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1. Introduction

Silicon carbide (SiC) is a representative wide-bandgap semiconductor material.¹ For conductive SiC, epitaxial layers are grown on a substrate and applied to power devices, which are widely used in electric vehicles, charging piles, photovoltaics, rail transit, smart grids, and so on. Due to its characteristics of low consumption, high efficiency and high power, SiC is expected to revolutionize the power electronics industry. With the attention on carbon neutrality, the application of SiC in the field of new energy has attracted the attention of researchers.²

The high manufacturing cost of SiC material limits its popularization and application.^{3,4} Its preparation is faced with the dilemma of low monocrystalline yield, especially when the size reaches 6 inches and above. For large crystal preparations, polycrystals, polytypes, and cracks are easily appear, and micro defects such as crystal micropipes and dislocations are also produced rapidly,^{5,6} which greatly restrict the performance of power devices.⁴ During the monocrystalline growth process, the temperature distribution of thermal fields plays a decisive role,^{7,8} and optimizing thermal conditions in the growth cavity is an effective way to improve crystal quality.⁹ However, due to the inability to directly observe the process parameters at high temperatures, we use numerical simulation methods to simulate the physical field, producing reliable results quickly and greatly reducing the experimental cost.¹⁰ In this paper, the STR

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Optimization of thermal field of 150 mm SiC crystal growth by PVT method

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The excellent physical properties of SiC as an electronic material determine its important application prospects, especially in the new-energy industry, but the preparation of large-sized materials with high quality is not easy. Therefore, the physical fields in the growth process were modeled and studied with the help of the numerical simulation software Virtual Reactor, and its accuracy was verified by the agreement between morphology of the experimental crystal and the simulation. Additionally, the effects of thermal insulation adjustment of crystal growth thermal fields, application of seed crystals with different diameters, and shelter structure on the crystal growth process were also studied. By optimizing the crystal growth conditions, a nearly flat and slightly convex crystal growth interface was obtained successfully in our lab. Crystal quality was significantly improved, and a 6-inch SiC crystal with single polytype, high quality and low defects was successfully prepared.

software Virtual Reactor (VR) was used to study the process of crystal growth.

In this work, a physical model of induction heating 6-inch crystal growth was established, and distribution characteristics of the main parameters based on growth time were obtained. Compared with the experimental parameters, the growth interface was optimized in the process of crystal growth, and regulation methods for interface shape were also revealed. Finally, experimental results show that high-quality single crystals with low defects can be obtained under optimized conditions.

Modeling and experiment

Physical vapor transport (PVT) method is the main process used to manufacture SiC crystals,¹¹ which are divided into induction heating type and resistance heating type. In this work, we used induction heating, and the basic design of the thermal field's physical structure is shown in Fig. 1a. The crucible is in the center position, surrounded by insulation modules, and it contains the seed holder, seed, growth cell, and SiC sources from top to bottom. For insulation module material, MFA (rigid felt) and GFA (soft felt) from SGL Group were respectively used in the bottom module and in other places. The crucible, made of SGL graphite R6510, also functions as a heater, and its physical parameters are shown in Table 1. The basic principle of the PVT method is that the SiC source in a high-temperature zone sublimates and then is transported to the surface of the seed as a condensation zone, where it becomes saturated vapor, nucleates, and becomes a crystal.¹²⁻¹⁴ To measure temperatures inside the crucible, two viewports on the upper and lower sides, respectively, were prepared.



Fig. 1 (a) Structure of hot zone and (b) mesh of model.

 Table 1
 Properties of crucible materials

Parameter	Value
Bulk density/g cm ^{-3} Resistivity/ μ Ω m Dynamic modulus of elasticity/MPa Thermal expansion (20–200 °C)/(K ^{-1}) Thermal conductivity (20 °C)/(W m ^{-1} K ^{-1})	$1.83 \\ 13 \\ 11.5 \times 10^3 \\ 4.2 \times 10^{-6} \\ 105$

SiC crystals grow slowly under high temperature at about 2500 K with suitable temperature gradient and a low vapor pressure of 100–4000 Pa, and usually, 5–10 days are needed to obtain a 15–30 mm thick crystal. In this simulation, we used 700 μ m production grade (micropipe density no more than 0.5/ cm⁻²), 6-inch 4H type polished wafer as seed material with 4° off-orientation angle. The temperature control point was placed at the center of the seed holder's upper surface and set to 2523.15 K. Vapor pressure in the growth cell was set to 2000 Pa with 90% argon and 10% nitrogen. Frequency of induction heating was set to 10 kHz, and heating power was calculated according to the temperature and thermal properties of the entire structure. To get crystal profiles at different growth time was set to 100 hours in the initial model.

There are more than 200 kinds of homogeneous isomers of SiC single crystal with little difference in physical properties, which makes it difficult to prepare single-structure crystals.¹⁷ 4H–SiC and 6H–SiC crystals are two different polytypes that can grow stably at present, and Table 2 shows their physical properties at 300 K. 4H–SiC has a wider band gap and better thermal conductivity than 6H–SiC, showing its advantages in manufacturing power devices.

Table 2 Properties of main polytypes ^{18,19}				
Parameter		4H–SiC Wurtzite type structure	6H–SiC Cubic β-ZnS structure	
Crystal structure				
Lattice parameters/Å	a	3.073	3.073	
	b	10.053	15.118	
Band gap/eV		3.230	3.000	
Melting point/°C		2830	2830	
Thermal conductivity/(W m ^{-1} K ^{-1})		480	360	
Electron mobility/($cm^2 V^{-1} s^{-1}$)		900	400	
Hole mobility/($cm^2 V^{-1} s^{-1}$)		120	90	
Breakdown field strength V cm^{-1}		$2 imes 10^6$	2.1×10^6	
Infrared refractive index		$2.55 (\perp c)$	$2.55 (\perp c)$	
		2.59 (c)	2.59 (c)	



Fig. 2 (a) Crystal shape predicted at different growth times, (b) actual crystal.



Fig. 3 Crystal growth rates in different positions at different growth times.

3. Results and discussion

3.1 Physics field distribution in the crucible

3.1.1 Morphology of SiC crystal. During SiC crystal growth, flat surface shape with a little convex is crucial to avoid polytype formation and decrease internal stress, which is fatal for crystal quality. After iterative computations, profiles of SiC crystal under different growth times were obtained, as shown in Fig. 2 together with the actual crystal for comparison. The crystal growth interface shows an obvious "M" shape with polycrystal surrounding the monocrystal part (Fig. 2a). As the SiC crystal grows thicker, the "M" shape becomes more obvious, and the polycrystal (area D) grows even faster, which compresses the space of the monocrystal. The diameter of the center monocrystal portion is less than 130 mm (around area C) after 100 hours, and its maximum thickness (area A) is 20 mm with 200 μ m h⁻¹ growth rate, while the thinnest part (area B), with 9 mm thickness, is 110 mm in diameter. Unfortunately, greater convexity always implies greater internal stress.²⁰ Meanwhile, we obtained 150 mm crystal in the lab with a convex and smooth surface within 130 mm in diameter, but polycrystal in the surrounding 130-140 mm area after the rounding treatment (Fig. 2b), which shows the consistency between experiment and simulation in shapes and profiles. This result suggests disadvantages of this hot zone in both interface shape and polycrystal propagation.

Growth rates in different stages also illustrate this conclusion, as shown in Fig. 3. The difference along the seed is large since the initial stage. The parameter is about $320 \,\mu\text{m}\,\text{h}^{-1}$ at the beginning in area A and decreases rapidly to below $130 \,\mu\text{m}\,\text{h}^{-1}$ in the range greater than 70 diameters, while it shows an abnormal behavior in area C between 130 mm to 150 mm diameters. As the crystal grows, growth rate in the center of the seed decreases to $190 \,\mu\text{m}\,\text{h}^{-1}$, $160 \,\mu\text{m}\,\text{h}^{-1}$ and $100 \,\mu\text{m}\,\text{h}^{-1}$ after



Fig. 4 Temperature distribution in the top of the crucible at initial stage.



Fig. 5 Temperature distribution in the growth cell at different growth times.

20 h, 50 h and 100 h. Meanwhile, the growth rate along the radial direction becomes relatively smooth, with fluctuations evident especially in the crystal edge, which suggests that the growth conditions are not unstable, and will be discussed later.

3.1.2 Temperature distribution in growth cell. Temperature has a direct influence on crystal growth, since temperature is the driving force for gas phase species transfer and deposition on crystal seed. For a general view, the initial temperature distribution of the hot zone's top part was obtained and is shown in Fig. 4. In the growth cell, the lowest value appears in center of the seed under the seed holder due to the existence of the viewport, which provides supercooling and shows a great influence on the crystal growth area in the growth cell.

The growth cell is the chamber where gas species flow and deposition happens. The temperature distribution here at different growth stages was also obtained, as shown in Fig. 5. The highest temperature appears in SiC powder near the crucible wall, and the lowest temperature appears in the center of the SiC seed, ranging from 2560 K to 2680 K. Following the white isotherms at different times, we can find that the hightemperature zone becomes closer to the crystal as the crystal grows, which is easy to understand, as both crystal and source play important roles. When the crystal gets thicker, the thermal resistance of powder source in different parts changes simultaneously. The center of the source surface starts recrystallization, and the part close to the crucible wall undergoes graphitization. Meanwhile, the process is affected by a certain crystal thickness because the growing crystal changes heat transfer, resulting in slight "M"-shaped isotherms, just like the crystal. In other words, the temperature becomes higher, then lower, and higher again from the center to the side in a horizontal direction, which increases temperature gradients in area A and decreases temperature gradients in area B.

To obtain the details of temperature distribution, Fig. 6 was generated by postprocessing to show temperature gradients along the axial and radial directions. Red and blue, respectively, suggest positive and negative parameters. From powder to crystal surface, the temperature gradient along the axial



Fig. 6 Temperature gradient distribution in the growth cell at different growth times: (a) along radial direction, (b) along axial direction.



Fig. 7 (a) Temperature distribution with velocity vectors, (b) temperature distribution with velocity streamlines, (c) gas velocity distribution at 20 h, (d) gas velocity distribution at 50 h.

direction is about 0 K cm⁻¹ to -15 K cm⁻¹ around the crystal, and -30 K cm⁻¹ to -45 K cm⁻¹ inside the crystal (Fig. 6b). The top part without SiC seed (area D in Fig. 2) has a similar temperature gradient of about -25 K cm⁻¹, which provided a greater driving force for crystal growth with seed or polycrystal without seed. Temperature gradients along the radial direction can also explain the poor crystal interface shape (Fig. 6a). The gradient increased to above 4 K cm⁻¹ from the center part to the side of the seed, creating a 100 mm to 120 mm diameter seed (area B) with the highest temperature and lowest temperature gradients along the axial direction. As the crystal grows, the side of the seed or the junction point of crystal, polycrystal and gas phase (area C) change violently in temperature distribution, which exacerbates the crystal edge growth state and causes small-diameter crystals with small effective thickness.

3.1.3 Flow field and velocity distribution. Guo²¹ put forward the field-coordination theory and pointed out that reducing the angle between velocity vector and temperature gradient is beneficial to enhance convective heat transfer and also mass transfer. Fig. 7a shows gas phase flow velocity distribution in the growth cell, with arrows in different lengths indicating the value, and Fig. 7c and d show velocity in axial direction. Since the crucible functions as the heater and the powder source locates in the high-temperature zone, gas phases such as Si, Si₂C and SiC₂ formed mostly in the powder side and flow to seed with low temperature. As a result, red color appears around the junction point (area C), and dark blue near the crucible wall close to the powder source surface. Meanwhile, velocity streamlines around the center of seed are perpendicular

to the growth surface (Fig. 7b). What is similar to Fig. 6 is that the areas around the junction point (area C) and center of crystal (area A) have greater parameters before 50 hours, and the former velocity even reaches above 0.03 m s⁻¹. In contrast, velocity is no more than 0.02 m s⁻¹ in other areas of seed surface. This suggests that the temperature gradient along axial direction deeply impacts species transfer, and the crystal growth here is suppressed due to the lack of coordination between velocity and heat flow field.

Gas flow velocity in the axial direction along the crystal diameter is shown as Fig. 8. Similar to the growth rate distribution in Fig. 3, the curve shows an inverted V-shape in beginning, and maximal value reaches 0.05 m s^{-1} in the center of seed. As the crystal grows thicker, velocity in the center decreases quickly and maintains at about 0.025 m s⁻¹ due to recrystallization in the center of the SiC source surface. The law of values change matches well with growth rate in different stages, which again proves the field-coordination theory and provides an effective way to optimize crystal growth condition considering both temperature field and flow field. To understand why the velocity curve looks uneven while the crystal growth interface is relatively smooth, we compared the changes of temperature gradient and flow velocity in the early stage. At 0 h, seed temperature distribution is strongly influenced by the whole environment (Fig. 4), but the junction of seed and polycrystalline growth surface (area C) have a special condition. Seed edge grows slowly and even decomposes, while outside the edge there is a very large temperature gradient beyond -25K cm⁻¹, which results in the formation of an inverted V-shape



Fig. 8 Gas flow velocity around the seed in axial direction.



outline, with the bottom of the "V" on the seed side. For the plane where the seed crystal is located, the outline becomes Mshaped, like wave lines. Because the temperature and flow velocity are affected directly by crystal shape, isotherms and contours are like two non-coincident wavy lines (Fig. 5 and 6), resulting in evident velocity fluctuation (from the junction to inside). Even if the figure shows a transient growth rate and changes gradually, we still can find residual traces (Fig. 2a).

3.2 Comparison of different optimization methods

3.2.1 Temperature distribution. Focusing on temperature distribution in the growth cell and flow field distribution around the crystal, three different designs are put forward to handle the "M" surface and side polycrystal, as shown in Fig. 9. Compared with the basic design (design A), design B has different top insulation to enhance thermal insulation performance around the viewport, and designs C and D are based on design B, but have a wider groove on top, small-diameter viewport and graphite ring on the powder surface, while design C uses full-size 180 mm diameter seed, and design D uses a shelter structure where the polycrystal forms. All growth conditions are the same for these designs during the simulation and experiment, except the height of coils in design C, which is raised by 40 mm.

Temperature distributions during crystal growth were obtained and are shown in Fig. 10a. Both designs A and B have greater temperature difference, with twisting isotherms along the seed especially in the center and edge of seed, and slight blue can be found outside the edge of the seed. Compared with the former, design B shows smaller temperature change along the seed, but the isotherm still has a conspicuous "M" shape. In contrast, design C and design D have smoother isotherms



Fig. 10 (a) Temperature distribution, (b) temperature gradient distribution of different designs.



Fig. 11 Distribution of flow field of different designs.

especially around the crystal, which present flat, with slightly convex shape. Meanwhile, it should be pointed out that designs C and D have lower temperature, about 40 K, than design A because that small viewport makes heating power smaller in the condition where we fixed the temperature parameter in the monitor point. Besides the heating condition change, graphite ring and shelter structure play an important role in helping transfer heat to the center and influence temperature distribution both in the center and side of seed.

Fig. 10b provides a clear view of the temperature gradient along the radial direction. Blue color in the crystal means temperature along the radial direction changes a little, while red color suggests a violent increase. Unlike design A and B with about 6 K cm⁻¹ around the seed, parameters in design C and design D in the whole crystal remain at about 1 K cm⁻¹, which will reduce inconsistency in different areas of crystal. Around the seed edge (area C), designs A and B have negative values leading to the lowest temperature, and design D has a positive value close to the structure, which means the temperature gradient in axial direction here is negative and harmful to crystal growth. Meanwhile, greater value changes around the graphite structure show great potential to influence crystal growth conditions by changing the structure inside the crucible.

3.2.2 Flow field distribution and crystal shape. To study the angle between velocity vector and temperature gradient, the temperature and velocity streamline in the growth cell were obtained and are shown in Fig. 11. Values of velocity are presented by different colors. In area A, designs C and D have similar velocity vectors, nearly perpendicular to the temperature gradient. In areas B and C, velocity streamlines in designs C and D appear in a more vertical direction, which would carry more species to the growth interface and enhance crystal growth here. Different from designs A and B, the polycrystal in area D will no longer grow in designs C and D, because the former ones use full-size seed and the latter cannot even supply the gas species needed, limited by the shelter structure. It should be pointed out that due to lack of shelter structure guiding gas transfer, the flow velocity in design C around the seed edge has a smaller angle than in design D.



Fig. 12 (a) Profiles of SiC crystals, (b) thickness of SiC crystals.

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Fig. 13 Crystals of different designs.

Subsequent simulation calculation predicts different thickness of crystals with different deposition uniformities under the same process conditions. The profiles of crystals and crystal thickness along the radius were obtained and are shown in Fig. 12. Compared with design A, design B has a slower growth rate, about 160 μ m h⁻¹ in the center of crystal, and the difference between the thickest part and thinnest part is 6.5 mm, while it is about 11 mm in the former one. It goes without saying that we get a more uniform growth condition along the seed by changing the top insulation. As result, the polycrystal in design B is also thinner. Designs C and D show real nearly flat interface; their convexity is much smaller than both designs A and B after two process times. As expected, design C prepared a 180 mm diameter crystal with little "M" shape; there are some polytype cracks and even polycrystals produced around the seed edge near the crucible wall. Design D has 14.8 mm in the center and 9 mm in the thinnest part, showing the best profile of the four designs, benefitting from the best conditions.

3.3 Experimental crystals and test results

Under different hot-zone conditions, four crystals were prepared as shown in Fig. 13. In designs C and D, a lower gas pressure of 500 Pa was used to increase the growth rate, and the growth time was set to 150 hours. A 170 mm diameter seed obtained by expanding diameter method was used in design C, and the crucible wall thickness was slightly adjusted to adapt to the size of the seed. It is a pity that we got a crystal with edge cracks using design C, which is 15 mm in length, and the effective diameter of the crystal is about 160 mm. However, this crystal with smooth surface still shows the advantages of optimized thermal field. Design D has the best crystal shape and appearance with 16 mm effective thickness and 21 mm maximum thickness, and its growth rate increases to about $100-140 \ \mu m \ h^{-1}$, almost twice as much as predicted by simulation due to different vapor pressures. Anyway, these results



Fig. 14 Raman mapping of substrate.

show good consistency in crystal surface convexity and edge morphology since we focus more on trends than thickness.

Considering that crystals A to C have polycrystals near the edge, increasing the crack risk in the process is a major concern. After multiple procedures, substrates from crystal D were obtained after chemical mechanical polishing (CMP), and the middle ones were further tested. Different SiC polytypes have different structures, so the phonon Raman spectra show obvious differences between the two polytypes. Raman spectrum scanning was used, and the mapping shows that the 4H polytype proportion of this substrate is 100% (Fig. 14), suggesting an optimized design with better field coordination plays an important role in providing stable crystal growth conditions.

Micropipes are the main factor leading to the performance failure of SiC power devices. We used SICA88 to measure the distribution of micropipes as shown in Fig. 15, which is a common method used without any damage to the wafer. Results show a low micropipe density of about $0.09/\text{cm}^{-2}$ in this substrate, which is far better than the production grade of the industry standard. These results indicate that maintaining stable step flow growth is conducive to healing micropipe defects. Since the formation of micropipes is related to the growth conditions and seed crystal, the research proved the feasibility of zero micropipe crystal preparation using low-grade seed, which requires more work to do.



Fig. 15 Micropipe distribution of substrate.

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Fig. 16 Resistivity distribution of substrate.



Fig. 17 Surface characterization of substrate: (a) bow/warp @3700×, (b) flatness @17 000×.

Conductivity is a key indicator of conductive crystals. We use a noncontact and nondestructive eddy current method to test its resistivity, as shown in Fig. 16. Results show that the average resistivity is 0.022 Ohm cm, meeting the industry requirements of 0.015–0.025 Ohm cm. 80% of surfaces have resistivity no more than 0.017 Ohm cm, and large values appear only at the edge of the wafer, about several millimeters, showing great distribution uniformity.

Geometric parameters of the wafer reflect both the postprocessing technology and internal stresses of the crystal. Surface geometric parameters were measured using the Tropel FlatMaster, as shown in Fig. 17. The wafer has small values for the bow and warp parameters, 1.8 μ m and 10.9 μ m, respectively. Meanwhile, it has good surface flatness too, with a total thickness deviation of 2.6 μ m and maximum local thickness deviation of 1.5 μ m. This result clearly suggests a high quality of this crystal with a nearly flat and slightly convex interface.

4. Conclusion

We have established a model of the growth process of large-size silicon carbide crystals using a numerical approach. The crystal morphology predicted by the simulation matches well with the experimental crystals. After studying the changes in the growth interface and the distribution of physical fields during the growth process, distributions of thermal field and flow field are optimized to meet field-coordination effects, and a nearly flat and slightly convex growth interface was obtained. A high-quality 150 mm single crystal with a center thickness of 21 mm was prepared using the optimized thermal field, with micropipes less than 0.1 cm^{-2} and a uniform resistivity distribution. Our work introduces the variation characteristics of temperature fields and so on during the growth of silicon carbide crystals, and provides a simple and feasible method to improve the quality of large-sized single crystals using the PVT method.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have influenced the work reported in this paper.

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