Investment casting using multi-jet modelling patterns: the thermogravimetric analysis of visijet® SR200 UV curable acrylate plastic

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Abstract. Rapid Prototyping (RP) technology is actively studied to be implemented in Investment Casting (IC) process. Nowadays RP techniques are studied for their feasibility as IC master patterns, in terms of pattern collapsibility and drainage during burnout. The purpose of the study is to determine the characteristic of Visijet® SR200 acrylate material during burnout process. Traditional IC patterns made from wax have properties that limit their application in precision casting, especially for parts with thin geometries that readily break or deform when handled or dipped in the refractory slurry. Furthermore, it is not economical when producing a small number of parts. Non wax patterns fabricated for IC process, revealed ceramic shell cracking due to excessive thermal expansions, incomplete collapsibility of pattern during burnout, residual ash and poor surface finish. Thermogravimetric analysis (TGA) was used to measured the weight loss of acrylate material as the temperature was increased. TGA measured the change of material’s mass as it is heated. It represents the decomposition temperature after being subjected to varying temperatures, as well as the amount of residual ash. In this experiment, the temperature range was from 20°C to 700°C with 5°C increment. Experiment results show the values of material’s optimum reaction temperature and decomposing temperature of Visijet® SR200 acrylate. The percentages of remaining materials were also monitored throughout the process to obtain the amount of residual ash. All of the temperature values obtained is a resemblance for the actual burnout process and can be used as references.

1. Introduction

Traditional IC process would require the master pattern which is made of wax to create the ceramic mould. RP technology has evolved the possibilities of implementing non-wax material as the master pattern of IC. Along with the ability to produce such complex geometry in shorter time, RP also has wide selection of material and approach to choose from. By using direct method of RP technique, a significant reduction of cycle times and production lead time in IC process [1-6]. Wax was traditionally used as it is rather easy to be shaped in various forms, either solid or liquid, for mass production, but it has quite low strength and could easily affected if exposed to high temperature
surrounding thus limiting the applications of wax pattern when it comes to high precision process [7-8].

One of the common RP process for IC implementations is Selective Laser Sintering (SLS) technique [9-16]. Usually wax is used to make IC pattern among the other SLS material because it’s thermal properties do compliments IC burn out process requirement. Pattern material should burn out completely without leaving any residue during the dewaxing process. Unfortunately, wax product may distort if it’s to be made using SLS process, affecting the final IC product. Plus, one of the serious problems of SLS is that since the solidified part is cooled rapidly, the model tends to be deformed and cracked due to the thermal stress [17].

The RP polymer pattern materials have a higher melting temperature and higher-melting viscosity thus it is very difficult to dewax [16,18]. To replace wax, it is necessary to use other material that is also easy to shape but have greater strength. Furthermore, the RP patterns need to have an excellent surface finish and must precisely recreate fine details with minimal burnout residue [14]. Besides SLS, other RP systems, such as Stereolithography (SLA) and Laminated Object Manufacturing (LOM) were tested for IC applications. However, since the epoxy material used in the SLA is a thermoset material, it does not melt like wax. It would expand and creates thermal stresses to the ceramic shell causing it to crack [9,19,20]. For the LOM application in IC, the paper which is the main material of the system would create very high amount of residual ashes during the burn out process [19].

Recent studies have shown the capabilities of pattern built from ABS using FDM technique. ABS pattern are capable to produce a clean burn out, it is robust and could be handled without damage, it also easy to be prepared and has good dimensional stability. However, the surface conditions are quite rough due to its surface layer and built condition, thus requiring surface finishing of the pattern before it could proceed with IC as the quality of final IC product are highly dependent to the quality of the pattern [21,22].

Another technique to produce the IC pattern is by using MJM. MJM technique could produce better surface finish compared to FDM [22]. MJM also be implemented in building part with internal structure as sacrificial IC pattern [23]. During burnout process, the main purpose is to completely remove the master pattern, leaving the vacant space for metal to be poured in. The temperature for burn out process depends on the type of pattern material. This paper will be focused on MJM part based on acrylate material. Thermal decomposition and weight loss are measure by using Thermogravimetric instrument. TGA is a technique in which, upon heating a material measure whether the weight increases or decreases.

2. Methodology
The implementation of RP in IC was on the part of the pattern making. The pattern is fabricated by the RP machine along with normal post processing of RP product. The RP pattern then will be dipped into a slurry then covered ceramic grain. The RP pattern will stucco a couple of the ceramic coating before the ceramic harden and turn into a shell. During burnout process, the acrylic pattern must be completely burnt out leaving the ceramic shell a perfect vacant for metal pouring. In this process, the acrylic material being heated and will start to decompose leaving just residual ashes. This part of the process is important as it is the last step before metal is being poured into a mould.

Since the materials reacts differently as it is subjected to heating, using Thermogravimetric (TG) will help to predict and understand how a material weight changes as temperature changes. It is a technique which the mass of the sample is monitored against time or temperature while the temperature of the sample, in a specified atmosphere is programmed.

Thermal analysis was carried out using Linensis Thermobalance TGA equipment. The samples were prepared in form of tiny flakes. For the testing, 70.0 mg of MJM Visijet® SR200 acrylate flakes sample was used. The sample was put into the crucible before it was inserted into the measuring system. In a thermobalance equipment system, a schematic diagram of the specific balance and furnace assembly is shown in figure 1. The sample was burnt under air atmosphere with the heating rate of 5ºC/min. The scale has to be balance with the least possible percentage of tolerance using the
potentiometer. It is very desirable to get the 0% of balance tolerance. The sample was heated to slightly 700°C with 5°C increment. The machine was left to run for 2 hours and 20 minutes.

![Schematic diagram of a typical balance and furnace assembly.](image)

**Figure 1.** Schematic diagram of a typical balance and furnace assembly.

The data obtained from the experiment is the raw data and will be manipulated using Simultaneous Thermal Analysis (STA) Software. The raw data include very high error thus it is necessary for the value to be corrected. Reference sample that was used to calibrate the system is Alumina Oxide (Al₂O₃) burnout under air atmosphere. The X-axis value will be corrected according to the weight absolute $1 \times 10^{-4}$. For the Y-axis value, the value will be smoothed according to the weight absolute $1 \times 10^{-4}$. The smoothing process should be done accordingly as the data that was too smooth will not resemble the actual character of the whole burning process. The data could then be derived into thermogravimetric data to obtain the decomposition value.

### 3. Result and Discussion

#### 3.1. Thermal decomposition

A derivative thermogravimetric (DTG) curve presents the rate of mass change (dm/dT) as a function of temperature against temperature (T) on the x-axis. Figure 2 shows typical DTG analysis curves for a sample of Visijet® SR200 acrylate heated in air atmosphere at a rate of 5°C/min. In this figure, the derivative of the curve is shown by brown line. A TG curve which is blue line represents the variation in the mass (m) of the sample with the temperature.

Within the graph, the line descends to resemble the lost of mass along the process. The material decomposition occurs in three stages. It could be seen from the graph by identifying the three curves
from the DTG that was derived from the initial TG curve. The first stage of decomposition (point 2-3) is the initial degradation in primary of the decomposition of the hard segment. During this stage, the rate of the temperature change is as much as 208.5°C, with the total weight loss of -33.85 mg. The second stage (point 4-5) is the polycondensation and poly of degradation mechanism and effected by the soft segment content [24]. For this stage, the rate of the temperature change is by 25.9°C with total mass lost of -14.15 mg. The third stage (point 6-7) is only attributing to the ash formation [21,25].

**Figure 2.** Derivative thermogravimetric (DTG) curves for Visijet® SR200 acrylate material.

In the early stage (point 1-2) of the decomposition, the material was not in stable conditions due to the presence of some volatile components [24,26]. The graph shows that there are two step of reaction which occurs at 312.8°C and 422.4°C. At 312.8°C, the rate of mass loss is 2.68mg/min while at 422.4°C the rate is -71.73mg/min. The crossing point between the both reactions created a point (point 5) which called the point of reaction which is at 387.4°C with the rate of mass loss of -3.14 mg/min. This particular point is where the maximum point of Visijet® SR200 acrylate material significantly decomposes. It caused by the break of polymer chain and polymer thermal decomposition [26,27].

### 3.2. Residual Ash

TG curves represent the variation in the mass (m) of the sample with the temperature as shown in figure 3. From the graph, the early stage of the experiment shows little changes by only -3.4% from the 70 mg sample. The line pose very small gradient up to around 200°C. From there towards 500°C, nearly -95% of the 70 mg sample has already decomposed which were resembled by a huge gradient in the graph within 200°C to 500°C and during this stage, only 3.9 mg of the sample remains. Further temperature increase completely burn the sample without leaving any residue. Though it took up to 622.1°C for the material to be completely burnt out, it is still well below the expected temperature of 700°C.
Figure 3. Thermogravimetric (TG) curves for Visijet® SR200 acrylate material.

There are many factors which influence a TGA curve. These factors may be due to instrumentation or nature of sample which could affects the shape, precision and accuracy of the experimental results, for example like the sample holder and the effect of the particle size [25]. Different TGA instrument may carry ranges of sample holder such as flat plates and deep crucibles of various capacities. The shape of the TGA curve may vary due to dissimilar heating conditions. Normally the thermocouple will be placed as near as possible without touching the sample, thus it leads to source of error due to the thermal lag and partly due to the finite time taken to cause detectable mass change. If the composition of the sample contains is such that it reacts either with the sample, or product formed or the evolved gases then this will cause error in recording the mass change of the sample.

Other than that, the particle size of the sample could also vary the result of TGA [21]. The smaller the particle size, the greater the extent of decomposition at any given temperature could occur, and the use of large crystal may result in apparent vary rapid mass loss during heating due to mechanical loss of part of the sample.

4. Conclusion
By using TGA, the burn out temperature to be used during IC for pattern made from Visijet® SR200 acrylate material could be determined. It also gives an idea how the material characteristic during the actual burn out process. It was found out from the TGA curves that the material would be able to decompose within the temperature of 700°C without leaving any residual ash which is exactly at 622.1°C. The material significantly decomposes at 387.4°C when the polymer chain breaks and the polymer thermal decompositions occur. As TGA could resemble the actual burn out process in IC, it shows the suitability of this material to be used as the pattern for IC. The material are able to be completely burnt out to make sure the quality of the casting will not be affected.

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