-crystalline fibers. The fiber crystals are picked from the sample freshly prepared, all of which are protected by the Paratone oil in order to keep away from the moisture during the measurement. The result shows that the DKDP single-crystalline fiber with the polygonal shape exhibits a 2/m point group symmetry with P2₁/c space group, which is different to the structure of the bulk monoclinic DKDP crystal reported (P2₁). The summarized crystallographic data of the monoclinic crystal phases of DKDP and KDP are shown in Table 1 for comparison. It shows that the unit cell parameters of the DKDP monoclinic single-crystalline fiber are a = 14.6571(7) Å, b = 4.5187(2) Å, c = 18.6962(9) Å and β = 108.030(2)°. The XRD spectra prove that the crystal structures of the polygonal shape DKDP fiber grown in different deuterium-content DKDP solution is the same monoclinic structure as shown in Table 1 (see Supplementary Materials). To our knowledge, the DKDP monoclinic P2₁/c is a new crystal phase of DKDP. According to the data in Table 1 and the packing diagram shown in Figure 2, the structure of the DKDP P2₁/c phase is identical to that of the monoclinic KDP single-crystalline fiber (P2₁/c) obtained under the same condition [3]. Therefore, the monoclinic phase DKDP single-crystalline fiber is isomorphic to the corresponding KDP monoclinic single-crystalline fiber crystal. The tetragonal-shaped single-crystalline fibers possess the identical space group and crystal lattice parameters to the bulk tetragonal DKDP phase (I-42d). Moreover, it is isomorphic to the
KDP tetragonal phase [12]. Except for the above two crystal phases, no DKDP monoclinic phase (P2₁) was detected in the obtained DKDP crystals grown under the present experimental conditions [5].

Figure 1. SEM images of the DKDP single-crystalline fibers obtained by rapid crystallization on glass substrate. (a,b) polyhedral solid-core/tubular single-crystalline fibers. (c,d) tetragonal solid-core/tubular single-crystalline fibers.

Table 1. Basic crystallographic data comparison of the monoclinic DKDP bulk crystal, monoclinic DKDP, and KDP single-crystalline fibers.

| Formula             | K(H₀.₁₆D₀.₈₄)₂PO₄ | KH₂PO₄  | K(HₓD₁₋ₓ)₂PO₄ |
|---------------------|------------------|----------|----------------|
| Space group         | P₂₁              | P₂₁/c    | P₂₁/c          |
| a (Å)               | 7.1400           | 14.599(5)| 14.6571(7)     |
| b (Å)               | 14.7100          | 4.503(5) | 4.5187(2)      |
| c (Å)               | 7.4500           | 18.651(5)| 18.6962(9)     |
| α (°)               | 90               | 90       | 90             |
| β (°)               | 92.310           | 108.041(5)| 108.030(2)    |
| γ (°)               | 90               | 90       | 90             |
| V (Å³)              | 781.83           | 1165.8(14)| 1177.46(10)   |
| Z                   | 8                | 12       | 12             |
| T (K)               | 293              | 293      | 297            |
| CSD No.             | 34415            | 427178   | 1963699        |
| reference           | [5]              | [3]      | This work      |

Raman spectra of the monoclinic and tetragonal single-crystalline fibers are measured in order to estimate their deuterium contents. The result is shown in Figure 3. The Raman spectrum of DKDP crystal typically possesses double peaks within 800 cm⁻¹ to 1000 cm⁻¹. The Raman peaks locate at 891.4 cm⁻¹ and 964.5 cm⁻¹ for the monoclinic phase single-crystalline fiber obtained in 75% deuterium content solution, while the peaks shift to 885.5 cm⁻¹ and 987.6 cm⁻¹ for the tetragonal single-crystalline fiber grown in solution containing 85% deuterium. The deuterium content for crystals is roughly estimated based on the peak maximum of 891.4 cm⁻¹ and 888.5 cm⁻¹ for monoclinic and tetragonal phases, respectively, based on the empirical formula [13]. The results are 55% for the monoclinic and 72% for the tetragonal single-crystalline fibers. In addition, the Raman scattering spectra are measured for multiple single-crystal fibers randomly chosen to show the deuterium-content fluctuation. The results are almost the same for these fiber samples. The statistical results for the deuterium-content estimation are shown Supplementary Materials.
Phase transition is one of the features of KDP-type crystals because of their hydrogen bond ferroelectric properties [14]. It is known that DKDP is characterized by the phase transition of the metastable tetragonal phase to the stable monoclinic phase ($P2_1$) [6]. However, according to the above description, there is no crystal phase of $P2_1$ detected both in solution and in solid state crystals under the present condition. The main reason for an absence of bulk monoclinic phase being detected should be that the solution with the deuterium content of 75–85% is located at the stability region of the tetragonal phase. The phase equilibrium transition point for this deuterium content solution is higher than 60 °C [6]. Since our solution crystallizes typically at a temperature lower than 60 °C, the bulk DKDP monoclinic phase is hardly detected in the solution. In our experiment, however, the monoclinic phase ($P2_1/c$) DKDP fiber appears in the growth solution and the final products. We found that the monoclinic DKDP single-crystalline fiber readily transits to the tetragonal phase if there is tetragonal crystal nucleus or particles in the solution. Figure 4 shows the evolution process from the crystallization to dissolution of the monoclinic DKDP single-crystalline fibers. The monoclinic fibers begin to grow in Figure 4a. Then the tetragonal particles appear in Figure 4b, which grow rapidly as shown in Figure 4c. One can also see the gradual dissolution of the DKDP monoclinic single-crystalline fibers in

Figure 2. Packing diagram of monoclinic ($P2_1/c$) DKDP crystal view along b axis. Grey: potassium, blue: phosphate, green: oxygen, yellow: hydrogen/deuterium.

Figure 3. Raman spectra of the monoclinic phase single-crystalline fiber and tetragonal phase single-crystalline fiber. The double peaks indicate the existence of deuterium in the crystals.
Phase transition is one of the features of KDP-type crystals because of their hydrogen bond. Deuterated KDP crystals have the advantage over KDP in decreasing the damage possibility when a non-centrosymmetric single-crystal (see the optical setup description in Supporting Information). The purpose of studying the DKDP low-dimensional crystals is to develop new functional materials with distinct optical properties. For single-crystalline fibers, they are characterized by the excellent nonlinearity, extended interaction length with laser light, tight spatial confinement of light in the waveguide structure, and natural physical flexibility, which benefit to high nonlinear conversion efficiency with much lower laser threshold [15–18]. Here, we further measure the second-harmonic generation (SHG) property of the tetragonal DKDP single-crystalline fiber since it is a non-centrosymmetric single-crystal (see the optical setup description in Supporting Information). Deuterated KDP crystals have the advantage over KDP in decreasing the damage possibility when they are used in type-II third-harmonic generation devices, which benefit from the split and reduced damage.
stimulated Raman scattering peaks as pumped by a high power laser [19]. In our experiment, the tetragonal single-crystalline fiber exhibits the excellent guided-wave for 532 nm laser (see Figure S4). No light leakage occurs during the laser propagating along the c-axial direction, which further indicates a high optical quality of the DKDP single-crystalline fiber. The second harmonic generation is achieved when a continuous wave 1064 nm laser being coupled into the entrance of the fiber along the c-axis. The spectrum is collected by a PMT grating spectrometer.

![Graph](image-url)

**Figure 5.** Comparison of XRD spectra of the DKDP monoclinic single-crystalline fibers before and after grinding. (a) the DKDP single-crystalline fibers on the substrate before grinding. (b) the DKDP single-crystalline fibers on the substrate after grinding. (c) the XRD spectra of tetragonal I-42d DKDP crystal phase. (d) the XRD spectrum of the monoclinic P2_1/c DKDP crystal phase.

The peak of the SHG spectrum in Figure 6a locates at 532.36 nm with the FWHM (full wide of half maximum) of ~0.18 nm. At the same time, a bright green light signal is captured by the CCD camera (Qimage MicroPublisher™3.3 RTV, Olympus, Japan) opposite to the exit of the fiber (Figure 6a inset). The curve of the SHG output intensity versus input power \( P(\omega) \) is shown in Figure 6b, which is well fitted by the function of \( P(2\omega) = C_2^2P^2(\omega) \). The fitting result shows a coherent propagation growth of the SHG along the c-axis of the fiber. The SHG efficiency calculated is about \( 5 \times 10^{-3} \) W\(^{-1}\). This frequency conversion efficiency is nearly the same level of that of the tetragonal KDP single-crystalline fiber we reported before [20]. The efficiency difference between them is not caused by the substitution deuterium for hydrogen in DKDP crystal. In fact, the diameter, length as well as the surface polishing play the key roles in the SHG efficiency improvement [21]. Here, the SHG of the tetragonal DKDP single-crystalline fiber should be an effective frequency conversion considering only about 2% of the pump light effectively has the right k vector for efficient phase matching inside the fiber because of the spread of the k.
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**Supplementary Materials:** The following are available online at http://www.mdpi.com/xxx/s1, Figure S1: PXRD spectra of the monoclinic DKDP single-crystalline fibers grown in different deuterium-content solution, Figure S2: The deuterium content analysis of the DKDP monoclinic single-crystalline fibers based on Raman spectra, Figure S3: The deuterium content analysis of the DKDP monoclinic single-crystalline fibers based on Raman spectra, Figure S4: The optical guided-wave of the DKDP tetragonal single-crystalline fiber for 532 nm laser.

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**Conflicts of Interest:** There are no conflicts to declare.

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