Liquid Crystalline Polymer

User’s manual for Sumikasuper LCP

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<contents>

1. Introduction of liquid crystalline polymer
   1-1 General properties of LCP 1
   1-2 Thermal resistance 7
   1-3 Moldability 10
   1-4 Mechanical properties 11
   1-5 Anisotropy 13
   1-6 Market situation of liquid crystalline polymer 14
   1-7 Conclusion 16

2. Application of liquid crystalline polymer and its technology 17
   2-1 Connector for PC, mobile, digital camera, etc. 17
      (1) Surface-Mount Technology (SMT) 17
      (2) Relationship between compound formulation and warpage 18
   2-2 Bobbin for back-light transformer of LCD 21
      (1) Proceeding of the back-light system of LCD 21
      (2) Requirement for the inverter bobbin 21
   2-3 OPU (Optical Pick-Up) actuator bobbin for CD-ROM, DVD, etc. 23
      (1) Requirement for OPU actuator bobbin 23
      (2) Relationship between modulus and thermal resistance 24

3. Injection molding technology for liquid crystalline polymer 25
   3-1 Control System of Injection Process
      (1) Open loop control 25
      (2) Closed loop control 26
   3-2 Thin-wall fluidity of LCP and characteristic of injection machine
      (1) Experiment 27
      (2) Result 29
   3-3 Flow hesitation
      (1) What is “Flow Hesitation”? 32
      (2) Flow behavior of LCP 32
   3-4 Metering of LCP
      (1) Principle of unstable metering 35
      (2) Solution of unstable metering 37

4. Improvement of warpage 39
   4-1. Theoretical background of the warpage problem 39
   4-2. Relationship between warpage and flow pattern 41
   4-3. Relationship between the depth of core-out and the warpage 44
   4-4. Case study
      a) Board to board connector (0.5mm pitch) 46
      b) S/O DIMM 46
      c) PGA socket 48

5. Trouble shooting during injection molding 50
   5-1 Outline for trouble shooting 50
   5-2 Itemized discussion 52
   5-3 Black spec  -burning & carbonizing- 53
   5-4 Blister & Bubble  -classification & improvement-
      (1) Comparison between “blister” and “bubble” 54
      (2) Reduction of the blister at soldering 54
         a. Effect of moisture 55

- iii -
b. Retention of resin at the inside of injection machine 57
c. Temperature settings of injection machine 58
   ~Difference between setting value and real resin temperature~
d. Relation between retention of LCP in cylinder and blister 60
e. Unbalance between cylinder size and molding volume 61
f. Purging method of Sumikasuper LCP 62
(3) Reduction of the bubble after molding 65
   a. Dragging of the air into the melt 65
   b. Suitable injection condition 65
   c. Suitable cavity design 65
5-5 Crack 68
   (1) Cracking problem for Coil Bobbin 68
   (2) Side-wall cracking for Case part 69
   (3) Weld crack at the hole of Board to Board Connector 70
5-6 Flash 73
5-7 Flow mark 74
5-8 Metering 75
5-9 Short-shot 76
5-10 Sticking 79
5-11 Warpage 80

Reference 81
Preface

This booklet was originally planned to provide the presentation to molding engineers of our customers, because there was little literature or references about improvement and solution of molding problems for Liquid Crystalline Polymer (designated as LCP hereafter).

In early 1990s, it was a period of actual growth of thermotropic LCP industry for electric and electronic parts such as connectors, relays or coil bobbins for personal computers, mobile phones or digital cameras, etc. It was also the same movement of developing the new industry for IT business. Many companies of this business area have wished to use this eccentric polymer for such new products, however, sometimes they faced several molding problems. It seems that the behavior of LCP is very different from conventional engineering plastic, so sometimes molding engineers found it unmanageable.

The word of LCP is attractive and many researchers have worked about this region. However, almost all result of such works merely provided the discussion between theoretical back-ground and obtained data, or disclosed some of interesting phenomena during molding, such as “relationship between shear rate and apparent viscosity”, “effect of statistic orientation to morphology”, etc. Those data or information, indeed, are very useful for researching or developing of new material based on LCP by specialists of polymer chemistry.

On the other hand, there are many useful and excellent literatures about plastic processing. Such literatures disclose not only principle of the mechanism or theoretical background of processing, but also the reason of molding trouble and its solution. However, such current knowledge sometimes prevents understanding the behavior of LCP and causes misunderstanding of improving immediate problem or hides the actual reason from the engineer.

In view of above situation, the first step of this booklet gave suggestions or solutions to solve molding trouble of LCP and to help understanding of LCP behavior. Thereafter we filled more contents of useful information over 10 years and it becomes workable brochure for not only experts but also beginners of LCP molding.

The 1st section expresses chemical and physical properties of LCP which will become the basic to understand LCP behavior theoretically. It also discloses the market situation of LCP industry.

The 2nd section discloses examples of LCP application and concept for electric and electronic parts. It will help you when you choose more suitable grade for your applications or items.

The 3rd section shows a methodology from a little different point of view. This section discloses that the performance and specification of injection molding machine are also very important for LCP molding. It tends to be ignore, but it sometimes influences whether you obtain molding part successfully or not.

The 4th section indicates several know-how to solve warpage problem for connectors with actual examples. This information will help you to develop the latest precision connectors or such electric and electronic parts successfully.

The last section, 5th is the most important section when you suffer the molding trouble. This section covers almost whole solution during using LCP.

December, 2006
Electronic Materials division
Sumitomo Chemical Co., Ltd.
1 Introduction of Liquid Crystalline Polymer

Liquid crystalline polymers (LCPs) are widely used in many types of electric and electronic parts due to their superiority in high heat solder resistance, high-temperature strength, dimensional stability, overall good chemical resistance, low flammability and low water absorption. LCPs especially exhibit better thin-wall fluidity and moldability than any other engineering plastics thanks to their extraordinary low melt-viscosity. This is also the reason why LCPs are now used for the latest designed and highest precision molding parts. In recent years, electric and electronic parts molded with LCPs have become more important to the IT related industries (as well as many consumer markets) because of Surface Mounting Technology (SMT). There is much information regarding LCP resins. However, its properties and performance are often not understood thoroughly. For this reason, summarizing the basic and general properties of LCP should be considered. In this section, we would like to survey the general properties of LCP by comparison with conventional engineering plastics.

1-1. General properties of LCP

Engineering plastics can be classified as plastics having over 100°C of TDUL (Temperature of Deflection Under Load: usually measured under 1.82MPa). Plastics having over 150°C of TDUL are called Super Engineering Plastics (Table.1-1).

![Classification of plastics](image)

Table 1-1. Classification of plastics

<table>
<thead>
<tr>
<th>General Purpose</th>
<th>Engineering Plastic (E.P.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP, PE, PMMA, PS, ABS, ...</td>
<td>PP, PE, PMMA, PS, ABS, ...</td>
</tr>
</tbody>
</table>

Fig. 1-1. Relationship between TDUL and continuous service temperature

Super engineering plastics not only have higher TDULs, but also superior long-term heat stability. TDUL only
indicates short-term heat stability. Fig. 1-1 indicates the relationship between TDUL and continuous service temperature. This parameter equals long-term heat stability and is measured under the conditions described in UL 746B. As shown in the figure, super engineering plastics indicate both higher TDUL and over 200 °C of continuous service temperature. This allows these plastics to be suitable for SMT soldering or other high heat treatments (in general, SMT soldering temperature is over 220°C). Conventional engineering plastics will exhibit similar TDULs when reinforced by glass or fibers, and thus perceived higher heat resistance. It should be noted; however, that these polymers do not have as high of continuous service temperature resistance. This means that conventional engineering plastics are not suitable for high heat treatment such as SMT soldering.

It is very important to think about the hierarchy of molecular structure when considering the macro dynamics of the polymer. The molecule of polymer can often be compared to a chain or a thread of yarn. In this case, let us presume that there are about 30 pieces of yarn cut at 30cm length and then crumpled into a ball. The crumpled yarn-ball is then put into a funnel that has enough spout diameter in order to pull the pieces out. At first, the yarn will not pull out because the diameter of yarn-ball is larger than that of spout. If you wish to pull the pieces out, you must push the ball with stick or rod from the topside of the funnel. This is very similar to the molecular situation during injection molding. Injection molding is the molding system where molten polymer molecules with random coil shapes are pushed into the mold by high pressure. They then take the form from the mold after cooling (Fig.1-2).

Parts of the molecule chains are sometimes folded regularly during the cooling process. Such folded polymer chains will assemble together through affinity. Assembled portions are called “semi-crystalline” or “crystallites”. Since these molecules form in lines, they are called “crystals” (Fig.1-3 a). Such polymers having both crystallites and ‘not crystallized’ portions (usually called amorphous) are called “crystalline polymer”. In general, the crystallinity of crystalline polymers is up to 30% before annealing or heat treatment (for increasing the crystallinity). Since the polymer molecules are very long and tangled, the entire polymer molecule cannot crystallize completely.

In addition, some chemical structures hesitate to form crystallites and the molecular chain will be packed randomly. These polymers are called “amorphous polymers” (Fig.1-3 b). By this reason, the crystallinity of amorphous polymer is estimated as 0%.

The crystallites of a crystalline polymer scatter the visible rays. Thus crystalline polymers are often opaque. On the other hand, amorphous polymers (and the crystalline polymers having smaller crystallites than the wave length of visible rays) are usually transparent. Of course filled or reinforced polymers should be excluded in this case even if they are amorphous.

Both crystalline and amorphous polymers have Tg (glass Transition Temperature) where the molecules start moving by heat energy. Only crystalline polymers indicate Tm (Melt Temperature) where the crystallites melt. Both Tg and Tm are the typical measurements to determine the polymer properties.

Some polymers have rigid rod molecular structures that are described by the liquid crystalline transition temperature (Tl,c). In general, these plastics are called Liquid Crystalline Polymers (or designated LCP). Here, we would like to presume the simple molding model again using matchsticks.

Let us imagine that a matchbox filled with matchsticks is overturned above the funnel. At first, the matchsticks will remain in the upper portion of the funnel. However, by adding a small vibration that puts the direction of the matchsticks in order, the matchsticks will flow out from the throat of the funnel very smoothly (Fig.1-4). In this case,
matchsticks are like the rigid-rod molecules of liquid crystal polymers. The action of adding vibration is like adding shear stress of molding to the polymer.

You can also imagine that the rigid rod structure of a matchstick will be more suitable than the yarn-ball due to the yarn never flowing out from the funnel throat without higher force. This resembles LCPs, which have extremely low melt viscosity and higher flowability and do not need the higher injection pressures required for conventional polymers.

Since the molecule of LCP is rigid, it will form semi-crystalline structures that are strongly oriented in one direction. Bundles of the molecules align in both directions so that the polymer matrix does not indicate the directional properties. The bundles are called the “domain” structures of liquid crystal. The domain structures are not detected easily due to the difference between the domains not being clear thermodynamically. In other words, domain like shape will be observed under the polarized microscopic investigation but the size of the domain is undetectable.

If shear stress is added to the melted LCP, the rigid-rod molecules are easily aligned in the same direction of the shear stress. After cooling, the melt will solidify and preserve this highly oriented three-dimensional structure (Fig.1-5). This directional property is called “anisotropy”.

The first scientist who named liquid crystals was Otto Lehmann in 1889 after discovering the birefringence phe-
nomenon by Friedrich Reinitzer in 1888. The low molecular weight liquid crystals; however, demonstrated relatively different history with polymer liquid crystals. Such investigations have helped to develop the current LCD (Liquid Crystal Display) etc. The first notable work of polymer liquid crystals started from the pioneering study by Onsager, Ishihara, and Flory. Flory predicted lyotropic liquid crystals that indicate liquid crystallinity in dilute solutions. This stimulated many researchers’ interest, and now there is much literature about liquid crystals.

It is important to point out clear fundamentals of commercially produced LCPs -

![Figure 1-5 Three-dimensional structure of liquid crystalline polymer](image)

The reason behind liquid crystallinity is not completely clear, but we know that the rigidity of the molecule and induced anisotropy of its shape are an important factor. Such molecules are called “mesogen”, which is derived from the former explanation of mesomorphic state. Basically, this is intermediate phase between solid and liquid states. Many chemical structures have been examined and we know that several chemicals indicate liquid crystallinity. The most important mesogen is p-hydroxybenzoic acid for LCP. This raw material can be polymerized by itself (self polymerization); however, it is obtained in an insoluble and unmelted form. It is usually co-polymerized by other raw materials to help reduce the process temperature. Of course, such copolymer still keeps its excellent heat and solvent resistance. In fact, few solvents attack LCPs with the most noted one being a mixed solvent of pentafluoropnenol and chloroform.

Another question is to determine when the liquid crystal first appears. Just like low molecular weight liquid crystal, LCP appears in tow kinds of circumstances - in the solvent as solution and in high temperature circumstances over the \( T_{LC} \). The former is called “lyotropic” LCP, and the latter is called “thermotropic” LCP. Some of the LCP are known as commercially produced products as shown in Fig.1-6. In this book, we would like to focus to thermotropic LCP, especially aromatic polyesters, because there are many kinds of commercial products in this field.

![Figure 1-6 Type of Liquid Crystalline Polymer](image)

The 3-dimensional structure of the LCP molecule is also important to consider - Fig.1-7. Most LCPs are nematic as the actual crystalline structure of commercially produced LCP has been studied. Some of the special chemical structures are known as cholesteric liquid crystalline, which comes from “cholesterol” with indicating light circularity. This sort of LCP is studied for the display application field, and it is not in scope of this book. 
The typical chemical structure of LCP is shown in Fig.1-8\textsuperscript{10}. Main-chain type LCP is now the most common. Side-chain and combined type LCP are proposed for LC display applications. They are used to improve problems with low molecular weight LC.

Among these structures, main-chain and nematic LCP is the most important, almost all LCPs produced commercially are included this category. Table 1-2 shows typical molecular structure with classification of TDUL. As seen in this table, all LCPs include a p-hydroxybenzoic group as a mesogen, but the other components are not equal. The different combination of reactive groups (monomers) gives the different thermal resistances. In general, LCP is classified in three types of categories. Type III is the lowest thermal resistance category and includes the early developed LCP X-7G manufactured by Eastman Chemical. Type I is the highest heat resistance category, and it indicates more than 260 °C of TDUL.

It needs to be emphasized that the chemical structure of each LCP is different from each other. This means that each LCP manufacturer may have different properties although total properties are categorized as “liquid crystalline polymer”. This is very similar to polyamides as well. For instance, PA6 and PA46 indicate much different thermal resistance; however, both of them have relatively higher water absorption and poor dimension stability vs polyesters. Different thermal resistance is derived by their own chemical structure, but the higher water absorption property is derived by the amide-bonding group.

LCP is similarly in the same situation. It is mostly affected to the difference of moldability. Several researchers have started to reveal this difference but no one has succeeded to describe it as yet. The difference of moldability is the number one cause of difficulty in molding shops.

Table 1-2. Typical molecular structure of commercial LCP
During molding process, the shear stress with the wall of mold induces the specific structure. In the case of injection molding, the melt polymer flows in what is called a “fountain flow” (Fig. 1-9). The shear stress will be generated between center flow and mold wall. After cooling, the record of this flow is highly characteristic.

As shown in Fig. 1-9, outer side of molded part is strongly aligned to the flow direction, although the center portion is aligned transverse or random direction. In general, this structure is called as Skin-Core structure. Fig. 1-10 shows the cross section of actual molded part and its schematic view to confirm this special structure. This is very similar as the structure derived by crystalline polymer; however, each skin and core layer of LCP indicates much more different mechanical and chemical properties from each other.

![Fig. 1-9 Schematic view of fountain flow](image)

The thickness of skin layer is usually 200μm under the appropriate molding conditions. However, molding conditions strongly effect the generation of this Skin-Core structure or thickness. If strong shear force is derived by high shear rate under appropriate temperature, the skin layer will be very defined until 200μm. However, if the shear force is lower, by the reason of lower shear rate and lower temperature, not only will the skin layer thickness be less, but also the Skin-Core structure can disappear. This will affect the fixed or stable character of LCP.

The core layer must disappear under 0.4mm (400μm) of the total thickness if the thickness of skin layer is always 200μm. However, the skin layer comprises many stratified sub-layers and micro-layers. Often some of the sub-layers or micro-layers must work like a core-layer during the route of melt flow.

This sublayer structure is like a shell of a pie or a bamboo sheath as shown in the picture of Fig. 1-10 left. Each layer is very thin and strong, but can peel of easily. For this reason, the skin material is generated when the surface of molded part is abraded or the gate is cut obliquely.

The reason of this abrasion is thought to be due to weak intermolecular attraction force (like van der Waals force or dipole-dipole attraction) between layers.

Since the generation of the skin layer and Skin-Core structure are the most important feature this must be optimized in the molding conditions.

---

<table>
<thead>
<tr>
<th>Type</th>
<th>Molecular Structure</th>
<th>TDUL</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type I</td>
<td><img src="image" alt="Molecular Structure" /></td>
<td>&gt;260°C</td>
<td>SUMIKASUPER Xydar</td>
</tr>
<tr>
<td>Type II</td>
<td><img src="image" alt="Molecular Structure" /></td>
<td>210 - 260°C</td>
<td>Vectra</td>
</tr>
<tr>
<td>Type III</td>
<td><img src="image" alt="Molecular Structure" /></td>
<td>&lt;210°C</td>
<td>Rodrun Novaccurate</td>
</tr>
</tbody>
</table>

*p-hydroxybenzoic group*
Fig.1-10 Cross section of molded part of LCP (right) and its schematic view (left)

We would like to compare the property among crystalline, amorphous and liquid crystalline polymers. As shown in Table 1-3, amorphous polymer indicates only Tg (glass-transition temperature), but crystalline polymer indicates both Tg and Tm (melting point). On the other hand, LCP indicates only T_{LC}, which is the transition temperature from solid to liquid crystal state as described above. This T_{LC}, however, usually depends on the shear stress during measurement. It means that a higher shear stress will decrease the T_{LC}, and lower shear stress will increase it. In the attached table, merits are listed in light blue zone and demerits are in light red zone. Thermal resistance, soldering resistance, and flame retardancy are mainly derived by chemical structure. Since the listed polymers commonly consist of aromatic monomers, all materials indicate relatively higher thermal resistance (>200°C).

Notable merits of LCP are higher soldering resistance and significantly higher flowability. On the other hand demerits are lower weld strength and strong anisotropy property.

<table>
<thead>
<tr>
<th></th>
<th>Amorphous</th>
<th>Crystalline</th>
<th>Liquid Crystalline</th>
</tr>
</thead>
<tbody>
<tr>
<td>SUMIKAEXCEL® PES</td>
<td>Polyphenylene Sulfid (PPS)</td>
<td>SUMIKASUPER® LCP</td>
<td></td>
</tr>
<tr>
<td>Tg 225 °C</td>
<td>Tg 85 °C</td>
<td>T_{LC} 320 to 400 °C</td>
<td></td>
</tr>
<tr>
<td>Thermal resistance (to 200 °C)</td>
<td>Thermal resistance (to 240 °C)</td>
<td>Thermal resistance (to 260 °C)</td>
<td></td>
</tr>
<tr>
<td>Dimension accuracy</td>
<td>Soldering resistance</td>
<td>Excellent soldering resistance</td>
<td></td>
</tr>
<tr>
<td>Creep Performance</td>
<td>Excellent solvent resist.</td>
<td>Solvent stability (except alkali &amp; steam)</td>
<td></td>
</tr>
<tr>
<td>Impact strength</td>
<td>High flowability</td>
<td>Superior high flowability</td>
<td></td>
</tr>
<tr>
<td>Boiled water resistance (160 °C)</td>
<td>High flame retardancy (V-0)</td>
<td>High flame retardancy (V-0 @0.3mm)</td>
<td></td>
</tr>
<tr>
<td>High flame retardancy (V-0 @ 0.4mm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Higher water absorption</td>
<td>Higher Degassing</td>
<td>Lower weld strength</td>
<td></td>
</tr>
<tr>
<td>Relatively lower flowability</td>
<td>Flash</td>
<td>Strong anisotropy</td>
<td></td>
</tr>
</tbody>
</table>

### 1-2. Thermal resistance

LCP realizes higher TDUL (Temperature of Deflection Under Load) than other plastics. Fig.1-11 shows temperature dependence of elastic modulus compared with crystalline (PEEK), amorphous (PES) and liquid crystalline (Sumikasuper LCP) polymers. As shown in this figure, it is clear that LCP keeps higher mechanical property over 200 °C.

LCP also does not have notable decrease of modulus as shown in PEEK at 140°C – meaning that LCP seems not to have glass transition behavior. Thermal analysis such as differential scanning calorimetry (DSC) also does not indicate the thermal transition as observed in other conventional crystalline and amorphous polymers. Careful examination will show a small slant of the modulus curve at around 120°C. S. Z. D. Cheng et al. recog-
1 Introduction of liquid crystalline polymer

zized that LCP having several kinds of chemical structures would indicate two different crystal structures in the temperature range of solid to nematic through annealing treatment. Other researchers have since expanded the region of study for several chemical structures of LCP. Some researchers have found that LCPs having Vectra like chemical structure have 2 or 3 kinds of relaxation. These are named as α, β- and γ-relaxation, which are very similar with conventional polymers. In this case, α-relaxation is very similar to conventional glass transition temperature (Tg). This change seems very similar with the change at Tg of crystalline polymer. In fact, a very small and dull peak is sometimes observed in tan δ curve of dynamic mechanical analysis at the same temperature region.

This means that it is not usually necessary to think about the Tg of LCP. This is also why any mold temperature up to molding temperature can be chosen for LCPs. For instance, the mold temperature of PEEK should be set at over 160°C, because its Tg is 144°C and higher than this temperature is necessary to achieve higher crystallinity after cooling. In this meaning, mold temperature has little impact on LCP properties. However, we should notice that at a little performance change occurred at Tg, which is pointed out by Ward et al.

Furthermore, LCP’s decomposition temperature is over 500 °C, which is much higher than most other plastics (Fig.1-12). This helps LCP to have very low out-gassing. This is due to the origin of outgas is usually in the heating of the decomposed material.
the tool or machine. This also helps to prevent several problems after heat treatment – such as IR reflow soldering or a post baking process. (Nevertheless sometimes the blister problem occurs after molding or after soldering. The reason and solution of this problem will be discussed later)

Fig.1-13 The evaluation method of outgas

In general, the main portion of outgas is an unknown organic portion\(^\text{18}\). Acetic acid, phenol, and its derivatives are also detected in outgas. The origin of such chemicals is further supported to be decomposition of LCP (Table 1-4).

![Fig.1-14 Comparison of total amount of outgas](image)

Table 1-4  Outgas portion of LCP

<table>
<thead>
<tr>
<th></th>
<th>unknown organic materials with low boiling point</th>
</tr>
</thead>
<tbody>
<tr>
<td>II</td>
<td>(\text{CH}_3\text{COH}) (acetic acid)</td>
</tr>
<tr>
<td>III</td>
<td>(\text{HO} \quad \text{O}) (phenol)</td>
</tr>
<tr>
<td></td>
<td>(\text{HO} \quad \text{C-OH} \quad \text{HO} \quad \text{C-OH}), etc. (phenol derivatives)</td>
</tr>
</tbody>
</table>

1-3. Moldability

LCP has very low melt viscosity property at molten state. For this reason, LCP’s exhibit excellent flowability. Fig.1-15 shows the temperature dependence of flow length at 1mm thickness (bar-flow) with conventional engineering plastics. As shown, LCP’s indicate much higher flow length but stronger dependence of temperature than the conventional plastics. The reason is due to its typical melt viscosity behavior as shown in Fig.1-16. After liquid crys-
1 Introduction of liquid crystalline polymer

At the transition temperature, the melt viscosity significantly decreases. As the temperature is increased more and more, there is lower dependency of the melt viscosity. It is called "plateau region" in general. The most preferable temperature region for injection molding is this transition temperature region, and its width is around 10°C. If the temperature is lower than this transition, the generation of Skin-Core layer will be insufficient because of insufficient melt of LCP. On the other hand, if the temperature is higher than this transition, the melt viscosity will be too low and uncontrollable for the molding. This is the reason why the molding condition of LCP is relatively narrow. The molding temperature should be set on this transition region of the melt viscosity.

A further important point is that the dependency of melt viscosity with temperature is relatively higher with LCP’s than other plastics. This tendency is similarly observed at the relationship between shear rate and melt viscosity. As shown in Fig.1-17, LCP’s show strong shear thinning property. Accordingly, LCP’s exhibit excellent flowability under moderate temperature and higher shear rate condition.

In contrast, this stronger dependence of melt viscosity with temperature and shear rate can cause other kind of problems.

![Graph 1](image1.png)

**Fig.1-15 Flow length (1mm thickness)**

![Graph 2](image2.png)

**Fig.1-16 Temperature dependence of melt viscosity**

The major problem is that it can bring quick solidification. After flowing into the mold cavity, heat will be removed from the melt polymer by the cooling effect of the mold. This small decrease of polymer temperature will cause of huge increase of melt viscosity. In extreme cases, polymer will not be able to flow. This effect is one of the merits of preventing flash and reason why LCP doesn’t exhibit this.

The second problem is that this effect can cause flow hesitation. Flow hesitation is mainly observed during injection molding of LCP. It can be seen to hesitate the flow into the cavity of the mold even if the cavity is near the entrance of the flow.
Since the flow of LCP is prevented by something (ex. small size gate, thin-wall portion, divergence, etc.), the shear rate will be reduced (Fig.1-18: corresponding to the shear rate change from $\gamma_1$ to $\gamma_3$). This reduction of shear rate induces great amount gain of melt viscosity (Fig.1-18: corresponding to the melt viscosity change from $\eta_1$ to $\eta_3$). As shown in Fig.1-18, the viscosity gain of LCP is comparably larger than that of amorphous polymer because of higher dependency between shear rates and melts viscosity of LCP. If this effect should occur, the melt viscosity of LCP gradually increases until the material will not flow any more. This effect doesn’t often correspond to the pressure loss, which is conventionally thought by general theory of material flow. Unfortunately, this is often observed as the molding problem like “short-shot molding” or “weld-line crack”.

To understand the melt viscosity property of LCP is very important to solve these kinds of problems.

**1-4. Mechanical property**

LCP’s have excellent mechanical properties. Table 1-5 shows Sumikasuper E6000 with an unfilled base resin compared with Polyethersulphone. Flexural modulus and TDUL of LCP are much higher than those of the PES. As described above, the orientation of rigid-rod molecule is induced by the shear stress with the wall of mold during the material flow in the cavity as so-called "fountain flow" (see Fig.1-9). Due to the fact that this orientation of the rigid-rod generates the skin layer, a molded article of LCP is reinforced by itself. That is the reason why LCP is sometimes called a "self-reinforcing material". This phenomenon derives the high rigidity and thermal properties of LCP’s. However, we should notice that the flexural strength is not high compared to the higher modulus. It suggests that LCP is more brittle then other plastics.

In addition, it should be noted that there are many differences between MD (Mechanical Direction) and TD (Transverse Direction) of both mold shrinkage and C.T.E. (Coefficient of Thermal Expansion) for LCP’s. Especially, C.T.E of MD is a negative value. This means that the dimension of MD will decrease with elevation of the temperature. This is induced by the orientation of rigid-rod molecule to the molding direction during processing. We have to note that anisotropy is the essential property of LCP.

On the whole, aromatic polymers have excellent flame retardant properties, and LCP is no exception. Both PES and LCP are classified as V-O via UL-94 regulations. This means that there is no chance to expand the flame through the article made by both materials. LCP’s exhibit such property at only 0.3mm thickness (actually under 0.3mm thickness,
but it is not proved because the thinner test piece less than 0.3mm is not obtainable) without any kind of flame retardant needed. This helps with ESHA and other environmental regulations and needs.

### Table 1-5 Comparison of mechanical properties for PES and LCP

<table>
<thead>
<tr>
<th></th>
<th>PES</th>
<th>LCP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity</td>
<td>1.37</td>
<td>1.38</td>
</tr>
<tr>
<td>Mold shrinkage (%)</td>
<td>0.6</td>
<td>0.22</td>
</tr>
<tr>
<td>Flexural strength (MPa)</td>
<td>129</td>
<td>106</td>
</tr>
<tr>
<td>Flexural modulus (MPa)</td>
<td>2,550</td>
<td>6,860</td>
</tr>
<tr>
<td>TDUL (°C)</td>
<td>203</td>
<td>262</td>
</tr>
<tr>
<td>C.T.E. ((10^{-5/°C})</td>
<td>5.5</td>
<td>-0.5</td>
</tr>
<tr>
<td>Flame retardancy</td>
<td>V-0</td>
<td>V-0</td>
</tr>
</tbody>
</table>

Furthermore, the thickness of skin layer is almost specific as 200μm as described above. As the total thickness becomes thinner, the ratio of skin layer comparatively increases against the total thickness. Since the skin layer is formed by integration of highly oriented fibrous semi-crystals of rigid-rod molecules, it derives high mechanical properties. As a result, the strength of LCP will gradually increase with decreasing of the thickness (Fig.1-19). This is typical and unusual property of LCP, which does not observed at conventional plastics.

The skin layer has another important feature - lower permeation of gases. Basically the crystal structure has a very high density and is strongly integrated by molecules. Accordingly the low molecular structures like gaseous organic chemicals or ions cannot easily permeate through crystalline portion. It is the same for LCP, because the skin layer of LCP is the highly crystallized portion of LCP molecule. For this reason, LCP’s exhibit excellent gas barrier properties. Fig.1-20 shows gas permeability data for many kinds of polymer films. It reveals that water and oxygen permeability of LCP is superior to any other plastic film and it is close to that of Aluminum Foil.
1-5. Anisotropy

As mentioned, LCP’s have strong anisotropic property and this often causes molding problems such as warpage, weld crack, etc. Here, we would like to focus attention on this issue.

Fig.1-21 shows the mold shrinkage for both MD and TD. The dotted line indicates unity shrinkage for whole directions, meaning that the material is isotropic if the plot is on this line. Although crystalline polymers such as PPS or glass filled material such as PES 4101GL30 indicate weak anisotropic behavior, LCPs exhibit stronger anisotropy, especially non-filled material (E6000). This means that LCPs essentially have stronger anisotropy, but GF filling reduces this property. This is opposite of most conventional plastics. This is one of the key factors when you select material and the formulation of fillers for the application.

Moreover, LCP has very low weld strength. As shown in Fig.1-22, the weld portion is the portion where the separated flow gathers together again. At this moment, the flow pattern should be the right side in the Fig.1-22. The direction of flow at weld portion should be transverse direction against the main flow direction (MD). In this case, the weld portion forms the very similar structure of skin layer against the flow direction. Since skin layer is derived by the integration of thin sub-layers and the affinity between layers is very weak, the layers are easy to peel off. In addition, there is a pulling force at the weld portion because the mold shrinkage at weld portion is TD vs MD for the main flow portion. This kind of different structure also becomes the cause of weakness. It seems like the kink of crystal, and it will be the over concentration of inner stress.

At last, the weld portion of a molded article of LCP is essentially very weak and unavoidable. In general, the mechanical strength of weld portion is 1/4 to 1/5 of normal portions. The filling of GF usually helps to improve the weld
strength, however, it should be noted that adding of minerals or having too much content of fillers can reduce it again. In general, longer GF or fillers works better for this property.

![Schematic view of flow pattern at weld portion](image)

Fig.1-22 Schematic view of flow pattern at weld portion

### 1-6. Market situation of liquid crystalline polymer

Several suppliers of LCP are known in the world. Fig.1-21 shows manufacturers of LCP and their history of commercial production. At the moment, there are 6 major suppliers, half of them are in USA and other are in Japan. Former customers of LCP were distributed in both USA and Japan 10 years ago, but now customers have expanded to Europe and Asia (south-east and east). In recent years, the Asian market share has become 80% of global demand, because customers have moved their production facilities to Asian countries such as China, ASEAN countries, Korea, etc.

On the other hand, the suppliers of LCP have reduced during this decade. Additionally several suppliers worked for marketing or trial production purposes in the past; however, the selection advanced. Now there are 4 major trade names and subsequent 3 or 4 trade names.

|   |                |     |            |           |            |
|   | Nihon Ekonol   |     | Resale Nisseki Chemical |
| II | 1984 Hoechst Celanese | 1998 Ticona | Production |
|   | Resale Polyplastics | 1994 Toray |

Fig.1-23 History of LCP production

Table 1-6 shows the capacity of major suppliers of LCP with their trading names. On 2004, total capacity of neat resin reached around 23,000 T/y, and this satisfies around 80% of demand. However, from 2003 to 2004, IT market including not only Personal Computer market but also OA applications expanded again with several 10% ratio than a year ago. For this reason, the demand and the supply of 2004 were tight. It seems that all suppliers are planning to expand their production capacity within a few years.

Table 1-7 summarizes the temperature range of each trade name of LCP. As shown this table, the strategy of each
supplier can be seen. Major suppliers make efforts to expand their portfolio line-up to cover wide temperature ranges for wide variety of applications. Usually, the suitable temperature range is determined by molecular formulation. Each supplier prepares to supply several base resin grades having different molecular structures. Since each molecular structure requires the most suitable temperature region during processing, keeping the temperature should be noted. Especially, since LCP has much excellent thermal property, the suitable processing temperature is also very higher than conventional plastics. Sometimes it reaches at over 400 °C. It is very important to confirm that your molding machine is suitable for this processing temperature before using.

Table 1-6 Production capacity of LCP supplier

<table>
<thead>
<tr>
<th>Company</th>
<th>Trade Name</th>
<th>Capacity(t/Y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sumitomo Chem</td>
<td>Sumikasuper LCP</td>
<td>6,000</td>
</tr>
<tr>
<td>Ticona/ Polyplastics</td>
<td>Vectra</td>
<td>7,600</td>
</tr>
<tr>
<td>DuPont</td>
<td>Zenite</td>
<td>5,500</td>
</tr>
<tr>
<td>Solvay</td>
<td>Xydar</td>
<td>4,500</td>
</tr>
<tr>
<td>Toray</td>
<td>Siveras</td>
<td>1,000</td>
</tr>
<tr>
<td>Others</td>
<td>(Ueno LCP, Rodrun, etc.)</td>
<td>1,000</td>
</tr>
<tr>
<td><strong>World Total</strong></td>
<td></td>
<td><strong>25,000 - 26,000</strong></td>
</tr>
</tbody>
</table>

For your reference, the portfolio line-up of Sumikasuper LCPs is shown in Fig.1-22. There are 3 or 4 different base resins (from lower side, E6000HF < E6000 < E4000 < E5000). Lower temperature grades such as E6000HF, E6000 and E4000 must be moldable by using a conventional injection machine having heating ability up to 400°C, however, the most highest temperature grade, E5000 needs the more higher ability for heating over 400°C. In general, such injection machine must be special specification during ordering.

Table 1-7 Temperature range of each trade name LCP

<table>
<thead>
<tr>
<th>SUMIKASUPER LCP</th>
<th>Xydar</th>
<th>Vectra</th>
<th>Zenite</th>
<th>Siveras</th>
<th>Rodrun</th>
<th>Novaccurate</th>
<th>Ueno LCP</th>
</tr>
</thead>
<tbody>
<tr>
<td>100 °C</td>
<td>200 °C</td>
<td>300 °C</td>
<td>400 °C</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In addition, each base resin has several compounding grades with special formulations of fillers. Each formulation is optimized for its special usage or application. We would like to reveal the relationship between application and suitable compounds in the next section.

1-7. Conclusion

LCP has the following merits;
1 Introduction of liquid crystalline polymer

- Superior thermal resistance
  This merit brings higher soldering resistance including Pb-free soldering for electronic application.
- Shear induce molecular alignment
  This effect brings very low melt viscosity for fine & precise mold.
- Self-reinforcement effect
  This effect brings higher mechanical properties.
- High flow ability
  This effect helps to mold in thin walls.

Barrier Resistance
This effect is from the skin layer and provides probably the best gas and liquid barriers.

LCP has the following demerits;
- Strong anisotropy
  This demerit brings the difficulty to control warpage problem.
- Low weld strength
  This property brings the difficulty of designing of molded article.

At last, LCP suppliers prepare several compound grades for fitting the many kinds of applications.

Fig. 1-24 Line-ups of Sumikasuper® LCP
2 Application of LCP and its technology

Several applications of LCP have been established. The purpose of this section is to introduce the latest technology related to each application, which is necessary to understand for the best material selection. The following descriptions will disclose tips for choosing the optimized material.

2-1. Connector for PC, mobile, digital camera, etc.\(^2\)

This application field is one of the largest and the most important for LCP. Over 60% of LCP materials are used for this market. The total amount of this application of LCP in 2003 is estimated to be approximately 8000MT/yr.

Before introducing the grades for this application, we will begin by considering the Surface Mount Technology (SMT).

(1) Surface Mount Technology (SMT)

This technology comes from the integration of electronic devices. This is needed due to the limited area or volume available when minimizing electronic equipment. It is said that Japanese company, SONY has developed this technology to realize the mobile gear, “Walkman”.

For integration of electronic circuits, all devices must be mounted and soldered on the same side of the printed circuit board. This realizes at least 2 times integration of device mounting, because both side of printed circuit board can be used instead of one side use for conventional soldering technology.

Fig.2-1 shows comparison between conventional technology and SMT for soldering. For conventional soldering, devices are mounted at the certain position where the terminal holes are set. Soldering will be carried out at the opposite side of mounted device. In this case, soldering heat will be added from the opposite side of circuit board. For this reason, the devices do not demand higher soldering stability over 240°C (240°C means the soldering temperature for conventional solder).

For SMT, however, solder is printed before mounting (usually paste solder is used), and devices will be mounted at the certain position. The printed circuit board with mounted devices will introduce into the IR (Infra Red) reflow oven, and both circuit board and devices will be heated until the soldering temperature of printed solder. The soldering temperature is usually 240°C for conventional solder and 260°C for Pb-free solder. Both of these temperatures are not equal to the case of conventional soldering, because the circuit board will act as a heat seal for the devices. In SMT, the devices themselves will be heated at the same temperature as soldering.

Accordingly, the devices for SMT are called Surface Mount Devices (SMD), and high heat resistance materials are needed for such devices. This is one of the main reasons why LCP is now used for this application field.

After cooling, we can obtain the assembled circuit board just like in conventional soldering; however, we will still be able to use the opposite side of the circuit board. This technology enables to use both side of circuit boards and use of multi layer soldering.

At the same time, we should think about the dimension precision of the device during SMT soldering. The deformation of device during soldering can bring a serious misalignment problem. Fig.2-2 illustrates the situation. Soldering will only be successful in the case of the molded part keeping its flatness and plane of dimension. (Fig.2-2 a). In the case of the molded part having warpage, the deformed upper side will be misaligned. (Fig.2-2 b).

In general, the tolerance should be under 0.10mm for soldering. This means that it should be under 0.05mm after molding due to the warpage increasing by the pin-tap process. In recent years, the customer’s needs have become more precise so that tolerances after IR reflow process also should be under 0.10mm. This is not easy to accomplish by the designer, molder, and polymer scientist. The details of warpage improvement will be explained later.
A. Conventional soldering procedure

B. Surface Mount Technological (SMT) soldering procedure

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Fig. 2-1 Schematic view of comparison between conventional and SMT soldering

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Fig. 2-2 Relationship between warpage and soldering

(2) Relationship between compound formulation and warpage

In general, the higher filler content indicates lower anisotropic property, which is the most important character for reduction of warpage. However, the relationship differs between filler content and anisotropy among different hyper-dimensional structures of plastics.

In conventional materials, anisotropy increases with the filler content. Anisotropy of LCP, however, decreases with the filler content. Table 2-1 shows the relationship between GF content and anisotropy for several plastics. Here, anisotropic character is evaluated by TD/MD ratio of mold shrinkage data. And these tendencies were illustrated in Fig. 2-3.

Table 2-1 Relationship between GF content and anisotropy (Data were calculated from TD/MD ratio of mold shrinkage at 3mm, ND: no data)

<table>
<thead>
<tr>
<th>Hyper-dimensional structure</th>
<th>Material Name</th>
<th>No filler</th>
<th>GF 10%</th>
<th>GF 20%</th>
<th>GF 30%</th>
<th>GF 40%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystalline plastic</td>
<td>PEEK (Victrex® PEEK)</td>
<td>1.0</td>
<td>ND</td>
<td>ND</td>
<td>2.31</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>PPS (Novamid® PA6)</td>
<td>1.13</td>
<td>ND</td>
<td>ND</td>
<td>3.25</td>
<td>ND</td>
</tr>
<tr>
<td>Amorphous plastic</td>
<td>PES (Sumikacecel® PES)</td>
<td>1.0</td>
<td>ND</td>
<td>1.33</td>
<td>2.0</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>PC (Iupilon®)</td>
<td>1.0</td>
<td>1.67</td>
<td>4.0</td>
<td>6.0</td>
<td>6.0</td>
</tr>
<tr>
<td>Liquid crystalline polymer</td>
<td>LCP (Sumikasuper® E6000)</td>
<td>8.45</td>
<td>ND</td>
<td>ND</td>
<td>3.9</td>
<td>ND</td>
</tr>
</tbody>
</table>
It is clear that LCP has a much different relationship between GF content and anisotropic character. It is thought that LCP itself has a strong anisotropic property. Introducing of fillers disrobes such original property by disrupting of the alignment of rigid-rod molecule of LCP.

In contrast, the flowability of LCP will decline with increasing of the filler content (Fig.2-4). Accordingly, this is the reason why GF 30-40% grades of LCP are the standard formulation, because this formulation is one of the best balance between anisotropy and flowability.

With this background we can turn now to the latest design of FPC (Flexible Printed Circuit-board) connector.

We should think about the flow balance with main flow portion and side rib. In generally, conventional design has more than 0.3mm thickness at not only the main flow portion but also the rib portion. The latest designed FPC connector often has under 0.2mm thickness for both main flow portion and side ribs (comparison between a and b in Fig.2-5).

In this case, long fiber is not able to flow from main flow portion to rib portion because the length of GF is not negligible. This unbalanced flow will induce serious molding troubles such as short-shots, weld-line cracks and warpage. In this case, short GF formulation or the formulation short GF using inorganic filler together is one of the improvements. Short GF has less ability to reduce the anisotropic character than long GF; however, as shown in this case, the flow balance needs priority more than anisotropy (Fig.2-5 b).
a. Conventional design having comparably thicker wall-thickness

b. The latest design having thinner wall-thickness

Long fiber: shortage of flowability to rib portion
Short fiber and/or inorganic filler: good flow balance to rib portion

Fig.2-5 Schematic view of comparison between conventional and the latest design of connector

For the above reason, LCP suppliers offer several kinds of formulations. Table 2-2 and Fig.2-6 show the portfolio of Sumikasuper® LCP for your reference.

Table 2-2. Grade line-up of Sumikasuper® E6000HF Z series for connector application

<table>
<thead>
<tr>
<th>Grade name</th>
<th>Filler formulation</th>
<th>Features</th>
<th>Recommendable application</th>
</tr>
</thead>
<tbody>
<tr>
<td>E6007LHF Z</td>
<td>c-GF 35%</td>
<td>standard grade</td>
<td>DDR, RIMM, DIMM</td>
</tr>
<tr>
<td>E6807LHF Z</td>
<td>c-GF/mineral 35%</td>
<td>low warpage</td>
<td>S/O DDR, CPU socket</td>
</tr>
<tr>
<td>E6808LHF Z</td>
<td>c-GF/mineral 40%</td>
<td>super low warpage</td>
<td>CPU socket</td>
</tr>
<tr>
<td>E6808UHF Z</td>
<td>m-GF/mineral 40%</td>
<td>best for FPC connector</td>
<td>FPC, b/b connector</td>
</tr>
<tr>
<td>E6810LHF Z</td>
<td>c-GF/mineral 50%</td>
<td>best for Card connector</td>
<td>Memory card</td>
</tr>
</tbody>
</table>

1) c-GF: chopped glass fiber, standard GF for conventional engineering plastics
   m-GF: milled glass fiber, short size GF for special purpose

Fig.2-6 Grade line-up of Sumikasuper® E6000HF Z series
2-2. Bobbin for backlight inverter of LCD

From the middle of 2003, the FPD (Flat Panel Display) market has had tremendous growth. LCD (Liquid Crystal Display) is one of the most promising items. Lighting in the LCD requires several CCFLs (Cold Cathode Fluorescent Lamp) used according to the panel size. LCP is now being used in the bobbin of the inverter for lighting of CCFL instead of former phenolic resins.

(1) Proceeding of the backlight system of LCD

The principle of perceiving of the image on LCD is to view the contrast that is generated by the backlight through a polarizing panel and liquid crystalline cell. We will omit the detail of technology of the liquid crystalline cell here (please refer to the certain literature about this technology). LCD needs light from the outside because it doesn’t emit light itself.

The former LCDs for small calculators or early mobile phones used only natural light. Recent LCDs, however, use active backlights. In general, CCFL (Cold Cathode Fluorescent Lamp) is used for the backlights. CCFL is very similar to the home-use fluorescent lamp; however, it uses the secondary electron emission from electrode made by Ni (nickel) or Ta (tantalum) instead of a filament. The merit of the electrode system is the ability of size reduction. The diameter of a CCFL was 3mmφ several years ago; however, recent diameters are now 1.8mmφ.

The life of LCD depends on the life of CCFL. Since the current CCFL has over 50,000hr life, the LCD life equals over 50,000 hrs.

There are 2 kinds of the backlight system as shown in Fig.2-7. Sidelight systems (Fig.2-7a) are used for small size LCD as mobile phone, PDA, or PC having less than 14’ size LCD, etc. The light for imaging is settled at the side portion of LCD, and the light is guided through light tube. Recently, white LED has begun to be used for these small size LCD applications. The number of CCFLs is 1 or 2 (in this case, CCFLs are settled at both side of light tube).

Under light systems are used for large sized LCD. In this case, several CCFLs are necessary due to unifying the brightness for all of the area of LCD.

The number of CCFLs is 4 to 6 tubes for PC having 14’ – 18’ display, and in the case of the flat TV having 27’ – 40’ reaches 16 to 20 tubes. Accordingly, since each CCFL needs an inverter for it’s lighting, the number of inverter bobbins is also increasing in proportion to expanding market of flat display.

![Fig.2-7 Schematic view of the backlight system](image)

(2) Requirement for the inverter bobbin

Fig.2-7 shows schematic view of assembling process of a transformer. Soldering process is needed to remove the overcoat of wire to establish electric contact with contact pins. The removing of the overcoat of wire is done through the thermal degradation of the overcoat material (in general, polyurethane, polyester, etc) because of the reduction of tact time.

Since this soldering temperature is 370 to 390°C, higher temperature resistance of the material is necessary. If the material thermal resistance is low, melting of the bobbin and slanting of contact pins will occur. Fig.2-8.

In addition, since all electronic parts including LCD should be small and thin, the material having both high heat
resistance and high moldability is demanded.

![Fig. 2-7 Schematic view of assembling process of transformer](image)

![Fig. 2-8 Schematic view of soldering process of transformer for LCD backlight](image)

Only a few LCP suppliers provide suitable grades for this application. Table 2-3 shows Sumikasuper® E4000 series as one of the best grades.

<table>
<thead>
<tr>
<th>Grade name</th>
<th>Filler formulation</th>
<th>Features</th>
<th>Soldering resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>E4008</td>
<td>m-GF 40%</td>
<td>standard grade for bobbin</td>
<td>370 - 390°C x &lt;3 sec.</td>
</tr>
<tr>
<td>E4006L</td>
<td>c-GF 30%</td>
<td>higher toughness</td>
<td>370 - 390°C x &lt;3 sec.</td>
</tr>
</tbody>
</table>

1) c-GF: chopped glass fiber, standard GF for conventional engineering plastics  
   m-GF: milled glass fiber, short size GF for special purpose
2-3. OPU (Optical Pick-Up) actuator bobbin for CD-ROM, DVD, etc.  

OPU (Optical Pick-Up) is the electronic part that enables the reading and writing of digital data, which is assembled in the CD-R or DVD media drive unit. Recent developments realized high-speed responses through use of LCP in the actuator bobbin for laser focusing.

(1) Requirement for OPU actuator bobbin

Fig. 2-9 shows schematic view of principle of OPU actuator bobbin. In order to focus the laser ray to the micron size pits on the optical disc medium that is rotating at high speeds, the lens actuator must be driven to follow the high resolution. This is especially true in the case where the resonance at the drive speed of the actuator induces reading & writing errors of the digital signal. Recent trends and requests are keeping the resonance frequency higher than the driving speed (frequency) of actuator. However, it is not so easy due to the recent 40 times speed CD-ROM and 10 times speed DVD-ROM demanding approximately 20kHz of resonance frequency. It is expected that the resonance frequency of the material should be improved to realize this specification.

In generally, resonance frequency, \( f_0 \), is described as follows;

\[
f_0 = \frac{1}{2\pi} \sqrt{\frac{k}{m}}
\]

Here, \( k \): comparative modulus, \( m \): specific gravity.

Accordingly, it is necessary that the comparative modulus (stiffness) of material must be higher and the specific gravity must be lower, in order to increase the \( f_0 \).

In addition, the metal terminal-less (resin terminal) bobbin system is proposed due to weight reduction. In this system, shown in Fig. 2-10, the conventional metal terminal part is switched to the terminal that is formed by the molded plastic. This system has merit to reduce the bobbin weight and emphasizes \( f_0 \) (see formula (1)). However, in this case, the plastic portion will be immersed in the soldering bath with much higher temperature than the above-mentioned OPU bobbin. The soldering resistance reaches to from 400 to 420 °C.

Due to the above reason, higher thermal resistance material is necessary for this application such as Sumikasuper 5000 series LCP resins.

Fig. 2-10 Schematic view of conventional and metal terminal-less bobbin structure
(2) Relationship between modulus and thermal resistance

As with conventional plastics, higher filler content of LCP will bring higher stiffness. However it will also bring the increase of specific gravity at the same time. Accordingly, the balance between filler content, higher stiffness, and lower specific gravity should be considered.

In addition, since LCP has the characteristic that thinner wall thickness will bring higher modulus, the thinner actuator will have merit of the improved design for higher resonance frequency (see section 1-4, Fig.1-15).

A few LCP suppliers provide some suitable grades having higher soldering resistance and higher comparative modulus with lower specific gravity. Table 2-4 shows Sumikasuper® E5000 series.

As show in Fig.2-11, in spite of Sumikasuper® E5006L having only 30% GF, it indicates approximately similar stiffness with E5008L which includes 40% GF. It means E5006L is one of the most suitable grade for this application.

Table 2-4 Grade line-up of Sumikasuper® E5000 series for OPU actuator bobbin application

<table>
<thead>
<tr>
<th>Grade name</th>
<th>Filler content1)</th>
<th>Specific gravity</th>
<th>Flexural modulus @0.5mmt (MD)</th>
<th>400°C dipping in solder bath</th>
<th>Moldability</th>
<th>Recommendable application</th>
</tr>
</thead>
<tbody>
<tr>
<td>E5006L</td>
<td>c-GF 30%</td>
<td>1.60</td>
<td>30 GPa</td>
<td>OK</td>
<td>Low</td>
<td>-X40 CD-ROM -X16 DVD-ROM DVD-Video CD-R/RW OM ,etc.</td>
</tr>
<tr>
<td>E5008L</td>
<td>c-GF 40% m-GF 40%</td>
<td>1.69</td>
<td>30 GPa</td>
<td>OK</td>
<td>High</td>
<td>CD-DA low price CD-ROM OM ,etc.</td>
</tr>
</tbody>
</table>

1) c-GF: chopped glass fiber, standard GF for conventional engineering plastics
m-GF: milled glass fiber, short size GF for special purpose

Fig. 2-11 Thickness dependence of flexural modulus
3 Injection molding technology for LCP

In recent years, there is strong demand for materials that can be used for lighter, smaller, and thinner products. However, thin-wall portions less than 0.2mm are difficult to mold even if improved LCP grades are used. In these cases, the thin-wall fluidity depends on the characteristic of injection molding machines. The purpose of this section is to investigate the relationship between thin-wall fluidity of LCP and the characteristic of injection molding machines.

3-1. Control System of Injection Process

Before investigating the relationship of injection machine, we will begin by summarizing the principle of the control system of injection machine. The condition of the injection process can greatly affect the quality of molding products. There are two control systems - "open loop control" and "closed loop control".

(1) Open loop control

![Diagram of open-loop control]

Open loop control is a non-feedback control system. Since this system has comparably lower cost and simple structure, many kinds of injection machines are used for a wide variety of plastic molding.

Open loop control, however, realizes lower reliability for the velocity (injection speed) due to it not feeding the actual velocity and pressure to the controlling unit. Injection speed is usually affected by the flow resistance induced by the material viscosity and tool cavity design (ex. gate, narrow portions of cavities, etc). Other factors such as the oil temperature and the viscosity of the molding material could also influence the real condition of the injection, and thus they might be often different from the set value.

Important setting parameters are as follows:

- Pressure setting: this means both injection pressure and holding pressure
- Velocity setting: “hydraulic flow” for hydraulic system vs. velocity.

Usually, this kind of machine uses lower response parts such as switching valves for hydraulic systems. It sometimes causes the generation of excess shock pressure.
(2) Closed loop control

The closed loop control is so-called “servo control system” which has a feed back loop circuit of control signals and comparison systems between setting value and actual value. This system not only indicates detected value of pressure or velocity, but also the feed to comparison systems to keep the actual value with setting value. Accordingly, the velocity controllability is more reliable than open loop control system.
However, this system is more expensive due to all mechanical parts requiring quicker and more accurate response. This system is more suitable to realize accurate molded parts. Important setting parameters are as follows:

Velocity setting: Required pressure is determined by CNC itself.

V-P switching position: This is one of the most important settings to determine the point from velocity priority control to pressure priority control. It affects to end position of injection and shock pressure.

Limiting pressure setting: Some machines don’t have this parameter. It limits the maximum pressure during velocity priority control (injection process). Basically, this parameter has no relationship with velocity; however, to low of a setting will affect controllability of velocity.

---

3-2. Thin-wall fluidity of LCP and characteristic of injection molding machine

On the basis of the principle above mentioned, we have carried out the investigation to clear the moldability dependency with characteristic of the difference of injection machines.

(1) Experiment

The 4 cavities tooling shown in Fig. 3-5 (cavity thickness: 0.2mm) was used for this study with Sumikasuper E6008 (GF 40%) as LCP. Injection molding machines were used shown in Table 3-1 and mold temperature was 130 °C.

The molding temperature of each injection machine was 360 °C measured by IR (Infra-Red) emission thermometer (IT-240S: Horiba Ltd., Japan) at 0.86 of the emission rate before each experiment. Details of measuring of the molding temperature are shown in Fig.3-6 (see the same manner of the description at 5-4-(2)-c).
3 Injection molding technology for LCP

Fig. 3-5  4 cavities model tooling

Table 3-1 the list of injection molding machines of this study

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>control system</td>
<td>open-loop</td>
<td>open-loop</td>
<td>closed-loop</td>
</tr>
<tr>
<td>clamping force</td>
<td>40T</td>
<td>30T</td>
<td>45T</td>
</tr>
<tr>
<td>cylinder diameter</td>
<td>25 mmφ</td>
<td>22 mmφ</td>
<td>22 mmφ</td>
</tr>
<tr>
<td>injection rate / injection pressure (cm³/s)/(kg/cm²)</td>
<td>1st 360 / 467</td>
<td>1st 99 / 770 (1)</td>
<td>151 / 2790</td>
</tr>
<tr>
<td></td>
<td>2nd 176 / 956</td>
<td>2nd 134 / 1312</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3rd 118 / 1431</td>
<td>3rd 195 / 896</td>
<td></td>
</tr>
<tr>
<td></td>
<td>4th 88 / 1920</td>
<td>5th 70 / 2388</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5th 70 / 2388</td>
<td>6th 58 / 2579 (1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>6th 58 / 2579 (1)</td>
<td>70 / 2388</td>
<td></td>
</tr>
</tbody>
</table>

1) for machine A, 6th mode is the standard position
2) for machine B, 1st mode is the standard position

Fig. 3-6 Schematic view of measurement of resin temperature with IR emission thermometer

Actual waves of injection speed and pressure were recorded for each injection condition and the flow length of
0.2mm cavity was measured at the same time. The schematic wave is shown in Fig. 3-7.

![Fig. 3-7 Schematic wave of this study](image)

(2) Results

Fig. 3-8 shows the relation between injection time and flow length. In the case of open-loop injection machines (A or B), flow lengths developed a tendency to increase with decreasing of injection time, however, injection time has not decrease under 0.10 second. In contrast, injection machine C had under 0.10 seconds of injection time easier than A and B, then flow length has greatly improved.

![Fig. 3-8 Relation between injection time and flow length](image)

The reason why these different characteristics were observed can be considered the difference of rise-up characteristic of injection machines. That is to say, compared with the rise-up characteristic of pressure, injection machine C is the most rapid vs. B and C shown in Fig. 3-9. This is due to the servo-valve of the closed-loop system showing quicker performance than the proportional controlled valve of open-loop system. It is clear when comparing the velocity characteristics shown in Fig. 3-10.
Fig. 3-11 shows the relation between maximum velocity and the flow length for various injection molding machines including A, B, and C. Closed-loop machines show better moldability than open-loop machines. It is reasonable to suppose that the closed-loop injection machines are profitable for thin-wall moldings using LCPs.
Fig. 3-11 Relation between maximum velocity and flow length for various injection molding machines
3-3. Flow hesitation

(1) What is “Flow Hesitation”?

We often face the strange problem as shown Fig.3-12. It is expected that the weld line is found at the opposite side of the gate. The flow end portion, however, is at the near side of the gate. This causes short-shot problems in this area. In normal situations, the short-shot problem should occur in the same portion with the weld lines, and the flow end portion should be the same.

Another case of flow hesitation can be observed in multi-cavity molds having sprue & runner structures drawn in Fig.3-13. In this case, the short-shot problem often occurs at near side of the sprue, and flash occurs at far side from sprue. It is usually impossible to improve those problems by optimizing the molding conditions.

(2) Flow behavior of LCP

Before considering the flow hesitation of LCP, it will be useful to discuss the flow behavior of conventional plastic. Fig.3-14 shows the relationship of pressure loss and flow length against the distance from gate using the model of example #2 of Fig.3-13.

As seen in the figure, inner pressure of material at the near side of the sprue is larger than far side, since the pressure loss will enlarge at far side. As the result, near side cavity of sprue (cav.1 and 2) indicate longer flow length and it means that those cavities have the tendency to indicate flash problem, since those are higher inner pressure than far side ones. At the same time, far side from sprue indicates shorter flow length and short-shot tendency.

In this case, if the flash problem at near side of sprue is not so serious, increasing the injection pressure or the same manner of condition optimizing is one of the best ways to improve unbalance flow length for these 5 cavities in Fig.3-15.

On the other hand, LCP indicates small dependence of pressure between near side and far side of sprue, since LCP has very lower melt viscosity Fig.3-15). In this case, until the space is filled with LCP at the runner portion, the LCP will not flow into each cavity through the narrow clearance of gate.

At the same time, however, if the pressure at the gate is not high enough there will be insufficient flow into the cavity through the narrow clearance of the gate. Since LCP has a high speed solidifying property, it is quickly cooled.
by the mold. For these reasons, plugging at the gate is generated especially at the near side of the sprue.

In addition, LCP flow does not reach the far side of the sprue on the first stage of material flow. Accordingly, LCP is rather easy to flow at far side of the sprue even if the plugging generated at the near side of sprue.

This situation is described as shown in Fig. 3-16. In fact, the inner pressure could not increase at the far side from the sprue. The flow lengths to the cavities are observed per the figure. The reflection of LCP flow at the end portion of the runner induce the opposite direction flow (upper part of Fig. 3-16).
This phenomenon is called “Flow Hesitation”, and it is often observed not only in large items but also small and thin portions of accurate designed items.

Once this phenomenon has occurred, no matter how the molding condition will be optimized, there is no way to improve the molding problem induced by this phenomenon. The most effective way to improve it is the modifying the flow pattern by moving the gate position, thickness of the cavity, or installing the core-out (depression), etc. to unify the flow pattern.

In general, this phenomenon is observed as short-shot, weld crack, or sticking to the mold. Since these problems seem no relationship to flow hesitation, they can be misdiagnosed. It is desirable to consult to such experts if your products have these kinds of molding problems.
3-4. Metering of LCP

Stable & short-term metering is an essential function of injection molding technology for plastic molding. In-line screw mechanism is a very sophisticated mechanism due to having both plasticizing and injecting functions of the plastic.

We often suffer unstable metering during LCP molding. This trouble is usually found via the unstableness of metering time and due to clumsy forwarding of LCP granules at the hopper or the pulsation spout of melt during purging.

We have to say that the unstableness of metering is one of the essential problems of LCP molding.

In recent years, LCP manufacturers have improved this phenomenon with their own technologies; however, it has not always been successful.

(1) Principle of unstable metering

We must think about the melting process of LCP during the metering of the injection machine. The heat in order to melt the LCP granule is only supplied from cylinder (or nozzle) heater. Since the heaters are installed outside of cylinder, the heat always comes from outer portion of cylinder shown in Fig. 3-17. Due to this reason, there is temperature deviation at small space between screw and cylinder.

Accordingly, the first step of melt of the granule starts from the side that is in contact with cylinder wall. If the melt process starts, the hardness of granule will be quickly lost, and melted portion will permeate into the every gap of un-melted granules. If such melt portion reaches to lower temperature region at screw side, it will solidify again and paste to the granules.

Since the melt of LCP has significantly lower melt viscosity than conventional plastic melt, it will easily climb over the screw edge through very small clearance between screw edge top and cylinder wall (Fig. 3-18). This means that back flow easily occurs for LCP.

If this pasted portion is generated, it will rotate with the screw and lose its transportation force of materials from hopper side to nozzle side. Eventually, this pasted portion will prevent and plug the transportation of the material.

This plug is usually generated at the portion where the heat supply arrives to the critical value for melting of the granule. This is especially at the compression zone of screw because heat comes not only from the outside heater but also inside by heat insulation compression.

Once the plug is generated, the latter material is not supplied from the hopper side due to the plug, although the melt material at the nozzle side from the plug will spend to the molding. Then the melt material at the nozzle side is lost gradually, and becomes hollow. This situation is called as the starvation state. It continues until the cancellation of the plug (Fig. 3-19). During this phenomenon, it observes that the granule at the inlet from hopper is slipping on the screw surface. The molding engineer often explains that “the granule of the LCP is slipping, so it is not supplied to the metering zone”.

This phenomenon can be confirmed by using the visible cylinder equipment or by de-assembling of the screw from the cylinder.
The metering time elongates during the starvation state, and it is cancelled due to the cancellation of the plug. Accordingly, the metering time usually indicates cyclical deviation in this case. Fig.3-20 shows typical deviation of the metering time for this phenomenon.

The cycle of the deviation or the period until cancellation of the plug depends on the situation or the mechanical reason caused by the screw design, screw diameter, clearance change, etc.

The schematic draw of this phenomenon for LCP should be as follows;
1) The granule transferred from hopper is pressed at the compression zone.
2) The melt material flows from nozzle side beyond the clearance between screw and cylinder due to its extraordinary low melt viscosity.
3) The melt material arrives to the compression zone, and infiltrates to the deep portion of screw groove.
4) Since the temperature at the deep portion of the screw groove is relatively lower than cylinder surface, the melt material infiltrated there is re-congealed and coagulates the un-melted granule. At last, it becomes the plug.
   The softening of the granule surface due to the temperature will promote this effect.
   It also means that the metering time elongates. Sometimes it reaches over 60 seconds.
5) The plug exists until the plug starts melting and broken by the pressure of transferred granule from hopper.
   Suddenly, the metering time decreases.
6) Repetition of above mentioned

![Graph showing typical deviation of metering time during unstable metering state](image)

**Fig. 3-20 Typical deviation of metering time during unstable metering state**

### (2) Solution of unstable metering

First of all, we should notice that the abrasion of the cylinder and screw parts causes the problem.

The abrasion of screw parts, especially the check-ring, is the most important. This will enhance the back-flow phenomenon during metering process due to enlargement of clearance between the screw part and the cylinder wall. It reduces the plug-ability of reverse direction flow of the melt. If the unstable metering occurs, we recommend replacing the screw parts.

The second portion is the inlet of hopper side. The abrasion of this portion causes excess supply of the granule to the compression zone of screw. Finally, such excess granule will adhere each other by generated pressure of its own compression. If you find a remarkable abrasion at this area, we recommend you to consider replacing of the cylinder part.

The third issue is the temperature setting of the cylinder. The nozzle and front heater should set at the most suitable molding temperature of LCP that is used, but the center heater must be set lower than 20°C than the molding temperature. Also, the hopper side heater should set lower than 40°C.

The reason of above recommendation is due to the suppression of excess melt of LCP before the compression zone of the screw. In general, H2 is at the compression zone of screw. If this portion set at 20°C lower than melting point of LCP, the melting and generation of melt film of LCP are suppressed, it is worthwhile preventing the generation of plug phenomenon. In this meaning, the hopper side heater also set at much lower than its melting point.

If the metering volume is larger than 60% of full scale of metering, we must be careful as lower settings of the center and hopper side heaters sometimes causes the insufficiency of melting of LCP. We must be careful about the balance between excess and insufficiency of melting of the material.
The final variable related to this problem is the screw design. It is not an easy question due to the screw being installed inside of hard and strong steel cylinder.

However, we can recommend several suggestions as the most suitable screw design for LCP molding based on our long-term experiences (see Fig.3-22).

We believe that the following recommendations have often succeeded. The differences are not only from basic differences of design of screw at each manufacturer of injection molding machine but also an individual difference with each product of injection machine.

Probably we need much more study about this region with injection machine manufacturers.

<Recommendable screw design>
Compression ratio: 2.0 – 2.2
L/D : 20 - 25

Fig.3-22 Recommendation of screw design
4 Improvement of the warpage

As mentioned, LCPs have excellent performance in many areas as compared with the other engineering plastics. Especially their excellent fluidity that is useful in the latest electronics parts having very thin-walls with complicated state-of-the-art 3-Dimensional designs.

However, one of the most common problems is warpage, which is often observed during the development of the part that requires accurate flatness. The reason of this problem is due to the progress of the Surface-Mount-Technology (SMT) for soldering and assembling.

In general, the conventional technology of soldering requires installing the terminals of the parts into the certain position of holes at the Printed-Circuit-Board (PCB). Soldering is then carried out at the opposite side of the PCB. In SMT, soldering is done at the same side on the PCB where the parts are assembled at the certain position.

In this case, the parts designed for SMT (usually known as a Surface-Mount Device: SMD) requires superior flatness and coplanarity as miss-soldering can occur if there is warpage or twist (Fig. 4-1). In general, the requirement of flatness is under 0.05mm after molding; however, it is not so easy to realize even if LCPs exhibit superior low shrinkage property.

In this section, we would like to show the principles of the reduction of the warpage for molded parts by LCPs.

![Fig. 4-1 Surface-Mounting Device with/without warpage during soldering](image)

**4-1. Theoretical background of the warpage problem**

In general, warpage of a molded part is caused by several reasons. For example, a) volume shrinkage, b) retention stress, and c) orientation of filler. However, we should point out that LCP itself is a strong anisotropic material.

According to our experiences and experiments, we found that above a to c have a relatively small affect on warpage of LCP molded parts. We have found that the direction of flow (in other words, it is a direction of orientation of LCP rigid-rod molecule) has the highest affect.

Generally speaking, melt polymers will prefer to flow into the thicker portion of the part more than the thinner portion. The reason is the pressure-loss of the thicker portion is lower than thinner portion. LCP is no exception to this physical rule. Since LCP indicates much higher shear thinning of viscosity, this tendency is much stronger than other kinds of plastics.

To take an example that has both thicker and thinner portions, as drawn in Fig.4-2, the polymer will first flow into bottom portion followed by its flow into the upper rib portion. This is due to the bottom thickness (b) is larger than the rib thickness (a)). In this case, the flow pattern of the material can be explained equivalently as the arrows drawn in Fig.4-2 a.

Since LCP has much smaller shrinkage for MD (Mechanical Direction) than TD (Transverse Direction), the bottom (the flow of LCP is MD) will keep its dimension but the rib (the flow of LCP is TD) will shrink. As a result, warpage expressed in Fig. 1-1b will probably become a reality.

In this theory, it is not necessary to think about the volume shrinkage in terms of thickness. This is different then conventional plastics as explained above. According to our investigation and experiences, almost all cases can be expressed by this flow pattern. That is, it is reasonable to think that the warpage of LCP molding is controllable by
fitting to more suitable flow patterns.

Three possibilities are recommendable to fit the flow pattern:
- First of all, thickness equalization of all portions should be considered. This is the first step and the most important. In the case of the part having both thick and thin portions, the reduction of warpage is usually difficult. Thicker portion should be thinner, and thinner portion should be thicker. This procedure is sometimes difficult to carry out due to restrictions such as product design or function. However, it should be noted that 0.01mm modification of the thickness sometimes could modify the flow pattern. The reason of this effect is that the latest designed part generally has under 0.5mm thickness and sometimes under 0.3mm. In these cases, 0.01mm difference is not negligible. We recommend that 0.01mm modification should be tested (usually, this kind of modification seems as “hair-line”, but it sometimes indicates noticeable improvement of warpage).
- The second recommendation is the installation of “core-outs” at the thicker portion (see Fig. 4-3a). This procedure is very similar to the conventional way of installing core-outs to improve not only warpage but also “subsidence”. However, the purpose of installation for LCP moldings is restriction and equalization of the material flow in thicker portions. If installing portions of the core-out is not correct, the warpage can get worse (Fig. 4-3c). For the equalization of flow pattern, installing the core-outs at the bottom will be suitable as shown in Fig. 2-3a.
- The third and last method is the moving of the gate position to the thinner portion (Fig. 4-3b). The purpose of this procedure is to make the material flow to thin portion earlier (in this figure, moving to rib portion). This procedure can be used not only for moving the gate itself but also moving the junction of portion to portion.

In the next section we will examine one of these procedures using the model mold.
4-2. Relationship between warpage and flow pattern

As described above, the anisotropy caused by flow pattern is the main reason of warpage in an LCP molded part. Evidence of this phenomenon can be seen following experiment.

Fig.4-4 shows our model mold, which has a thinner wall section at the contact portion and a thicker wall section at the bottom portion with one side gate. The material is introduced from the gate and flows into the base portion and the contacting portion.

The dependency characterization between warpage and injection speed was investigated using ultra-high speed injection machine, UH-1000 manufactured by Nissei Plastic, under following molding conditions.

- Molding temperature: 360°C (determined by IR emission thermometer, see section 3-2)
- V-P switching pressure: 39.2 MPa
- Holding pressure & time: 19.6 MPa for 5sec

Injection speed conditions were changed within the range from 50 to 800 mm/sec.

After molding, the warpage of molded connector was measured on the flat board as described in Fig. 4-5. Since the gap between molded part and the surface of flat board was not so large, a multipurpose projection microscope was used.

The result is shown in Fig. 4-6. According to the increase of injection speed, the warpage was quickly decreased for the slow speed region up to 100 mm/sec and gradually decreased for the faster speed region over 200 mm/sec.

This result indicates that higher injection speed must be applicable to improve the warpage and it should be more than 100mm/sec.
After this, the anisotropy at the surface of molded part was evaluated by measuring of the infra-red dichloic measurement technique. Fig.4-7 shows the infra-red spectra with using polarizer directed to both MD (Molding Direction) and TD (Transverse Direction) of the molded part by using FT-IR equipment, type FTS-40 with microscopy unit manufactured by DIGILAB. The 1500 and 1600cm⁻¹ bands are ascribed to aromatic ring vibration and the 1735cm⁻¹ band is ascribed to the stretching of ester carbonyl group.

Since the surface of LCP molded part is oriented to the MD direction, the transition moment of the benzene ring must be coincident with the direction of the molecular axis, which indicates the flow direction of LCP at the measured portion.

In this case, the dichloic absorbance ratio, D, and the secondary orientation factor, f, forms the following relationship:

\[
f = \frac{(D - 1)}{(D + 2)}
\]

\[\text{here } D = \frac{A_{//}}{A_{\perp}}, A_{//} \text{ is absorbance of MD, } A_{\perp} \text{ is absorbance of TD}\]

The result is shown in Fig. 4-8. The result of ’f’ at each measured portion was described by vector with its angle and relative length.

The result is shown in Fig. 4-8. The result of ’f’ at each measured portion was described by vector with its angle and relative length.
When using a slower injection speed of 50mm/sec, LCP flows to the longitude direction at the thicker portion and flows to thinner portion from thicker portion. The equivalent flow pattern (light red) is quite similar as Fig.4-2. Higher and higher injection speed, the flow direction of thinner portion inclined to the longitude direction and became very close to the thicker portion at the bottom. This means that equivalent flow patterns will change and equalize the difference of anisotropy between thinner and thicker portions.

Since increasing the injection speed reduced warpage, it is clear that flow pattern changes can enable improvement in the warpage.

As seen in this experiment, equivalent flow pattern is the most important factor in finding reasons for warpage.

![Diagram](image_url)

**Fig. 4-8** Result of IR dichloic spectra study and equivalent flow patterns
4-3. Relationship between the depth of core-out and the warpage

We have also investigated the effect on warpage of the depth of core-outs. Fig. 4-9 shows a modified model mold for this investigation. The core-outs were installed as shown and their depth could be modified from 0mm to 0.8mm (1/2 of bottom thickness). Results are shown in Fig. 4-10.

The warpage was reduced with increasing the core-out depth. This was observed for all grades of LCP tested. It is expected that this is due to the changes in the flow pattern. This was confirmed by observation of short-shot samples seen in Fig. 4-11. The difference of the flow length between thick portion (bottom) and thin portion (top side rib) became smaller and smaller with the core-out depth changed deeper and deeper. This means that the main flow at the thick portion (bottom) was inhibited by the core-outs, and the flow pattern was equalized gradually.

So far we have outlined the relationship between flow pattern and warpage using our experimental results. It seems that the observation of short-shot molding is one of the most useful and important step to evaluate the flow pattern.

Of course, recent computer simulation technology progresses very quickly and we expect that prominent software will enable more accurate evaluation in near future.

However, we cannot yet recommend to only believe in the simulation results without further empirical test since the accuracy of simulation software is not high enough yet. We should still pay attention to short-shot moldings. These must bring us very important information about flow pattern and the cause of warpage.
Fig. 4-11 Photo of short-shot samples
4-4. Case study

In this section, we would like to show you actual examples of warpage improvement for several connectors. For proprietary reasons, we have changed the design detail in these examples so that they do not reflect any actual parts.

a: Board to board connector (0.5mm pitch)

Fig. 4-12 shows schematic view of 0.5mm pitch board to board (b/b) connector. This connector had a warpage of about 0.10mm at original design. From the observation of short-shot molding, the flow pattern was thought as shown in the figure.

In this case, it is thought that the main cause of the warpage is the flow difference of top portion and bottom portion in the side wall. That is, the top portion indicates MD flow properties vs the bottom portion indicating TD flow properties.

To equalize the flow pattern between top and bottom portions, depressions were installed at top portion of side walls as seen in Fig. 4-13 (the reason of this installation is limitation of top portion flow). The warpage was improved with a reading under 0.05mm.

![Fig. 4-12 Schematic view of b/b connector](image1)

![Fig. 4-13 Installing of depressions](image2)

b: S/O DIMM

Fig. 4-14 shows S/O DIMM that was designed having average thickness as 0.03mm. In this case, 3 types of warpage were observed.

- mode 1: warpage of longitudinal direction
- mode 2: twisting
- mode 3: arm tumbling to the inside

![Fig. 4-14 Schematic view of S/O DIMM](image3)
i) Mode 1 (warping) and Mode 2 (twisting)

We started from the observation and evaluation of short-shot moldings. As the result, it was found that the flow patterns between top and bottom were not uniform - the top flow was slower than bottom flow (Fig. 4-15 “mode 1”). The reason of “mode 1” was thought to be that the thickness of the top was thinner than the bottom (Fig. 4-15 down left). In this case, the flow pattern at the arm (opposite side of the gate) was thought as shown in Fig. 4-15 (down right: arrows). Thus, the arm (opposite side) deformed to the top direction, and this molding indicated mode 2 warpage.

We recommended the equalization of the thickness at top and bottom portion. After that, the warpage of mode 1 and 2 were improved less than 0.10mm (see Fig. 4-16).

![Fig. 4-15 Flow pattern analysis 1](image1)

![Fig. 4-16 Idea for solution 1](image2)

ii) Mode 3 (crooking)

The molding still had warpage of 6~7/100. The reason for this was due to the unbalance flow of both top and bottom planes. Fig. 4-17 shows the flow pattern at top portion. We recommended installing a protrusion portion at the front side of both top and bottom planes (see Fig. 4-18). With this, the warpage was completely reduced.
c: CPU socket

Fig. 4-19 shows the conventional 2 gates system for CPU socket. In this case, the warpage will be unavoidable, especially at the gate side due to the inequality of the flow pattern at each side as seen in the figure. The 4 gate system from the center portion shown in Fig.4-20 is the one of the best ways to solve the issue. This system, of course, still has a chance of warpage at the grip portion (see Fig.4-21 a); however, it can be improved by installing suitable depressions (as described in above sections (see Fig.4-21 b)).
Liquid Crystalline Polymer

hardly shrinkable

Fig. 4-20  The 4 gates system from center portion

Fig. 4-21  Idea for improving the warpage for CPU socket
As mentioned, Liquid Crystalline Polymer (LCP) has different properties in its molten state during molding as well as the finished part. This often makes it difficult to solve several technical problems involving molding. This difficulty is due to less systematization of the polymer processing technology. Many molding engineers depend upon traditional experiences, which are often not based on chemical and rheological backgrounds. In the case of conventional plastics, this is not an issue as the rheological properties for each polymer depend upon similar theoretical characters even if the plastics having the different chemical structure will be used. However, the rheological properties of LCP are again very different, thus the molding of LCP requires a somewhat different background or knowledge.

For example, consider the crack problem of weld-line. In general, this problem is induced by separate flow fronts from separated flow, holes, insert pins or un-uniform thickness. The text book polymer processing proposes the optimization of molding conditions including modification of mold design. By increasing of pressing force between melts, often this will have an effect of increasing of the weld-line strength. Accordingly, almost all molding engineers try to improve this problem with optimization of molding conditions, since the modification of mold design requires much time and cost.

In the case of LCP, is this manner possible? The answer is No. In general, the optimization of molding condition for LCP often has no meaning to increase weld line strength.

The reason is clearly explainable. Since the difference between melt and molten state of LCP is thermodynamically very small, the rheological property of melt state and solid state of material is quite similar. This means that the melt of LCP is nearly equal to the solid of LCP. Accordingly, the once separated flow of LCP will not stick to each other any more, because solid materials will not stick each other. For this case, the modification of mold design is necessary.

As mentioned above, the reason of almost all problems are explainable and improvable. However, the thinking should be modified based on the chemical and the rheological properties of LCP.

In this section, we would like to examine of the reasons of molding problems and try to propose the improvement with experiment or case study.

5-1. Outline for trouble shooting

When there is a molding problem, we recommend that you should gather the correct information about the situation at first. For the materials engineer or researcher the following information is the most important. If you gather the information and disclose it quickly, the problem will be improved more quickly.

The important characteristics are labeled in Table 5.1. These 6 items will be necessary when you consult about your current molding problem to the LCP manufacturer.

<table>
<thead>
<tr>
<th>a) What has happened?</th>
<th>b) What kind of part?</th>
<th>c) What kind of grade and lot No.?</th>
<th>d) When and how often has it happened?</th>
<th>e) How about cavity dependency?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of trouble &gt;&gt;&gt;</td>
<td>Drawing, sketch, or molded part is very helpful to think about the reason of trouble.</td>
<td>Of course, this is the main basic information, however, the molding engineer often forgets to inform this. The lot No. is the most helpful to chase the production history at the manufacturer’s factory. If this information is missed, it will be more difficult to understand material related issues.</td>
<td>It is also important to identify the cause and duration of the trouble, or whether the problem has happened suddenly or regularly, during production or during trial testing. Also the percentage of fault is important, as it will give the material engineer more clues to solve the problem. Higher percentage (&gt;10%) means that the problem is essential, but lower percentage (&lt;1%) means that the problem may be caused by some error.</td>
<td>Sometimes this is neglected, but it gives us clear image about the problem. There should be the description of cavity No., and the dependence of cavity No. to the</td>
</tr>
</tbody>
</table>

Table 5-1 Information list for quick improvement of molding problem
issue should be noted. The cause of trouble is sometimes induced by un-uniformity during mold manufacturing for multi-cavity molds.

f) How about molding conditions? The following settings should be recorded: cylinder temperature, mold temperature, injection speed & pressure, holding pressure & time, metering position, decompression, and V-P switching position. Also filling time during injection process, minimum cushion, metering time, and cycle time and drying conditions should be recorded also. Information about injection machine (manufacturer, catalogue number, and duration of use) is also necessary.

We also recommend preparing the attached fixed form in order not to lose the important information when molding trouble has happened. (Fig.5-1).

![Fig.5-1 Sample of “Trouble shooting form”](image)

5-2. Itemized discussion

Let us begin our discussion about trouble shooting for LCP molding by defining vocabulary about molding problems.
Table 5-2 shows the definition of molding troubles using LCP in this section. The surface impression of these words are very similar to those used for conventional plastics; however, sometimes the actual cause or meaning is quite different. Accordingly, the remedies are also somewhat different from conventional procedures.

Table 5-2 The definition of molding trouble using LCP

<table>
<thead>
<tr>
<th>Item</th>
<th>Cause</th>
<th>Remedies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black spec</td>
<td>Carbonization by burning</td>
<td>- Enforcing the purging procedure</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Optimizing the circumstance of injection molding</td>
</tr>
<tr>
<td>Blister(^1)</td>
<td>Decomposition of material (includes contamination)</td>
<td>- Specifying the target material, and removing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Optimizing the circumstance of injection molding</td>
</tr>
<tr>
<td>Bubble(^1)</td>
<td>Swallow of air</td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td>Crack</td>
<td>Weakness of weld-line</td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td>Flash</td>
<td>Excessive pressure</td>
<td>- Optimizing the molding condition in order to reduce the shock pressure</td>
</tr>
<tr>
<td>Flow mark</td>
<td>Essential property of LCP</td>
<td>(almost impossible to improve)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Increasing the mold temperature</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td>Metering</td>
<td>Plug generation at compression zone</td>
<td>- Modifying the cylinder temperature setting</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Optimizing the screw design</td>
</tr>
<tr>
<td>Short-shot</td>
<td>Flow hesitation(^3)</td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Changing to high performance type injection machine</td>
</tr>
<tr>
<td>Sticking</td>
<td>Flow hesitation(^3) and excessive pressure</td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Changing to high performance type injection machine(^3)</td>
</tr>
<tr>
<td>Warpage</td>
<td>Unbalance of flow pattern</td>
<td>- Choosing the most suitable grade</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Modifying the cavity design(^2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Changing to high performance type injection machine(^3)</td>
</tr>
</tbody>
</table>

1) Distinction between Blister and Bubble is not so clear, but the reason is different. The detail about this issue will be discussed at section 5-4.
2) This explanation doesn’t indicate the same meaning at each item. Please refer the certain explanation from next section.
3) Please also refer the section 3.
5-3. Black spec

The generation of black spec is often observed in higher thermal stability grades of LCP, for example Sumikasuper E4000 series and E5000 series. The cause of this is due to the viscosity properties of LCP and the thermal stability.

As explained at section 1-3 and 3-3, melt LCP is difficult to be processed from flow prevention structures, dead zones, etc. Examples of this are the dead space of a manifold in hot runner systems, or the valve structures of a shut-off nozzle. In these areas, the former LCP will not be pushed out or replaced, and it will remain for a long duration.

LCP has excellent thermal stability; however, this remaining portion of LCP will be damaged gradually. In time, it will become carbonized material by thermal decomposition reaction. This is one of the reasons for black specs.

Another reason is the relationship between molding temperature and decomposition temperature. As explained at section 1-2, LCP decompose at over 500°C, and this temperature is thought as the upper limit of organic materials. Higher thermal stability grades such as Sumikasuper E4000 series and E5000 series have molding temperature of 380 or 400°C. Since such higher molding temperature is very close to its decomposition temperature, this circumstance is very severe for the material, even if it has excellent thermal stability. If the material is exposed at these conditions for long time, the decomposition can not be prevented.

Accordingly, the best way to improve this problem is to remove the possibility of retention under molding temperature (please also refer the section 5-4-(2)-b and 5-4-(2)-c).

The first step of improvement is to remove the reason. In general, we search and find the flow prevention structure. In this meaning, we recommend not to use the hot runner system and shut-off valve structure of injection machine. If you wish or plan to use these, you must think about the risk of black spec.

The second step is the cleaning of inside parts of cylinder, especially nozzle and the tip parts of screw. Usually, replacing to the new parts is strongly recommended. Since the tip parts of screw must be worn out with long term using and molding with glass filled material, the clearance between inside surface of the cylinder should become bigger. This clearance difference will be also the cause of unstable metering and short-shot.

The third step is the purging. To make a habit of purging regularly is preferable to prevent the generation of black spec. The details are described at section 5-4-(2)-f.
5-4. Blister & Bubble

Blistering is a very common molding problem for LCP. This problem causes serious complaints from end customers since it appears on the surface of molded parts.

The general perception is that this problem occurs when the thermal stability of the material is insufficient or that the procedure to remove moisture is not suitable - similar to the reason of “silver streaking” in conventional plastics. Often people doubt the thermal stability of LCP and are nervous about the drying temperature setting.

Indeed there is merit to these concerns; however, with careful observation you will note that there are 2 cases. One is mainly observed just "after molding", and another is mainly observed "after soldering".

We call that the former is "bubble", and the latter is "blister", because we think both of them are induced different reason.

First of all, we would like to classify "bubble" and "blister".

(1) Comparison between "bubble" and "blister"

Table 5-3 shows the comparison between a "bubble" and a "blister". From this table, both can be easily distinguished with careful observation. This difference is based on the actual reason of each case.

The reason of the “bubble” is mainly swallow of cavity air. On the other hand, the reason of the “blister” is mainly the decomposition of material (in this case, “material” does not always mean “LCP”).

Table 5-3  Comparison between "the bubble" and "the blister" *

<table>
<thead>
<tr>
<th>Item</th>
<th>the bubble</th>
<th>the blister</th>
</tr>
</thead>
<tbody>
<tr>
<td>after molding</td>
<td>yes</td>
<td>rare</td>
</tr>
<tr>
<td>after soldering</td>
<td>rare</td>
<td>yes</td>
</tr>
<tr>
<td>arising spot</td>
<td>fixed</td>
<td>random</td>
</tr>
<tr>
<td>the decomposition of material(without regard to causes)</td>
<td>rare</td>
<td>yes</td>
</tr>
<tr>
<td>the dependence of molding condition</td>
<td></td>
<td></td>
</tr>
<tr>
<td>temperature</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>speed</td>
<td>yes</td>
<td>a little</td>
</tr>
<tr>
<td>pressure</td>
<td>a little</td>
<td>no</td>
</tr>
<tr>
<td>the dependence of tooling design</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cavity design</td>
<td>yes</td>
<td>rare</td>
</tr>
<tr>
<td>air vent</td>
<td>yes</td>
<td>a little</td>
</tr>
</tbody>
</table>

*: the order of frequency is yes > a little > rare > no

(2) Reduction of the blister at soldering

The reason of the “blister” is the gas generated by decomposition of organic material. The source of this decomposed material is thought as follows:

A. Decomposition of LCP
   A-1. Production fault of LCP
       This means that the average molecular weight or distribution may be out of specification.
       In general, this cause is lot No. dependency.
       If you cannot find out any other fault described below, you should consult to LCP manufacturer with all information that you collected. The most important information is lot No. in this case.

   A-2. Hydrolysis of LCP
       Insufficient drying is the cause of this problem.
       There is the case that the actual drying temperature differs from the setting temperature.
       You should refer following section:
       a: “Effect of moisture”

   A-3. Thermal decomposition of LCP
       The cause is longer retention time in cylinder or local over heat of cylinder.
       In general, since LCP has excellent heat stability, it is hard to decompose itself except the case of causes above mentioned.
       You should refer following sections:
b: “Retention of resin at the inside of injection machine”
c: “Temperature settings of injection machine
   ~ Difference between setting value and real resin temperature~”
d: “Relation between retention of LCP in cylinder and blister”
e: “Unbalance between cylinder size and molding volume”

B. Contamination of other material
B-1. Contamination of former material
   Insufficient purging is the cause of this problem.
   It is important to carry out the recommendable purging procedure and to use the suitable purging material.
   You should refer following section:
   f: “Purging method of Sumikasuper LCP”

B-2. Contamination of purging material.
   Sometimes the purging material also becomes the cause.
   Only recommended purging material should be used for this purpose.
   You should refer following section:
   f: “Purging method of Sumikasuper LCP”

a: Effect of moisture

LCPs have extremely low water absorption (0.02%) compared with general plastics. However, the moisture in the air
can condense on the surfaces of resin granules. This moisture can cause blistering or decomposition of the LCP by de-
hydrolisis reaction. The moisture should be removed by drying the material before molding.

   recommended drying condition: 120 to 140 °C, 4 to 24 hr.

   It is often found that the hopper dryer is not hot enough in spite of the indicator of the dryer showing high tempera-
ture (for example 130 °C). There is 2 points for the cause of this trouble;

   a) The hot air is not able to circulate because the filter of the dryer is stuffed.
      -In this case, it is sometimes found that the inside temperature of dryer hopper indicates low temperature
      (for example 40 °C).

   b) The granule retain in the air service line between hopper dryer and hopper of the injection machine, and
      those granule mix with newer granule.
      -If the granules once dried by heat are cooled, then the moisture in the air condenses on the surface of gran-
      ule again. Therefore, those cooled granule must be removed before molding.

   In general, people believe that the material will keep dry if the piping system is filled with dried air supplied by air
dryer (ex. dew point: - 40°C). However, in long piping systems, as shown in Fig. 5-2, moisture can enter from outside
through small apertures of the piping. .
Accordingly, the following is recommended if the atmosphere contains high humidity such as east and south China,
east-southern Asian countries, etc.
   -To install air condition system for drying the molding room
   -To shorten the piping system length (installing drying machine just beside the injection machine)
   -In the case of blow-up machine of material, to stock into the certain vessel filled with dried air before blowing

   In addition, as indicated by our experiment it is difficult to dry the granules of material completely in high humidity
conditions using conventional procedures. Accordingly, we recommend that the resin bag should be packed or
wrapped after taking it out from the bag.

<Test method>
The granule of Sumikasuper E6807L has been left in the stabilized humidity oven at a certain condition described be-
low. After treatment, the material was molded to test-piece by injection molding machine for evaluation of soldering
resistance.

Test Sample : E6807L
5 Trouble shooting during injection molding of LCP

Molding Temperature : 360 °C
Test-piece : 0.8 mmt mini-dumbell test piece
Humidification : 40 °C, 85% RH

Temperature and Humidity Camber: PL-1GM (Tabai Espec Co.)
(Sample was set in a humidity chamber)
Drying before molding: 80, 100, 120 °C for 3 hours.
(Sample was drying for just 3 to 3.5 hours before molding at each temperature.)
Evaluation: The blister was detected after one min. dipping into the soldering bath.

![Diagram of hopper dryer and service line of pellet]

Fig.5-2 Schematic view of hopper dryer and service line of pellet

<Result>
The result is shown in table 5-4. The molded parts indicated initial performance under the certain drying condition after 1 week; however, the soldering resistance gradually was reduced by lower drying temperature or shorter drying times (Table 5-4, 1st week).

If the material left under above high humidity condition over 1 week, the situation significantly became worse. After 3 weeks, all molded parts did not keep their initial performance with conventional drying machine and conditions (soldering resistance designated under 260°C, which is not enough performance for recent non-lead soldering with 260°C IR-reflow condition).

This result means that the material should not leave under the high humidity atmosphere before drying, even if the drying machine is installed with air dryer.

Table 5-4 Relationship between humidity and blister (+: positive, no change, -: negative, blister has generated)

<table>
<thead>
<tr>
<th>Drying condition</th>
<th>Soldering temperature (N=2, 1 min.)</th>
<th>270°C, 1 min.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1st week</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>80°C, 3hrs.</td>
<td>- - - - - - - + + + + + +</td>
<td>20 / 20</td>
</tr>
<tr>
<td>100°C, 3hrs.</td>
<td>- - - - - - + + + + + +</td>
<td>14 / 20</td>
</tr>
<tr>
<td>120°C, 3hrs.</td>
<td>- + + + + + + + + +</td>
<td>0 / 20</td>
</tr>
<tr>
<td>ref. 120°C, 8hrs.</td>
<td>+ + + + + + + + + +</td>
<td>0 / 20</td>
</tr>
<tr>
<td><strong>Drying condition</strong></td>
<td><strong>Soldering temperature (N=2, 1 min.)</strong></td>
<td><strong>270°C, 1 min.</strong></td>
</tr>
<tr>
<td><strong>2nd week</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>80°C, 3hrs.</td>
<td>- - - - - - - - - - - - - - -</td>
<td>20 / 20</td>
</tr>
<tr>
<td>100°C, 3hrs.</td>
<td>- - - - - - - + + + + + + +</td>
<td>20 / 20</td>
</tr>
<tr>
<td>120°C, 3hrs.</td>
<td>- - - - - - - - + + + + + + +</td>
<td>20 / 20</td>
</tr>
<tr>
<td><strong>Drying condition</strong></td>
<td><strong>Soldering temperature (N=2, 1 min.)</strong></td>
<td><strong>270°C, 1 min.</strong></td>
</tr>
<tr>
<td><strong>3rd week</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>120°C, 3hrs.</td>
<td>- - - - - - - - - - - - - - -</td>
<td>20 / 20</td>
</tr>
<tr>
<td>120°C, 6hrs.</td>
<td>- - - - - - - - - - - - - - -</td>
<td>20 / 20</td>
</tr>
<tr>
<td>120°C, 12hrs.</td>
<td>- - - - - - - - - - - - - - -</td>
<td>20 / 20</td>
</tr>
</tbody>
</table>
Liquid Crystalline Polymer

| 120 °C | 24hrs | - | - | - | - | - | - | + | + | + | no data |

1) 2 test pieces were used at each temperature condition. The temperature of the soldering bath was increased by 5 °C starting from 240 °C.
2) Indicated the number out of 20 pieces, which the blister was detected after 1 min. dipping into the soldering bath set up at 270 °C.
3) The 3rd week’s experiment was carried on under the drying condition at 120 °C only.

b: Retention of resin at the inside of injection machine

The apparent viscosity of LCP strongly depends on both temperature and shear rate, in addition, its dependence is stronger than general crystalline and amorphous plastics. Because the apparent viscosity becomes very low under the proper injection conditions, thin-walled parts (0.3mm or less) are easily molded.

However, long term running of molding generally causes the retention of resin at dead-spaces of injection machine and this residue causes some deteriorations, for example, the contamination of black spots, the increase of gas, and the reduction of soldering resistance temperature (including the blister).

Especially, LCP easily remains in dead-spaces of injection machine because of its low viscosity. For similar reasons, the purging of previous resin is somewhat more difficult than with general plastics.

Test method

After cleaning up a screw and cylinder, the black color grade of Sumikasuper LCP was molded up to 200 shots. After this, natural color grade was molded up to 400 shots. The screw and cylinder were then taken apart for analysis of the residuary condition of the black grade.

Test Sample: black color material Sumikasuper E6008 B
natural color material Sumikasuper E6008
Molding Temperature: 350 °C

<Result & Conclusions>

As shown in Fig.5-3, we should pay particular attention to the retention of the resin at the inside of the nozzle, the tip parts of the screw and the compression zone.

Especially the inside portion of the nozzle is the most important. The carbonized LCP formed carbon pipe was often observed after long-term production. Moreover, the diameter of the carbon pipe was usually 4 to 5mmφ in general. It is significantly narrow for the conventional plastic because of the pressure loss, but it is sufficient for LCP molding.

On the contrary, the inner diameter of the nozzle should be 4 to 5mmφ for LCP molding. If the inner diameter is relatively wide, the nozzle tip behaves one of the barriers of material flow due to extraordinary high dependency to the shear rate of melt viscosity of LCP.

In the same meaning, the shut-off nozzle mechanism behaves the same kind of barrier of melt flow. Open-nozzle should be used (this was confirmed by comparison between open and shut-off nozzle under the cooperation of an injection machine manufacturer).

Recently, many injection machine manufacturers provide special designed nozzles for LCP molding.

Following are the recommendations;

a) The open-type nozzle should be used for Sumikasuper LCP.
   The shut-off type nozzle is not suitable (the shut-off valve and its surrounding portion causes the residue which will change to the cause of black-spots and the blister).

b) The internal diameter of the nozzle should be 4–5mmφ (the standard size: ca. 8mmφ, which is not suitable for Sumikasuper LCP).
5 Trouble shooting during injection molding of LCP

![Diagram of nozzle, cylinder, and metering zone with labels: Nozzle, Cylinder, Metering Zone, Compression Zone, Feed Zone, and Metroing Zone.]

**Fig. 5-3 Situation of retention resin at nozzle, screw, and cylinder**
(The recommendable design of nozzle is also drawn in the bottom of this figure)

---

**c: Temperature settings of injection machine**

~ Difference between setting value and real resin temperature~

One of the most important causes for various molding problems is the difference between the set temperature of the injection machine and actual resin temperature. Modern injection machines have digital indicators for the nozzle and cylinder temperatures so that the operator has confidence that the values recorded are equal to the actual resin temperature.

Even with this, the indicator value often differed from the actual resin temperature in almost machines that we measured.

**<Measurement procedure of resin temperature>**

The measurement of resin temperature is usually done by sticking the pin-type probe (thermocouple) into a round ball of purged resin. In the case of the resin having the molding temperature over 300°C; however, the temperature reduction of the resin ball by the radiation of heat is significant and not able to ignore. Thus it is difficult to measure the resin temperature precisely.

Because of the above reasons, some researchers have proposed several kinds of measurement. For example, Murata et al.\(^1\) proposed the measurement using the Infra-red (IR) emission thermometer and the seethermocouple for inside temperature of cylinder, and the supersonic measurement and the integrated thermocouple for mold temperature\(^29\), etc.

Among these measurements, IR emission measurement is one of the most convenient and accurate ways for evaluating the actual resin temperature, although this measurement has several defects. For example, the emission ratio should be adjusted not only for the difference of material, but also for the difference of its color.

Several researchers have applied to evaluate the actual temperatures of plastic molding by IR emission measurement\(^30\). We have also decided to install this measurement for evaluation of the relationship between actual resin temperature and setting temperature (see Fig.5-4).
apparatus: Infrared emission thermometer (IT-240S: Horiba Ltd.)
spot diameter: 1.2mm

The setting of emission ratio is very important for the infrared emission thermometer\textsuperscript{31}. We have ascertained that reasonable value of emission ratio for many kinds of plastics is 0.86 by measuring at the production factory of resin pellets.

resins used for ascertainment:
- PP (non-filler: natural color )
- ABS (GF filled: natural color, black)
- PES (non-filler, GF filled: natural color, white)
- PEEK (GF filled: natural color, black)
- LCP (GF filled: natural color, black, white)

<Result>
The result is shown in Table 5-5. Almost all cases show that the actual resin temperature greatly differs from the setting temperature.

Table 5-5 Result of temperature measurement

<table>
<thead>
<tr>
<th>No</th>
<th>Injection machine</th>
<th>Injection rate (cm\textsuperscript{3}/sec)</th>
<th>Temperature control system</th>
<th>Molding material</th>
<th>Setting temp. (\degree C)</th>
<th>Actual temperature (\degree C)</th>
<th>Metering\textsuperscript{1)}</th>
<th>Purging\textsuperscript{1)}</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Vertical type</td>
<td>27</td>
<td>ON-OFF type</td>
<td>PES 3601GL30 (GF 30%)</td>
<td>380</td>
<td>400 ~ 407 (+20 ~ +25)</td>
<td>400 ~ 440 (+20 ~ +60)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(A company)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>ditto</td>
<td>27</td>
<td>PID</td>
<td>ditto</td>
<td>380</td>
<td>375 ~ 380 (-5 ~ 0)</td>
<td>400</td>
<td>(+20)</td>
</tr>
<tr>
<td>3</td>
<td>Vertical type</td>
<td>89</td>
<td>ditto</td>
<td>ditto</td>
<td>370</td>
<td>395 ~ 400 (+25 ~ +30)</td>
<td>407</td>
<td>(+37)</td>
</tr>
<tr>
<td></td>
<td>(B company)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Horizontal type-1</td>
<td>114</td>
<td>ditto</td>
<td>PES 3601GL20 (GF 20%)</td>
<td>380</td>
<td>380 ~ 385 (0 ~ +5)</td>
<td>360 ~ 364 (-20 ~ -16)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(B company)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>ditto</td>
<td>114</td>
<td>ditto</td>
<td>PBT GF30%</td>
<td>270</td>
<td>275 ~ 280 (+5 ~ +10)</td>
<td>270</td>
<td>(0)</td>
</tr>
</tbody>
</table>
The reason why the actual resin temperature differs from the set temperature is thought to be the position of the probe (thermocouple) to control the nozzle heater is not suitable as shown in Fig. 5-5 (upper side). In this case, the thermocouple measures the temperature at the installed point, but there is often heat loss due to radiation at the non-covered area by the heater. Accordingly, the measured temperature decreases significantly when compared with the temperature at the heater position. Since the heater controller works to keep the “temperature” at measured point, the “actual temperature” at heater position must be higher than its “set temperature” (see Fig.5-5 upper-right).

From our examination, the thermocouple should be installed below the nozzle heater (see Fig.5-5 lower-left). In this case, the “actual temperature” is almost consistent with the “setting temperature”.

We should take care when we use the word “temperature”. As described above, sometimes the “temperature” is not consistent with the “temperature” which we would like to discuss. In general, the word of “molding temperature” should be used as the meaning of the “actual temperature” of the resin or the temperature below heater.

![Fig. 5-5 Comparison of thermocouple position in various types of injection machine](image)

**d: Relation between retention time of LCP in cylinder and blister**

As mentioned above, the deterioration of resin occurs when the actual resin temperature is higher than the most suitable one. This deterioration may cause blistering, which may lead to retention in cylinder.

It was found that there is strong dependence between blister and molding (actual resin) temperature or the retention time of resin in cylinder.

**<Test methods>**

Sample: E6807L
Molding machine: PS-40E5ASE (Nissei Plastic Industrial Co., LTD)

The transformation and blister of test pieces were observed after immersing them into the soldering bath set at 210 and 250°C each for 60 sec. The test pieces have been molded with various conditions of:
- molding temperature
- retention time

Also test pieces were molded with following conditions:
- different decompression degree
- low pressure / low speed

<Results>

The result is shown in Table 5-6. Neither transformation nor blister was observed when the thickness of the test pieces was thin (0.5mmt). Meanwhile, transformation and blister were observed at thick test piece (0.8mmt), and its tendency depended on the molding conditions.

It was ascertained that the blister occurred at soldering temperatures of 250°C when the resin was retained in cylinder for 15 minutes. Also the blister appeared in soldering at 210°C when the resin was retained and molded at 380 - 400°C. Change of decompression degree had no influence on the soldering resistance in this time.

As a result of the above test, the molding temperature and the retention time can be considered as main cause of blister. Unfortunately, the color change of the test piece is quite small even if it is molded after retaining in cylinder at 380°C (in case of 400°C, the color change can be recognized somewhat easily).

The purging procedure is inevitable when the resin is retained in the cylinder. Regarding the hot runner system, the situation will be more severe than above mentioned.

If the decompression degree is too high, the air may be taken into nozzle from nozzle touch portion. This air may be contained to moldings and the blister may occur in the soldering test.

Table 5-6 Result of retention test

<table>
<thead>
<tr>
<th>Thickness of testpiece (mm)</th>
<th>Molding condition</th>
<th>Condition after molding</th>
<th>Soldering Temp. test</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>210 °C 6 sec</td>
</tr>
<tr>
<td>0.5</td>
<td>Standard condition</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>0.8</td>
<td>Standard condition</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Retention for 5 min</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Retention for 10 min</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Retention for 15 min</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Retention for 5 min</td>
<td>Slightly colored</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Retention for 10 min</td>
<td>Slightly colored</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Retention for 15 min</td>
<td>Blistered</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Retention for 5 min</td>
<td>Blistered</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Retention for 10 min</td>
<td>Blistered</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Retention for 15 min</td>
<td>Blistered</td>
<td></td>
</tr>
<tr>
<td>□</td>
<td>Decompression 5mm</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Decompression 9mm</td>
<td></td>
<td>□</td>
</tr>
<tr>
<td>□</td>
<td>Low pressure/low speed</td>
<td></td>
<td>□</td>
</tr>
</tbody>
</table>

Standard condition : decompression degree 2mm

\( V = 50\%, P = 30\% \), cycle time = 20 sec

Low pressure/low speed : \( V = 30\%, P = 15\% \)

"O" shows that neither change in form nor blister was found.

\textbf{e: Unbalance between cylinder size and molding volume}

When using too large of an injection machine vs molding volume, the measuring length becomes too short. The retention time in cylinder becomes too long. In addition, excessive shearing power may be added to the resin at the screw providing zone or the compressing zone due to the high power of the large injection machine. In such situations, the deterioration of resin can easily occur.
Using 2 injection machines having different screw diameters described Table 5-7, test pieces were molded, and the soldering resistance measured.

Table 5-7  Result of the soldering test using 2 injection machines having different screw diameter

<table>
<thead>
<tr>
<th>Sample</th>
<th>Amount of gas (ppm)</th>
<th>Soldering resistance (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG-150 (Sumitomo)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Condition 1</td>
<td>10.8</td>
<td>280 (285: blister)</td>
</tr>
<tr>
<td>Condition 2</td>
<td>12.1</td>
<td>280 (285: blister)</td>
</tr>
<tr>
<td>PS-40E5ASE(Nissei)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal condition</td>
<td>6.8</td>
<td>&gt;300 (no blister)</td>
</tr>
</tbody>
</table>

SG-150 (150T, 40mmφ)     condition 1: N C1 C2 C3 C4 330 330 320 310 300 °C actual resin temp.: 350°C
condition 2: N C1 C2 C3 C4 360 360 350 340 330 °C actual resin temp.: 380°C
PS-40E5ASE normal condition: N C1 C2 C3 360 360 350 330 °C actual resin temp.: 350°C

<Result>

Fig. 5-6 shows the relationship between the metering for 1 shot and the max. metering range. If the metering for 1 shot is too short compared with maximum metering range, the retention time of the polymer inside the cylinder becomes too long, and it will be the cause of deterioration of polymer as mentioned above.

![Diagram](image)

Fig. 5-6  The relationship between the metering for 1 shot and the max. metering range

Since most of articles molded by LCP should be small, the size of suitable molding machine should be small. Recommendable specification of injection machine is as follows;
- clamping force of the mold: 50 - 70 Tons
- screw diameter: 24 - 27 mmφ
For more smaller products:
- clamping force of the mold: 15 Tons
- screw diameter: 17 mmφ

f: Purging method of Sumikasuper LCP

Several kinds of material are known for purging when using LCP resins. The most popular materials are PP(polypropylene), glass-filled PC, and commercially provided purging materials. According to our investigation, almost all materials indicated the ability of purging, but did not indicate the ability
of self-purging by LCP. In this case, the purging material will remain in the cylinder of injection machine and it will decompose by the heat due to the molding temperature of LCP being higher than 300°C. Such purging material doesn’t have higher thermal stability as LCP.

What needs to be emphasized is that we should notice not only purging ability of LCP by purging material, but also self-purging ability of purging material by LCP.

For example, glass-filled grade of PC is widely used as purging material. However, the amorphous polymer such as PC has comparably higher melt viscosity (it means that the purging material will not be removed by LCP because lower viscosity material has less ability to purge higher viscosity material). In addition, the melt viscosity of PC will enormously increase if it cooled under around 140°C. This increase of viscosity also makes the purging by LCP difficult. Accordingly, glass-filled PC is not suitable for purging of LCP.

We would like to now introduce our investigation regarding purging of LCP.

<Test method>

Test method is very close to the description at 5-2-2-(2)-b: i.e. after cleaning up of a screw and a cylinder, the black color grade of Sumikasuper LCP was molded up to 200 shots. Then, a purging material listed below was used for purging of black color material, and immediately the purging material was purged again by natural color material. After that, the screw and the cylinder were taken apart for analysis of the residuary condition of the black grade and purging material itself.

Test sample: black color material Sumikasuper E6008 B
natural color material Sumikasuper E6008
Purging material: listed in Table 4-8 with results
Molding Temperature: 350°C

<Result>

The results were shown in Table 5-8.

Table 5-8  Result of investigation about purging material for LCP

<table>
<thead>
<tr>
<th>Grade name</th>
<th>Manufacturer</th>
<th>Property of purging material</th>
<th>Test results</th>
<th>Total assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Base polymer</td>
<td>type</td>
<td>Decomposition temperature(°C)</td>
<td>Purging ability</td>
</tr>
<tr>
<td>HF21D</td>
<td>Idemitsu Petro</td>
<td>HDPE</td>
<td>- 494</td>
<td>++</td>
</tr>
<tr>
<td>Z clean</td>
<td>Chisso co.</td>
<td>PST</td>
<td>form</td>
<td>436</td>
</tr>
<tr>
<td>S11</td>
<td>Chisso co.</td>
<td>HDPE</td>
<td>GF 496</td>
<td>+</td>
</tr>
<tr>
<td>S29</td>
<td>Chisso co.</td>
<td>PP whisker</td>
<td>441</td>
<td>++</td>
</tr>
<tr>
<td>Amteclean</td>
<td>Matsushita</td>
<td>PP</td>
<td>whisker</td>
<td>437</td>
</tr>
<tr>
<td>Ex(AP-10)</td>
<td>Matsushita</td>
<td>PP</td>
<td>whisker</td>
<td>446</td>
</tr>
<tr>
<td>Asaclean</td>
<td>Asahi Kasei</td>
<td>PST</td>
<td>GF 430</td>
<td>++</td>
</tr>
<tr>
<td>Plasclean</td>
<td>Hoshi plastic</td>
<td>HDPE</td>
<td>form</td>
<td>495</td>
</tr>
<tr>
<td>Tycelean P</td>
<td>-</td>
<td>-</td>
<td>390</td>
<td>++</td>
</tr>
</tbody>
</table>

1) Information were adopted by published document of each manufacturer
2) Decomposition temperature was evaluated by ourselves using TGA equipment (TGA-50 of Shimadzu co.).

Thus, we recommend following listed materials as purging material for LCP (see Table 5-9). We also recommend that you should not use any other materials for this purpose, even if you have much experience to use LCP.

Table 5-9  Recommendable purging materials

<table>
<thead>
<tr>
<th>Purging Reagent Product</th>
<th>Z Clean S11</th>
<th>Amte Clean Ex</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Chisso Corp.: JAPAN)</td>
<td>(Matsushita Amtech : JAPAN)</td>
<td></td>
</tr>
</tbody>
</table>

2) HDPE (High Density PolyEthylene)
We also recommend that the purging procedure described below (see Table 5-10) should be followed. In general, since the purging materials have less thermal stability than LCP, it is not preferable to leave the injection machine filled with purging material at the higher LCP molding temperature. It is important to quickly remove the purging material when using LCP molding.

We also recommend that this kind of purging procedure should be carried out at least once a week, even if you are not switching to other plastics. The 5 to 10min investment in time is very worthwhile. Carrying out this procedure frequently reduces the generation of materials in cylinder and black spec generation.

Table 5-10 Purging process

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Temperature setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) End of molding</td>
<td>Same as molding (Ex. 360-365-330-290°C)</td>
</tr>
<tr>
<td>2) Feeding of purging material</td>
<td>Same as above</td>
</tr>
<tr>
<td>3) Decreasing setting temperature</td>
<td>Notice: Do not stop purging</td>
</tr>
<tr>
<td>4) Continuing the purging</td>
<td>Same as above</td>
</tr>
<tr>
<td>5) Increasing setting temperature</td>
<td>Same as molding (Ex. 360-365-330-290°C)</td>
</tr>
<tr>
<td>6) Resuming of molding</td>
<td>Same as above</td>
</tr>
</tbody>
</table>

- This purging method should be execute once a week preferably.
- Please stop the injection machine after the above step 4) is completed.

Starting procedure: 1) Set the temp. as same as 3).
2) As soon as reaching the molding temp., then feed the purging material in the hopper.
3) Follow the same step from 3).
(3) Settlement of Bubble problem

It is necessary to consider both the suitable molding conditions and tooling design in order to reduce the dragging of air into the cavity and possibly creation of bubbles.

a: Dragging of the air into the melt

In some cases, dragged air into the melt causes bubbles (the oxidation of polymer will occur and color change or black spots will be often observed in this case: please refer (2) "Reduction of the blister at soldering" also). The reasons of the air dragged into the melt are as follows;

i) Too low of back pressure
   LCP does not need high back pressure, however, too low back pressure causes insufficient removal of the air dragged from hopper.

ii) Too high decompression degree of screw on metering process
   The air is dragged into the melt from nozzle touch portion.

iii) Too high screw rotation on metering process (over 200 rpm)
   This causes insufficient removal of the air dragged from hopper similarly with i).

b: Suitable injection condition

As described above, dragging air into the melt will cause both the bubble and the blister. In addition to the description, the following procedure is more effective to reduce bubbles.

To remove air in the sprue and runner smoothly, the injection speed should be set lower (ca. 20~50 mm/sec) as the polymer passes through the gate. After this, the injection speed could be increased if necessary (Fig. 5-8). As described at section 3, products having thinner wall thickness (<0.2mm) will need higher injection speed. However, it is often difficult to remove air not only in the cavities but also the sprue and runner from the air vents. Usually the injection speed is very high (>100mm/sec) and the injection time is too short (ca.<0.1 sec).

c: Suitable cavity design

After improving the injection conditions, the bubble may still remain at the thinner wall of the products. The cause is thought to be air in the cavity that is dragged with the polymer flow as shown in Fig. 5-9. A similar phenomenon was observed for calcium carbonate and long glass fiber reinforced PP by Yokoi et al32. According to Yokoi et al, in the case of higher injection rate, And Tsuji et al. expanded this phenomenon to LCP theoretically by computer simulation using expanded Doi equation33. In this case, the bubble is often observed at the thinner position beside the thicker portion (Fig. 5-10).

We often recommend both installing enough numbers of air vent (see Fig. 5-11: not only cavity but also runner is more effective) and installing the edge at the corner of cavity (see Fig. 5-12).
5 Trouble shooting during injection molding of LCP

Fig. 5-8 Suitable injection condition (example)

Fig. 5-9 Schematic view of dragged air in the cavity
Fig. 5-10 Arising spot of the bubble

Fig. 5-11 Install the enough numbers of air vents

Fig. 5-12 Install the edge at the corner
5-5. Crack

The main cause of cracks in LCP parts is from crack generation at the weld lines. The reason why the weld strength of LCP is so weak is discussed in section 1-5. In general, there are only a few choices to improve this problem. For example, to shift the weld line to other sections in order to prevent excess force, or to increase the thickness and to equalize the surrounding portions of the weld line.

In addition, the generation of weld is fairly complicated and we must consider the flow pattern induced by the difference of wall thickness.

In this section, we would like to show you our approach to solve this problem using actual case studies.

(1) Cracking problem for Coil Bobbin

Coil bobbins are part of electrical inductance by winding the wire around the bobbin. During the assembly process, the coil bobbin was subjected to a strong winding force by the wire (Fig.5-13). After wiring, the problem occurred at the body of the bobbin (Fig.5-14). The crack was observed at around center of the cylinder portion horizontally. In addition, the cracked line was elongated at the rib installed inner side of the cylinder.

In this case, the outer diameter of cylinder between brims equals d, the inner diameter (a) is for vertical direction and is larger than horizontal direction (b) as seen in Fig. 5-15. It means the thickness of cylinder is not equal.

In this case, the flow pattern is as shown in Fig.5-16. The LCP flows from the gate to upper brim, then downward. Accordingly, the LCP flow downwards through thicker side-wall portions and rib portions. The melt LCP arrives at bottom brim and will reflect upward where the LCP is delayed to flow (Fig.5-16 i). That is, it generates the flow hesitation and the weld at the portion where both downward and upward flow bump.

Due to eliminating the flow hesitation, the equalization of wall-thickness is recommended. Diameter a and b should be the same. After equalization of the thickness at cylinder portion, the LCP flows more uniformly. (Fig.5-16 ii).

In addition, there is the case that the short shot occurs at the weld portion and causes lower strength. Higher holding pressure is sometimes available to improve this problem. In general, LCP requires relatively higher injection speed but lower and shorter holding pressure due to its quick solidification property. However such higher injection speed sometimes enhances the flow hesitation. For these cases, allowing relatively higher holding pressure conditions is effective after packing, under ordinary higher speed and lower pressure condition in order to push the LCP to the weld portion:

Holding pressure: 80 to 90 MPa  (generally 40 to 50 MPa)
Holding time: 1.5 to 2.0 sec.  (generally up to 1.0 sec.)
(2) Side-wall cracking for Case part

The case for electronic parts has a relatively simple design, but sometimes the weld crack is not negligible. Especially if the mechanically important section has a thinner wall and also the weld lines.

Fig.5-17 shows a typical designed case having a thinner and thicker section. If the gate is installed at the portion described in the figure, the weld line generates at the thin section. In this case, the weld crack problem cannot be avoided.

The LCP introduced from the gate flows the top portions and spreads in all directions and then flows downwards. Since only the front portion has a square hole, the flow is divided, and bumps at the bottom portion of the hole. In this
case the generation of weld is not avoidable (Fig. 5-18 i), without change of gate location.

Fig. 5-18 ii) shows one of the most preferable solutions for this problem. The LCP will flow in the same manner as the original position, but after flow downwards there is less chance to generate the weld at the weak section. Of course, the weld will generate at the opposite side-wall, but there is much less probability to break because this section is much stronger.

Incidentally, the weld line having Y character shape at the side-wall indicated in the figure is one of the most preferable to reduce the weld crack problem for case parts. It means the whole flow of LCP from top and both sides are balanced. If the weld line shape is far from Y character, you must consider modifying the part design in order to control the flow.

![Flow Pattern Diagram](image)

i) estimated flow pattern       ii) after moving the gate

Fig. 5-18 Estimated flow pattern: i) current flow pattern, ii) recommendable gate position

### (3) Weld crack at the hole of Board to Board Connector

Fig. 5-19 shows a typical housing part of a board to board connector with 0.5mm pitch contact pins. Core-outs are already installed in the base section to improve warpage.

If the crack is caused by the low weld strength, there is only way to improve it as shown in Fig. 5-20. This is to change the direction of springs and to remove the stress from the weld portion.

![Connector Diagram](image)

Bottom view

The crack trouble has occurred at this portion

Fig. 5-19 Schematic view of the board to board connector (0.5mm pitch)
However, if the only reason of the crack is the low weld strength, why does the crack occur near the gate side in this case? And this is not a rare case when once the crack problem occurs.

If you observe the flow pattern carefully (usually using the short-shot molding), you will be able to understand the flow pattern of this case as Fig. 5-21. The formation of the weld lines is not equal for the whole portion of the holes. Fig. 5-22 shows the difference of the weld portion between gate side and after center portion. Since the flow pattern between both side walls and the spine is nearly equal near the gate, the polymer will flow in the side wall and the spine. Then the weld will form at the center of thin ribs between the holes (A in Fig. 5-22). At the center portion of this connector; however, the polymer flow from the side wall will become faster than at the spine, due to the effect of the depressions (B in Fig. 5-22).

Since it is thought that the weld strength of A and B is the same, the reason why the crack occurs must be related to the existence of the weld line at the rib portion between both holes. As shown in Fig. 5-22 A, the strength of the weld section is very weak because the wall thickness is very thin. However, as shown in Fig. 5-22 B, the weld line is formed
at the spine section and the weld strength will become stronger than that in Fig. 5-22A. It is easily understood that the crack will not occur at the rib portion between both holes.

From these points, some ways to improve the situation (also see Fig. 5-23);

1) Changing the gate system from 1 point gate to 2 point gates
2) Installing the additional depression near the gate side
3) Installing the additional depression at the side wall of anti-gate side
   (to remove the back flow as seen in Fig. 7-3 upper 3)

Fig. 5-23  Schematic view of the idea to improve the crack based of its flow pattern control
5-6. Flash

Sometimes a molded part has an extra amount of material by flowing of a polymer melt into the clearance between parts in a mold. This is called “flash”\(^{35}\). In general, flash is caused by excessive injection pressure, low melt viscosity, excessive rapid injection, or inadequate mold venting.\(^{35}\) LCP usually doesn’t have this flash problem even though it does have significantly lower melt viscosity and is quicker filling into the cavity with rapid injection velocity.

Wissbrun has tried to explain this paradox in his former work\(^{36}\). He demonstrated following evidences:

- LCP has very close solidification temperature to injection molding temperature, its latent heat is almost zero, and its thermal conductivity is high.
  This means that the large thermal diffusivity with small amount of heat induces quick solidification, and its order of solidification speed is significantly larger than conventional plastics.
- The power-law exponent \(n\) is higher for LCP than that of the isotropic polymers tested by Richardson\(^{37}\). End corrections of LCP are significantly larger and those depend on both shear stress and temperature, although those of isotropic polymers depend on stress, but not very much on temperature.
  This means that the flow length of LCP at narrower sections from the wider areas is comparably shorter than that of conventional plastics, and it also means that LCP has less tendency of flashing.
- The flow length at constant pressure depends upon the power of the coefficient \(\eta_0\) of power-law equation. This value of LCP is also significantly larger than that of isotropic polymers. This higher value affects excellent moldability, but does not affect the flash tendency.

Finally he concluded that quick solidification behavior and stronger dependence of melt viscosity with shear rate of LCP relates to minimizing flash and the smaller melt viscosity of LCP induces higher moldability. This examination is completely consistent with the usual observation at molding and is quite correct explanation of LCP behavior.

However, we sometimes suffer the flash trouble for the LCP molding beyond the basic behavior of LCP as described above.

In these cases, we should notice that LCP has very low melt viscosity, and it favors not only flow to the mold cavity but also to a narrower gap such as air vent if excessive injection pressure is added. We would confirm following conditions.

1) Higher actual resin temperature
   The molding temperature should be set at moderate, which is disclosed on the technical issues provided by the LCP manufacturer. Since LCP is very sensitive to the temperature, the most suitable range is within \(+\text{−}5^\circ\text{C}\) from the designated molding temperature.
   For your reference, the measurement of actual resin temperature and related problems are explained at the section 5-2-2-(2)-(c).
2) Lower clamping force
   If the start inspection should be carried out daily, this error must be prevented. This trouble sometimes happens since the clamping force setting does not change during mass-production and by using the same mold and the same material, and no one pays attention to this.
   Re-adjustment of the clamping force is necessary if the accuracy of the parting line or mating surface is enough for accurate molding.
3) Excess injection pressure
   The “injection pressure” means not only the injection pressure setting or the holding pressure setting, but also the shock pressure at the end of injection process. In general, LCP does not need higher injection pressure or holding pressure. If these setting values are too high, changing and choosing of lower value is preferable to solve.
   In the case that these values are correct, we should adjust and reduce the shock pressure. In this case, we should look at other ways. One way is by adjusting the V-P switching point and observing the molding wave monitor. The details are described at section 3-1.
5.7. Flow mark

The surface of molded parts of LCP is not smooth and appears to be a typical pattern, which is very similar with a so-called "flow marks". It often becomes an obstruction when an LCP molded part is used for exterior parts whose appearance is regarded an important property. The reason of this phenomenon is not clear, but it may be due to the reflection of light upon the LCP molecular alignment at the surface. This difference of alignment should be induced by the flow pattern of LCP during the flow.

In general, the melt polymer flows in the manner of so-called "fountain flow" (Fig. 5-24). However, it should be observed under the ideal condition or some special circumstances.

Indeed the thin walled part indicates a comparatively smooth surface. It seems that the reason for this is from the melt polymer flowing in the fountain flow as mentioned above.

For LCP molding, such unbalanced flow is the reason of the flow mark phenomenon shown in the figure. This is due to the difference of orientation of the LCP molecules on the surface and different reflectivity of the light. In this case, the eddy portion (as indicated "dark" in the figure) should be the portion where the glass fiber appears to the surface. Again, this is due to the insufficient touching of melt polymer to mold wall by the eddy. (Because the sufficient touching to the mold wall will bring the smooth surface. This means that sufficient pressure of the melt polymer covers the fillers as Glass Fiber.)

In general, higher mold temperature is useful for improving the surface condition. Indeed, it is known that the surface of an LCP part molded with over 150°C of mold temperature can be improved. Such parts have relatively shining and clear color surface (of course the mold surface must be polished like a mirror). Black color grade indicates darker black than the case molded by the condition with much lower mold temperature. However, the higher mold temperature induces another problem, the sticking problem to the mold because of enlargement of the mold release force.

Since leveling of meandering during LCP flow is very difficult, this phenomenon is one of the essential for LCP molding. In other words, the flow mark on the surface for LCP molded article is unavoidable.
5-8. Metering

This problem is induced by the starvation phenomenon of the melt during the metering process discussed at the section 3-4.

Almost all LCP manufacturers are applying additives to their compounded granule to prevent adhesion of granules to themselves\(^{38}\). However, sometimes this is not enough to improve the starvation caused by this.

The first recommendation to eliminate this is by modifying the temperature setting of the cylinder. Since excess melting at the compression zone is one of the main causes of plugging, decreasing the set temperature at the hopper side is recommended. In general, the nozzle and front section of the cylinder should be the same temperature as the molding temperature. The middle section of cylinder should be 20 to 30 degrees less from the molding temperature, and the hopper side section should be at least 60 degrees from the molding temperature.

Ex.1) in case of the standard molding temperature as 350°C
   For Sumikasuper E6807LHF is as follows;
   Nozzle: 360°C, Front: 360°C, Middle: 340°C, Hopper side: 300°C

Ex.2) in case of the standard molding temperature as 380°C
   For Sumikasuper E4006L is as follows;

Ex.3) in case of the standard molding temperature as 400°C
   For Sumikasuper E5008L is as follows;
   Nozzle: 400°C, Front: 400°C, Middle: 360°C, Hopper side: 340°C

The cylinder heating system having 4 heating zones (including nozzle) is obviously necessary for LCP molding. If the current injection machine does not have fewer than 3 heating zone (nozzle + 2 heaters for cylinder), you should consider replacing the heater system of the cylinder to a more modern one.

Of course, the above temperature setting at the hopper side is under its melting point. Accordingly, it should be noted that the flow of insufficient melt due to lower temperature setting might cause other problems. We believe that the above example is one of the most popular and effective ways to eliminate unstable metering, but we also take care about the other possibility of problems.

We should also take care of using a relatively high melt viscosity material and the case that the metering amount is larger than the ordinary capacity. Such cases sometimes induce insufficient melting.

The second recommendation is the optimization of the screw design. Now we have insufficient knowledge about the screw design, and waiting for the many researchers’ studies about the relationship between the metering and the screw design. We recommend referring to the section 3-4.

Also the abrasion of metal parts of the screw should be noted. Especially the check-ring part must be kept clean and replaced often. If the abrasion advances and the clearance become too large, it often causes plugging.

This part should be replaced each 3 or 4 months. The screw should also be replaced every 6 or 12 months whether or not any other problems occur.
5-9. Short-shot

Most people misunderstand the actual reason of short-shot problem during LCP molding. Usually the molding engineer thinks that the reasons of short-shot involves the flowability or melt viscosity of the material. It is only one of the reasons; the cause is really due to 4 reasons as shown in Fig.5-26.

First, it is a higher melt viscosity, which is the most conventional thinking way.

Second, it is due to the flow hesitation as described at section 3-3. It is very troublesome because it is difficult to distinguish between shortage of fluidity or generation of flow hesitation for non-expert engineers of LCP molding. The appearances of both reasons are very similar, however, the remedy is different. In the case of shortage by fluidity, optimization of the molding conditions will usually work. In the case of flow hesitation, it will not be soluble by the same way.

The third reason is due to the starvation derived by the plugging described in the section 5-8.

The last reason is due to dragged air. Installing the air vent is one of powerful solution if the cause is insufficient exhaust of cavity air.

We must consider about each reason and improvement.

![Fig.5-26 Classification of short-shot problem](image)

**Short-shot**

- **insufficiency of flowability**: higher viscosity
- **flow hesitation**: unbalance of flow (section 3-3)
- **starvation**: plugging (section 3-4)
- **dragging of air**: bad exhaust (section 4-4(3))

1) Insufficiency of flowability

In this case, molding temperature should be increased slightly. We usually recommend increasing 5 to 10 °C from the original setting at the nozzle and H1 (Front portion heater of cylinder). You must take care about the generation of Flash because excessive increase will produce unexpected decrease of the melt viscosity of the material.

At the same time, slight increase of the injection speed and holding pressure are also effective.

For injection speed, it must be increased more than 100 mm/sec if it is lower than such magnitude. In the case of over 100 mm/sec, every 10 mm/sec increase is recommended.

For holding pressure condition, elongation of holding time up to 2.0 sec is recommended. Since the solidification of LCP is so quick, over 2.0 sec of holding time usually has no effect. The gate will seal by solidification of LCP prior to 2.0 sec.

All procedures of this section must be done slowly because sudden changes of those parameters induce a Flash problem and it will link to break the mold part.

If no change is observed, you should consider the following reasons and procedures.

2) Flow hesitation

Sometimes this problem involves Crack (weld-crack) problem. You should also refer the explanation of "flow hesitation (section 3-3)" and "crack (section 5-5)" within this section.

It should be done carefully as flow hesitation may occur even if the thickness difference is 0.01mm for relatively thinner parts having average thickness under 0.3mm. If you observe the cavity dependence or the case that the problem generated fixed portion, you should reconfirm the distribution of cavity dimension during assembling of the mold, the dimension change of the gate due to abrasion or runner length difference from sprue.

Let us explain this kind of short-shot and its solution by using an example.

Fig.5-27 shows a bobbin item for induction coil. This item had not only a short-shot problem (generation of hole), but also a weld crack problem. The molding engineer asked us the most suitable molding condition because changing the molding temperature or increasing the injection speed did not improve these problems.

The reason for these problems was thought to be that the flow of LCP from the gate filled the upper rim first.
(Fig.5-28 a), then flowed to the bottom through inside ribs installed inside of cylindrical portion (Fig.5-28 b). After that, the material filled into the bottom rim at the same time of filling the cylindrical portion without inside ribs.

In this case, the opposite side of the gate of the cylindrical portion (especially bottom side) was the last part of filling due to flow hesitation (Fig.5-28 c).

Due to the above mentioned, all the energy due to increasing of temperature or injection speed was lost by filling of the material to the other portion.

At last, the hole and weld portion existed at indicated portions.

![Diagram](image)

Fig.5-27 Short-shot problem of bobbin for inductance coil

Now, we should consider how to equalize the flow pattern. In this case, there were 3 possibilities.

1) Thickening of cylindrical portion

Due to equalizing the flow difference between the inside ribs and cylindrical wall, thickening of the wall of cylindrical portion was recommended. The inside rib is relatively thicker than the other sections, and that is the most important reason of flow hesitation.

2) Installing of core-outs at out side of inside ribs

Installing of core-outs at out side of inside ribs is also effective for equalizing of the flow pattern because such core-out will prevent and control the flow at this thicker portion.

3) Re-allocating of the gate

Since the bottom rim is thicker than upper rim, moving of the gate to bottom side is somewhat effective. Because thicker portion should be filled firstly to prevent losing the filling energy from melt flow.

Above improvement ideas are shown in Fig.5-29. In this case, our customer carried out all procedure according to our recommendation, and the problem has improved.

There are still 2 reasons for the problem; we recommend referring to the listed sections.

![Diagram](image)

Fig.5-28 Schematic view of the LCP flow into the bobbin cavity
5 Trouble shooting during injection molding of LCP

1) thickening of cylindrical wall
2) installing of core-outs at outside of inside ribs
3) re-allocating of gate

Fig. 5-29 Solution of the problem
5-10. Sticking

Sticking to the mold is one of the most troublesome problems because it is easily linked with the mold damage, especially the damage of delicate core-pins. This is due to the stronger frictional resistance between metal part of mold (ex. core-pin) and the material.

The spraying of mold releasing reagent or replacing to an improved grade of mold release property should be considered to improve this problem. However, we should also point out that the short-shot problem could be followed by flow hesitation.

(1) Relation between “sticking” and “short-shot”

As explained in section 5-9, flow hesitation is one of the reasons for a short-shot. In this case, most molding engineers recognize that the cause of short-shot is insufficient filling of the material. They realize that they must increase the injection pressure, speed, holding pressure, or holding time. Sometimes they must try to increase the cylinder temperature. As a result, the cavity is added excess pressure and it will induce over-packing.

However, this is not successful due to the actual reason being not flow ability of the material, but with the flow hesitation.

Accordingly, we must recognize and improve the flow hesitation phenomenon for this case.

In general, sticking problems occurs when trial molding for a new mold or material replacement. In these cases, we must take care whether short-shot occurs at the flow end of the mold (usually opposite side from the gate) or not.

If the short-shot occurs at the flow end, it should be the reason of the insufficient moldability. For this case, the improvement method is very similar with normal procedure with conventional plastics.

If it occurs at the other sections, it must be induced by the flow hesitation. For this case, normal procedure to improve the short-shot problem is not suitable.

(2) Improvement of sticking

At the same time, the response of the injection machine is also considerable. If the machine is not quick in response, excess shock-pressure also induces over-packing (Fig. 5-33).

Accordingly, we would like to recommend the following:
- Replacing the material to a mold releasing grade (this must be the basic manner to improve this problem)
- Using of high response injection machine (see section 3-1)
- Modifying of the flow pattern by moving of the gate position, thickness balance of the cavity, installing of core-outs, etc. (see section 3-3).

![Fig. 5-33 Comparison of shock pressure during injection molding between high and low response machine](image-url)
5 Trouble shooting during injection molding of LCP

5-11. Warpage

As discussed in section 4, the main reason of warpage for LCP molding is the flow pattern, which brings the high orientation of LCP molecule to the flow direction.

Since LCP does not indicate Tg (glass transition temperature), the mold temperature will not influence warpage and also the dimensions. In addition, injection conditions such as cylinder temperature, injection speed & pressure or holding pressure will also not influence warpage.

In other words, the warpage of LCP molding is not changeable by molding conditions.

If the warpage problem occurs, we should investigate whether the flow pattern is appropriate or not. If the problem suddenly occurs, we should also investigate whether there is a possibility of changing the flow pattern.

For designing an appropriate flow pattern, we have already discussed in section 4.

In this section, we would like to discuss the case where the warpage trouble suddenly occurs during production – the initial warpage was improved before starting production.

Since the warpage is only influenced by flow pattern, we consider the reason why the flow pattern changes.

The reasons are considered as follows;

a. Viscosity change of material
   Each lot No. material has a melt viscosity. Usually, such viscosity difference is controlled within a certain range, which is well considered by the material company.
   However, if the specific lot No. material indicates different result of viscosity; we recommend asking to material company for help.
   On the other hand, there is a possibility that the material viscosity changes at the customer side.
   The main reasons are contamination of other material and insufficient drying before molding.
   Both reasons and improvements are discussed in section 5-4. Blister & Bubble. Please refer the section.

b. Abrasion of mold part
   Dimension change should also be considered. We often observed that only 0.01mm difference of dimension significantly changes the flow pattern. This is especially so if the dimension change occurred at the thickness of main flow portion.
   This case should be considered when the mold is maintained or additional mold is constructed. In this case, such small difference of dimension often brings different results of warpage.

c. Unsuitable molding condition
   As described above, in general molding conditions does not influence the warpage of LCP. However, if original molding conditions were not suitable, sometimes the deviation of the injection machine will influence to the flow pattern and cause warpage.
   In this meaning, the following cases are sometimes the cause of the problem;
   - too slow injection speed
   - too low cylinder temperature
   Both reasons prevent the formation of the orientation layer of LCP (so-called skin layer). In this case, the anisotropy must be lower than the case of suitable injection condition.
   The most difficult case is where the initial warpage has not been improved by modifying of the mold design.
   Sometimes modification of the original design of the mold must be necessary (please refer the section 4).
   Suitable injection molding conditions is also one of the most important responses to reduce the problems.
   Please refer the section 3 and use it under appropriate condition.
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