Reprinted from

Space Processing of Materials

4–5 August 1996
Denver, Colorado

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conditions of low gravity where there would be less stirring taking place and the effective compositional gradient in
Directional Solidification of Mercury Cadmium Telluride During the Second United States Microgravity Payload Mission (USMP-2)


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ABSTRACT

As a solid solution semiconductor having a large separation between liquidus and solidus, mercury cadmium telluride (MCT) presents a formidable challenge to crystal growers desiring an alloy of high compositional uniformity. To avoid constitutional supercooling during Bridgman crystal growth it is necessary to solidify slowly in a high temperature gradient region. The necessary translation rate of less than 1 mm/hr results in a situation where fluid flow induced by gravity on earth is a significant factor in material transport. The Advanced Automated Directional Solidification Furnace (AADSF) is equipped to provide the stable thermal environment with a high gradient, and the required slow translation rate needed. Ground based experiments in AADSF show clearly the dominance of flow driven transport. The first flight of AADSF in low gravity on USMP-2 provided an opportunity to test theories of fluid flow in MCT and showed several solidification regimes which are very different from those observed on earth. Residual acceleration vectors in the orbiter during the mission were measured by the Orbital Acceleration Research Experiment (OARE), and correlated well with observed compositional differences in the samples.

KEYWORDS: Microgravity, Crystal Growth, HgCdTe, Solidification, Fluid Flow, Furnace

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instrument location are transposed to the AADSF position. As can be seen, the first attitude was of very low total residual acceleration, but the vector in the Z direction points from the solid to the liquid. The second attitude has a large component caused by the gravity gradient effect of the AADSF being 13 feet above the c.g., but the vector normal to the interface is stabilizing from liquid to solid. The final attitude also has the large transverse vector, and in this case has the destabilizing component from solid to liquid.

Figure 7 illustrates the history of the growth of the mercury cadmium telluride in terms of the mission timeline. The top scale is in Mission Elapsed Time (MET), and tells in days, hours and minutes the duration of the mission from launch. The y-axis shows the cadmium telluride mole fraction (x in Hg$_1$,Cd$_x$Te$_{1-x}$), while the x-axis is from measured values along the crystal. The solid line is a modeled diffusion controlled composition profile based on the D value shown. Post growth analysis indicated a supercooled region of 2.2 cm and this has been taken into account during the modeling. This distance is longer than the ground based crystals demonstrate. The MET scale is not linear, because the growth rate is not linear; it has been plotted to simulate the real time for a given distance, and in particular those times at which attitude changes took place. The translation rate used was 0.79 mm/hr, which was thought to be slow enough to avoid constitutional supercooling even in the space environment with no stirring adjacent to the interface. To provide a time indication of the changes in attitude, it was decided to increase the translation rate by a factor of 15 for 5 minutes, followed by a return to the initial rate. This change was made at MET 6/15:37 and was planned to give 4 hours of “recovery” time prior to the attitude change shown in figure 8. This would involve a translation of almost 1mm in five minutes. It was anticipated that breakdown and dendritic growth would occur thus enabling the position and shape of the interface to be determined. As will be shown later breakdown did not in fact occur.

There are five sets of compositional readings present on figure 8. The solid triangles are calculated from the densities of 2 mm wide wafers cut from the boule. These represent an average composition over that entire volume. Due to the need to cut in other orientations, density reading were made only near the beginning and end of the boule. The ampoule is conical so that the first to freeze material is of smaller diameter; density reading were not made. The other four sets of readings were taken with energy dispersive analysis, utilizing the HgM, CdL and TeL x-ray emission lines. The scanning electron microscope was used in the spot mode with 10 keV accelerating potential. Despite the fact that the surfaces were curved good totals were obtained between 99 and 101% after ZAF (atomic number, absorption, fluorescence) corrections were made. The J7 position referred to in figure 8 indicates an electrical jack on the furnace which pointed directly toward the front of the orbiter. This location was carefully tracked during the opening of the cartridge and the subsequent dissolution of the fused silica ampoule in hydrofluoric acid. Of specific interest is that each surface exhibits a different compositional trend, particularly over the first half of the growth. The J7 + 90 position consistently has a high x value, while the J7 and J7-90 positions are consistently lower. It should also be noted in this region that the density readings are lower than any of the surfaces and that they do follow the calculated diffusion data. In this sense they are similar to the ground testing.

At the position corresponding to the rate change, there is an obvious drop in the x-value corresponding to the inability of the boundary layer to be replenished by cadmium fast enough to maintain equilibrium solidification. Figure 8 is an etched region through this part of the material, and there is no indication of dendritic growth or breakdown; in fact the grains continue to grow through both the rate changes. Following the initial drop in composition, there is an increase in the cadmium content when the material reverts to the original slower rate. Following the second attitude change at MET 8/00:30 it can be seen that all the surface curves deviate not only from the position of the diffusion controlled tendency, but also from the shape. Furthermore, the EDS readings indicate that the surface CdTe concentration is lower than the mean value indicated by density. This is the opposite of the normal tendency, and is indicative of complete mixing. Cross sections of material from this area have not been made yet, but the indication is that there is more radial homogeneity than would normally be seen in this material during the final transient.

Figure 7 shows the OARE data transcribed to the cross sections of the crystals during the 3 main attitude periods of the mission. The data are shown in the form of the resolved component parallel to the interface, and also the vector perpendicular to the interface. As can be seen in the first and third attitudes, this latter was unfavorable. The results exhibit considerable radial asymmetry in the first attitude; the degree of inhomogeneity is in fact worse.
than in the ground based equivalent experiment. Figure 9 shows a cross section of the crystal grown in the first and second attitudes. The first attitude (+YVV/-ZLV) gives rise to very inhomogeneous material despite the low total vector. This sample is, in fact, worse than normally seen on the ground and worse than the ground truth sample run to simulate the mission. Clearly the second attitude with the vector in a favorable direction is more homogeneous. In both attitudes one can see a tendency for the heavier HgTe-rich material to be aligned with the vector. This a clear indication of three dimensional fluid flow. In this case of the material grown in the -XLV/-ZLV attitude the material is much more homogeneous. Even though the transverse vector is three times as large (1.55 µg₀ versus 0.55 µg₀), the perpendicular vector is stabilizing and so the material is more uniform. The ratio of perpendicular to transverse is very low, on the order of 0.3. Ideally this ratio would be at least 4:1. The material is still nothing like as uniform as would be achieved with diffusion control, and, for the same temperature profile, is inferior to that produced by growth in an axial magnetic field.¹⁰ Nevertheless, the improvement is clearly the result of a reduction of the fluid flow. With improved microgravity conditions, including a much reduced transverse vector and a perpendicular to transverse ratio of 4:1, conditions closer to diffusion control can be obtained. This is one of the objectives of further microgravity missions. The first mission has clearly shown the effect of three dimensional fluid flow on the solidification characteristics of a material such as this having a wide separation between the liquidus and the solidus.

6. ACKNOWLEDGMENTS

The authors are indebted to C. Bahr, D. Lovell, J. M. Jones and G. Nelson for their dedicated technical assistance, particularly in the fields of furnace fabrication and operation and glass-blowing. The design and fabrication of the AADSF by personnel at MSFC and Teledyne Brown Engineering has made this project possible. The support, both moral and financial of the Microgravity Science and Applications Division at NASA Headquarters has been enthusiastic, warm and helpful.

7. REFERENCES

Figure 1. Interface Region showing Convective Cells

Figure 2. Gravity Effects in the Solidification of Pseudobinary Solid Solution Systems. (a) Rejected component less dense than bulk liquid results in thermosolutal convection throughout the liquid volume. (b) Rejected component more dense than bulk liquid results in interface shape instability in presence of radial temperature gradient.
Figure 3. CdTe Compositional Distribution for Two Different Growth Rates

Figure 4. Typical Quenched-in Interface in HgCdTe
Figure 5. Temperature of Solidification at Interface

<table>
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<th>Orbiter Attitude</th>
<th>Transverse Residual Acceleration</th>
<th>Residual Axial Acceleration</th>
<th>Resolved Transverse</th>
<th>Axial/Transverse Ratio</th>
<th>Total Resolved</th>
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<td>Hot</td>
<td>1.45</td>
<td>-0.31</td>
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Figure 6. Orbiter attitudes and residual acceleration vectors AADS during USMP-2
Figure 7. Mission Timeline for USMP-2 Showing Composition of Crystal Grown

Figure 8. Microstructure of HgCdTe during the Rate Change (etched in Polisar II)
Figure 9. Compositions of Material Grown During two Attitudes of the Orbiter