EFFECT OF CU-CO MIXED METAL OXIDES ON THE COMBUSTION OF PSAN-HTPB BASED SOLID PROPELLENTS

Rajan Lakra¹, Eshwar Reddy Cholleti², Puran Chandra Joshi³, Narendra Yadav⁴

¹ Student, M.E in Space Engineering and Rocketry, Birla Institute of Technology-Mesra, Ranchi, India
³ HOD & Professor, Space Engineering and Rocketry, Birla Institute of Technology-Mesra, Ranchi, India
⁴ Research Scientist-III, Space Engineering and Rocketry, Birla Institute of Technology-Mesra, Ranchi, India.

Abstract

Composite propellants based on ammonium perchlorate (AP) as oxidizer are state-of-the-art. However, global environmental impact restricts AP-based propellants because of their high chlorine exhaust. Therefore, efforts are on to innovate propellants with clean exhaust. Ammonium nitrate (AN) has, thus, regained importance in the field of propellants and explosives, more specifically in insensitive formulations, because of its clean burning and low hazard. However, the stumbling block for the application of AN - as solid propellant oxidizer has been its dimensional instability caused by phase transformation which results in increase of volume and porosity of the propellant grain. This can be overcome by the use of phase stabilized ammonium nitrate (PSAN). In the present study AN phase stabilized by incorporating 10% KN to prepare composite solid propellant. Copper-Coalt mixed metal oxides have been synthesized via citric acid complexing method. Three different catalysts were prepared with different Cu-Co molar ratios 1, 0.5 And 2 in the initial reactants. The synthesized catalysts were then added to the PSAN-HTPB (Phase Stabilized Ammonium Nitrate- Hydroxyl Terminated Poly-Butadiene) composite formulations. These propellant samples were subjected to burn rate measurement in a Crawford High Pressure Strand Burner and thermal degradation studies in Simultaneous Thermal Analyzer (STA). The thermal decomposition and burn rate of the propellant is observed maximum on the addition of Copper-Coalt mixed metal oxide catalyst with molar ratio 2. A total of six propellant samples catalyzed and non-catalyzed and one preheated virgin sample were undertaken for the above analysis and results obtained are discussed. It has been observed that except the propellant sample with Cu-Co-III and virgin propellant where PSAN was prepared by solid mixing, other propellant samples were hard to burn.

Keywords: Solid propellants, Ammonium nitrate, Phase stabilization, Burn rate, Thermal decomposition, Mixed metal oxides, Catalytic combustion.

1. INTRODUCTION

AN has drawn a considerable amount of attention due to its low cost, smokeless, availability and comparatively less toxic exhaust. It is also a chlorine free oxidizer. Its principle use is with low burning rate, low performance rocket and gas generator as it is having poor ignitability and low performance. Another drawback of ammonium nitrate is its phase instability with temperature. So it is essential to make it phase stabilized before it is used as solid oxidizer. In the present work the phase stabilized ammonium nitrate (PSAN) was prepared by adding 10% KN in saturated solution of AN and subsequently evaporating the water to get a crystalline PSAN. This PSAN was used in all the propellant samples prepared in the present work except in one virgin sample where PSAN was prepared by solid mixing method. An attempt has been made to synthesize the copper cobalt mixed metal oxides in three different molar ratios by citric acid complexing method. Three mixed metal oxides were synthesized using molar ratios of copper nitrate tri-hydrate and cobalt nitrate hexa-hydrate.

The total six propellant compositions were processed using HTPB fuel binder, PSAN oxidizer and mixed metal oxide catalysts. These propellants were subjected to burn rate study using Crawford High Pressure Strand Burner. Thermal analysis was also carried out for AN, PSAN, virgin propellants and catalyzed propellants. Attempts have been made to analyze the data and some important conclusions drawn from present experiment are highlighted.

2. EXPERIMENTAL

The experimental section of the current research includes phase stabilization of ammonium nitrate (AN), grinding and sieving of AN, synthesis of mixed metal oxide catalysts, formulation and processing of composite solid propellant with and without catalysts, burn rate measurement and thermal decomposition study of the oxidizer and prepared solid propellant.

2.1 Phase Stabilization of Ammonium Nitrate (PSAN)

Use of AN in the field of rocket propulsion may be a better choice as it is cheap, easily available and non-toxic ecofriendly smokeless products which is the need of present
scenario, but it has some disadvantages like low burning, phase changes at different phase transformation temperature and hygroscopic. And if we neutralize these negative natures of AN, it may become the best choice as an oxidizer for the solid rocket propellant.

There are lots of chemicals used to stabilize the phase changes of AN such as potassium fluoride, potassium nitrate (KN), boric acid, ammonium sulphate, ammonium orthophosphate, magnesium nitrate, etc. The amount of chemical used to stabilize the AN varies from 0.5% to 15% of the AN. These chemicals can be added by solid mixing, solution mixing or by spray.

As AN changes its phase at several phase transformation temperatures, causes the slight change in volume so to avoid that the phase stabilization has been done by adding 10% KN in aqueous solution of AN and then the solution was kept for drying to get crystalline solid form.

The process involved weighing of AN and KN which are provided by Central Drug House New Delhi. Digital weighing machine was used to weigh accurately. 1kg AN was added to 250ml of water and the mix was stirred till all the AN dissolved. To facilitate the mixing, the mixing temperature was kept 80°C. 10% KN was added and continued stirring for again 15-20 min or till it dissolved. A clear and colorless aqueous solution was obtained. The solution was spread in a steel tray and kept in oven at 55-60°C for drying for 14-18 hours. A dry solid crystalline was obtained. The same process and the weight percentage of AN and KN was repeated whenever the PSAN was required. To make a separate propellant sample 200gm of AN was also stabilized by solid mixing method. Here also the 10% KN was added same as in solution mixing.

As the dry crystalline solid PSAN was obtained in the phase stabilization process, so to make it usable for processing of the propellant and to obtain the bi-modal of AN, It was grinded by using micro-pulverizer. It was sieved by using different mess sieves in a sieve shaker. PSAN retained on 60 mesh and 100 mesh was used for processing the propellant. The approximate particle size of PSAN used was 250 microns and 150 microns.

2.2 Synthesis of Cu-Co-O Mixed Metal Oxides

All the chemicals for the experiment were of analytical grade and were used without further purification. Three different mixed metal oxides were synthesized containing different Cu-Co molar ratios in the initial reactants. In a typical synthesis, copper(II) nitrate trihydrate (Cu(NO\textsubscript{3})\textsubscript{2}.3H\textsubscript{2}O) and cobalt(II) nitrate hexahydrate (Co(NO\textsubscript{3})\textsubscript{2}.6H\textsubscript{2}O) were dissolved in 25ml distilled water in china dish to obtain a mixed metal nitrate aqueous solution. Then citric acid was added to these solutions. After stirring for 10 min each, the solutions were heated at 105°C for 4 hours in hot air oven to evaporate the water solvent to produce viscous gels.

The gels were then dried at165°C in a muffle furnace for 2 hours. The resulting mass was the bubbly dark powders. The precursors were successively heated at 600°C for 3 hours to obtain final black copper cobalt mixed metal oxides. The same synthesis process has been carried out to get the required amount of catalysts. In a single synthesis around 0.3 to 0.4 grams of one sample catalyst was collected. These three metals oxides were then stored in separate sealed containers to avoid external effects. Table 1 shows the catalysts prepared in the present work. And above plate 1 shows the initial stage of catalytic synthesis.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Copper nitrate(gm)</th>
<th>Cobalt nitrate(gm)</th>
<th>Citric acid (gm)</th>
<th>Molar ratio(Cu/Co)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu-Co-I</td>
<td>0.6</td>
<td>0.72</td>
<td>3.15</td>
<td>1.0</td>
</tr>
<tr>
<td>Cu-Co-II</td>
<td>0.6</td>
<td>1.44</td>
<td>3.15</td>
<td>0.5</td>
</tr>
<tr>
<td>Cu-Co-III</td>
<td>1.2</td>
<td>0.72</td>
<td>3.15</td>
<td>2.0</td>
</tr>
</tbody>
</table>

2.3 Processing of Solid Propellants

The required amount of Ammonium Nitrate in coarse to fine ratios 75:25. HTPB, DOA, IPDI were properly weighed. The mixing was done at 60°C. HTPB was heated in a bearer using vext bath. Mixing was started by dehumidifying the HTPB and DOA and then mixing thoroughly by stirring for 10 minutes. The IPDI was added in the mix and stirred for again 10 minutes. Ammonium Nitrate coarse and then fine was added in slots and the mixing was done for around 30 minutes to get consistent slurry.

The propellant slurry was casted in Aluminum flat plate mold which was properly cleaned, greased and was covered by aluminum foil for easy removal of propellant sheets. After casting the slurry, the mold was kept on vibrator for around 30 minutes to remove voids which were formed during casting. The mold was kept in an electric vacuum oven at 60°C for a period of 6 days to allow the propellant to cure and attain the required mechanical strength. The mold was then taken out from the oven and the propellant was removed from mold. It was cleaned and stored in desiccator to avoid the absorption of moisture from atmosphere.
2.4 Materials and Methods

Simultaneous Thermal Analysis refers to the simultaneous application of Thermo Gravimetric Analysis (TGA) which is also called Thermograviometry and Differential Scanning Calorimetry (DSC) to the same sample in the same instrument. In the present experiment Netzsch STA 409 PC Luxx was used. The temperature range of STA is ambient to 1100°C for the model Netzsch STA 409 PC Luxx. Heating rate is 0.01 – 60°C/min and uses nitrogen atmosphere.

Crawford Burner is an apparatus used to determine the burn rate of a solid rocket propellant in an elevated pressure environment. The propellant that is being tested is called a strand and is burned in a pressurized tank called a firing vessel. The strand is a thin bar of propellant which is coated so that the burning cross-sectional area is restricted. The strand is electrically ignited at one end and the duration of time for which it burns along the length is measured with the help of fuse wires embedded in the strand. These wires are connected to the timer via an electric relay. When the first wire cuts, the timer unit starts and when the second wire is cut, it stops. The burn time is recorded and the burn rate is determined with the help of burn time and burn length. The schematic of the Crawford strand burner setup is given in figure-1. The burn rate measured is typically 4 – 12% less than the actual burn rate observed in rockets because the high temperature condition that is prevalent in actual rockets is not simulated.

Details of the Crawford Strand Burner

Model & Make: Indigenously made by BIT, Mesra, India

Micro Palvarizer was used to grind the PSAN to get the two different size ranges of oxidizer particles which were 150µ and 250µ to get the maximum fraction loading while propellant processing.

Sieve shakers are used to collect the different collect the oxidizer particle sizes which can be defined by mesh sizes i.e., hole per squire inches in the sieves.

Table 2: Composition of solid propellants

<table>
<thead>
<tr>
<th>PSAN</th>
<th>HTPB</th>
<th>DOA</th>
<th>IPDI</th>
<th>Cu-Co-I</th>
<th>Cu-Co-II</th>
<th>Cu-Co-III</th>
</tr>
</thead>
<tbody>
<tr>
<td>75</td>
<td>18.92</td>
<td>4.6</td>
<td>1.46</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>75</td>
<td>18.92</td>
<td>4.6</td>
<td>1.46</td>
<td>2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>75</td>
<td>18.92</td>
<td>4.6</td>
<td>1.46</td>
<td>-</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>75</td>
<td>18.92</td>
<td>4.6</td>
<td>1.46</td>
<td>-</td>
<td>-</td>
<td>2</td>
</tr>
<tr>
<td>80</td>
<td>15.14</td>
<td>3.68</td>
<td>1.168</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

All quantities are in weight Percentage

3. RESULTS AND DISCUSSION

Mixed metal oxide catalyst has very high potential to enhance the decomposition of composite solid propellants. The present study aims to synthesize copper-cobalt mixed metals oxides with different Cu-Co molar ratios (1, 0.5 and 2) in the initial reactants and evaluate the effect of these mixed metal oxides on the combustion characteristics of PSAN-HTPB based composite solid propellants. Citric Acid (CA) complexing method was used to synthesize the copper cobalt mixed metal oxides. Well-0 synthesized mixed metal oxides were obtained after the CA-Cu-Co precursors were calcined at 60°C for 3 hours. The propellants used in the present study consisted of 80% PSAN and 75% PSAN. The remaining were fuel binder, plasticizer, curing agent and catalyst in a definite proportion. The ammonium nitrate was made phase stabilized by incorporating 10% potassium nitrate.

The present work centered around the study of thermal behavior of propellant sample as well as oxidizer and burning rate of propellants formulated.

3.1 Thermal Analysis

Thermal decomposition studies of any substance gives valuable information about the condensed phase and gaseous phase combustion processes in the combustion reaction. In the present work, the thermal degradation analysis was carried out for all the propellant samples i.e. virgin PSAN-HTPB propellant, virgin preheated PSAN-HTPB propellant, (Solid mixed) PSAN-HTPB propellant, PSAN-HTPB 80/20 propellant, PSAN-HTPB with 2% Cu/Co-I, PSAN-HTPB with 2% Cu/Co-II, PSAN-HTPB with 2% Cu/Co-III and AN, PSAN, PSAN prepared by solid mixing, PSAN stored for 3 months and PSAN stored for 3 months and preheated. The results are presented in figure-2-5. The heating rate in all the experiments was kept at 10°C/min and the heating was done up to 1000°C. The individual TGA and DSC test have been carried out and the comparisons graphs are shown here for the both oxidizer, modified oxidizer, propellant samples and catalyzed propellant samples as mentioned above.

![Crawford high pressure strand burner](image-url)
TGA of pure AN describes weight loss of it on heating it from room temperature 28°C to 600°C. It is observed that it starts decomposing from temperature 220°C and the almost complete decomposition occurs by 303°C. A total of 90.22% mass is decomposed in this temperature range. Further heating does not show any weight loss till 600°C.

TGA of PSAN describes weight loss of PSAN with the temperature up to 900°C. It is observed that the weight loss occurs in two different temperature ranges. Weight loss about 81.32% occurs in the temperature range 180-330°C. Then the next weight loss about 11% occurs in the temperature range 540-780°C. The total weight loss is around 92.32% till 800°C. This two temperature decompositions may be due to the different temperature for the decomposition of AN and KN.

TGA of preheated PSAN describes weight loss of PSAN in the temperature up to 1000°C. In this case the PSAN was heated at 60°C for 3-4 hours before subjecting it for thermal analysis. It is observed that the weight loss occurs in the two different temperature ranges. Weight loss about 89% occurs in the temperature range 200-340°C. The next weight loss about 8.70% occurs in the range 575-800°C.

TGA of solid mixed PSAN describes the weight loss of PSAN up to 500°C. In this case the PSAN was prepared by solid mixing process. Weight loss occurs in one time decomposition. Decomposition starts at 150°C and ends by 350°C then no change in weight occurs. About 88.87% of mass loss occurs in this temperature range.

A comparison of TGA thermogram in the temperature range up to 350°C for AN, PSAN (Prepared by solution mixing/solid mixing) and PSAN preheated is presented in fig 2. Pure AN decomposes up to 90.22% by 305°C. The PSAN made by solid mixing method the 88.9% decomposition takes place by 285°C. The decomposition temperature range changes when AN was made phase stabilized by the addition of KNO₃ solution mixing method/solid mixing method. Freshly prepared PSAN decomposes up to 81.32% by 290°C however PSAN stored for 3 months, the decomposition temperature range extends further. About 81.32% and 89% decomposition occur by 310°C for preheated and non-heated samples.

The description of TGA thermograms for all the propellant samples prepared in the present work are mentioned below and a comparison TGA thermogram showing the different weight loss of propellant samples as shown in fig-4. These thermograms highlight that propellant sample shows the % weight loss in 3-4 different temperature ranges. The rate of weight losses in these temperature ranges is different. The weight loss in the first temperature range may be attributed to the removal of moisture and some low temperature volatile ingredients present in the propellant sample. The major weight loss occurs in the temperature range 175°C to 600°C as a result of decomposition and removal of decomposition products of PSAN as well as the evaporation and degradation of polymeric fuel binder, plasticizer and curing agents and also the reaction products. It is noticed that the rate of decomposition is higher in the first two temperature ranges.

TGA thermogram for virgin PSAN-HTPB propellant sample describes the weight loss in the temperature range 28°C to 1000°C. The weight loss occurs in three decomposition temperature ranges. The decomposition starts 175°C and about 24.32% mass loss occurs by 225°C. About 42.82% weight loss occurs in between 225-280°C. The finally about 26.09% weight loss occurs in the temperature range 280-525°C.

TGA thermogram of virgin PSAN-HTPB propellant sample weight loss of it in the temperature from 0-600°C which was preheated at 60°C for 2-3 hours before subjecting to thermal decomposition studies. Weight loss occurs in four stages. Decomposition starts at 175°C and about 12.82% weight loss occurs by 220°C. Second stage decomposition continues from 220°C and about 56.82% weight loss occurs by 280°C. On comparing above two thermograms of propellant unheated and heated, it is seen that nearly the same weight loss (about 93%) occurs by 600°C. It is noticed that preheated sample reveals four temperature range of varying decomposition range whereas non-heated sample three temperature range of different decomposition rate.

TGA of PSAN-HTPB 80/20 virgin sample describes weight loss of it in the temperature up to 700°C. Weight loss occurs in three temperature ranges. Decomposition starts at 175°C and about 12.08% weight loss occurs by 220°C. It continues decomposing and about 58.85% of weight loss occurs by 260°C. Then further weight loss about 17.97% occurs at slower rate till 600°C.

TGA of PSAN-HTPB propellant sample describes the weight loss of it in the temperature up to 700°C. In this case PSAN was prepared by solid mixing method. Weight loss occurs in three stages. Decomposition starts at 175°C and about 18.60% weight loss occurs by 220°C. The decomposition continues and about 57.39% weight loss occurs by 260°C. It continues at slower rate after 260°C and about 22.57% weight loss occurs by 600°C.

A comparison of above four virgin PSAN-HTPB propellant samples have been taken in the first two stages of
decomposition where the rapid weight loss occurs. It is observed that the weight loss of propellants differs markedly with shifting the decomposition temperature. The total weight losses are 67.14%, 69.64%, 70.93% and 75.99%. 67.14% is up to 2nd temperature range for virgin PSAN-HTPB propellant. When the same was preheated for 2-3 hours at 60°C before subjecting it to thermal analysis about 2.5% higher mass loss was seen as compared to unheated propellant up to 2nd temperature range. 3.79% higher weight loss occurs when the oxidizer % increased by 5%. 8.85% higher weight loss occurs up to 2nd temperature range when the PSAN was prepared by solid mixing method for the virgin PSAN-HTPB propellant formulation.

TGA for PSAN-HTPB propellant with 2% Cu-Co-I catalyst shows the weight loss of it in the temperature from 0-600°C. Three stages decomposition is noticed. The weight loss starts from 150°C and about 27.54% weight loss is completed by 230°C. There is 35.84% decomposition between 230°C and 265°C. Next the slow decomposition is noticed in the temperature range 265°C and 600°C where about 27.85% weight loss occurs. It is also noticed that nature of weight loss curve is similar to curve of non-heated and preheated virgin PSAN-HTPB propellant in the temperature range 150-260°C. However the rate of weight loss slowed down in the nest temperature range. These results indicate that Cu-Co-I de-accelerates the decomposition rate of PSAN-HTPB based solid propellant.

TGA for PSAN-HTPB propellant containing 2% Cu-Co-II catalyst represents the weight loss with temperature up to 1000°C. In this case decomposition occurs in four stages. There is 24.72% weight loss in the temperature range 150-230°C. It is observed that the decomposition process is faster in the second temperature range and 40.89% mass loss occurs in between 230-275°C. There is 16.73% decomposition between 275°C and 425°C. It continues decomposing at slow rate and 14.95% weight loss occurs between 425-600°C. The weight loss rate in the 2nd temperature range is very fast for a short duration then slows down asin the propellant sample with 2% Cu-Co-I.

TGA for PSAN-HTPBpropellant with 2% Cu-Co-III describes the weight loss of it in the temperature up to 600°C as shown in fig-3. Three stages decomposition process is observed in this case also. In the first stage about 18.57% decomposition occurs between 180°C and 230°C. After that the decomposition rate increases and about 58.09% weight loss is noted in between 230°C and 250°C. Finally there is 13.50% decomposition between 250°C and 510°C. A total about 92.16% decomposition occurs by 600°C.

A comparison of weight loss results in all above catalyzed and non-catalyzed propellant shows that the weight loss is highest in the second temperature for all the propellant samples studied in the present work. The percentage of residues left for above mentioned propellant samples are 5.77%, 7.01%, 11.10%, 1.44%, 8.77%, 2.71%, 7.84% respectively after 600°C.

The TGA Thermograms of all the propellant samples have been presented for comparison purposes in figure 4. The TGA Thermograms of solution mixing virgin propellant samples and propellant sample with Cu-Co-I and Cu-Co-II catalysts are almost similar whereas the TGA thermograms of solid mixed virgin propellant and the propellant with 2% Cu-Co-III differs from remaining all other thermograms but their nature is similar. All the thermograms show either a three or a four stage decomposition in which the catalytic activity is highest in the second stages. The weight losses in the virgin solid mixed PSAN-HTPB propellant and propellant sample with 2% Cu-Co-II have higher and also higher decomposition rate in the second stage as compare to other catalyzed and non-catalyzed propellant samples. It is also noticed that decomposition temperatures have shifted for PSAN-HTPB propellant with 2% Cu-Co-III in the lower temperature side. These results indicate that the solid mixing is effective method for phase stabilization of AN and to improve the decomposition behavior of PSAN-HTPB solid propellant the mixed metal oxide catalyst Cu-Co-III has potential effect.
3.1.1 DSC of Oxidizer

In Differential Scanning Calorimetry, the instantaneous temperature of a sample is compared with the same sample in inert reference material during the programmed change of temperature. The temperature should be the same until thermal event occurs, such as melting, decomposition or change in the crystal structure. In an endothermic event takes place within the sample, the temperature of the sample will vary with that of the reference so a minimum variance will be observed on the curve with respect to reference. On the contrary, if an exothermal event takes place, then the temperature of the sample will exceed that of the reference and a maximum variance will be observed on the curve.

DSC test shows that AN reveals four endothermic peaks on the heating from room temperature 28°C to 600°C. The first two endothermic peaks with the onset temperature 59.14°C and 134.4°C, respectively, are observed due to two phase transformation of pure AN. The third endothermic peak having an onset temperature of 170.6°C represents the absorption of heat for the melting of AN. The 5th endothermic peak whose onset temperature is 302.2°C represents the absorption of heat for the final and complete decomposition. These results are in confirmation of TGA curve.

DSC curve for PSAN reveals five endothermic peaks on heating it from room temperature 28°C to 500°C. The first two endothermic peaks with onset temperature as 120.2°C and 130.2°C are observed, respectively to the two phase modification of PSAN. The third endothermic peak, having an onset temperature of 159.7°C represents the absorption of heat for the melting of PSAN. And the fourth and fifth endothermic peaks with onset temperature 207°C and 282°C shows the heat absorption for the final and complete decomposition of PSAN.

DSC curve for preheated PSAN also shows the different phase changes of it. In this case also four endothermic peaks are obtained on heating from room temperature 28°C to 1000°C. The first two endothermic peaks with onset temperature as 120.5°C and 132.9°C are observed respectively due to the phase transformation of PSAN. The third endothermic peak with onset temperature 163.4°C represents the absorption of heat for the melting of PSAN. Fourth endothermic peak with onset temperature 303.5°C shows the heat absorption for the final and complete decomposition of PSAN.

DSC curve for PSAN made by solid mixing reveals four endothermic peaks from room temperature 28°C to 550°C. The first two endothermic peaks with onset temperatures as 93.1°C and 129.2°C are observed respectively due to the two phase transformation of PSAN. The Third endothermic peak with onset temperature 163.8°C represents the absorption of heat for the melting of PSAN. And the fifth endothermic peak with onset temperature 275°C shows the absorption of heat for the final and complete decomposition of the PSAN.

Figure 5 shows a comparison of DSC curves for AN, PSAN, preheated PSAN, PSAN prepared by solid mixing and fresh PSAN. These DSC curves show that all the oxidizer endothermic peaks from room temperature 28°C to 600°C. The first two endothermic peaks represent phase transformation but the differences are in peaks temperature or the onset temperature shifts depend on the type of PSAN. Third peak represents heat absorption for melting of treated and non-treated AN. The last peak shows absorption of heat for final and complete decomposition of treated and non-treated AN. It is noticed that melting temperature of treated AN has shifted. Fresh PSAN and stored PSAN shows nearly same result but little shifting occur in their melting and decomposition onset temperature. In the present research solid mixed PSAN shows faster decomposition as compared to pure AN and solution mixed PSAN.

3.1.2 Differential Scanning Calorimetry of Propellant Samples

Table 3 presents the temperatures at which endothermic and exothermic process of virgin PSAN-HPTB and the propellant samples with catalysts takes place. The endothermic peaks are almost at same temperature for six propellant samples. Only PSAN-HPTB propellant sample with Cu-Co-III has somehow lesser endothermic peaks. But there is a clear difference in the exothermic peak temperatures between the virgin PSAN-HPTB sample and the PSAN-HPTB samples with catalyst. There are three or more than three exothermic peaks for all the propellant samples except, Propellant sample with 2% Cu-Co-III catalyst which has two exothermic peaks.

From the individual DSC thermograms for virgin PSAN-HPTB and PSAN-HPTB propellant with 2% Cu-Co-I, Cu-Co-II and Cu-Co-III mixed metal oxides respectively, we can notice that the endothermic peak temperature is almost similar for all the propellant samples. However, there is a marked difference in the exothermic peaks which suggest that the decomposition of the propellant samples with copper cobalt mixed metal oxides is rapid at a certain temperature.
DSC curve of virgin PSAN-HPBP Propellant sample which illustrates the endothermic and exothermic behaviors of it. The heating rate was 10k/min. Two endothermic peaks of onset temperature as 133.3°C and 164.2°C are observed. The first endothermic represents phase changes of virgin PSAN-HPBP propellant sample and the second endothermic peak represents absorption of heat for the melting of propellant sample. On further heating the three exothermic peaks are absorbed with onset temperatures as 220.0°C, 239.1°C and 416.0°C respectively. The net enthalpy evolved is 1282J/g for the first two exothermic peaks and 2542J/g for third exothermic peak.

The DSC thermogram of virgin PSAN-HPBP preheated propellant illustrates the endothermic and exothermic behaviors of it, which was heated at 60°C for 3 hours before subjecting to DSC analysis. Nearly the same processes evolve. The same two endothermic peaks with onset temperature 133.6°C and 164.2°C are observed. Three exothermic peaks with onset temperatures are 213.5°C, 243.3°C and 413.1°C are also observed however the net enthalpy change is 1103 J/g for first two exothermic peaks and 1926J/g for third exothermic peak found lower as compared to the unheated propellant.

The DSC thermogram of virgin PSAN-HPBP propellant sample illustrates the endothermic and exothermic behaviors of it in which the PSAN was prepared by solid mixing of AN and KN. The two endothermic peaks with onset temperatures 133.9°C and 173.5°C are observed where first one is nearly same but the 2nd one is higher. The exothermic peak temperatures are 226.3°C, 391.6°C and 446.2°C are observed which differ from solution mixing PSAN-HPBP propellant sample. The heat evolved during first exothermic peak is 1244J/g which is higher as compared to solution mixing PSAN-HPBP virgin sample. The 2nd exothermic peak shifts to higher temperature side and is very close to 3rd exothermic peak with total 1267J/g heat evolved.

The DSC thermogram of PSAN-HPBP 80/20 sample illustrates the endothermic and exothermic behavior of it. The endothermic peaks with onset temperatures 134°C and 166.3°C are nearly same as compare to the solution mixing virgin propellant samples. It is noticed that the three exothermic peaks with onset temperatures 221.4°C, 383.6°C and 413°C differ from the exothermic peaks of solution mixed PSAN-HPBP virgin samples but they are similar to the solid mixed PSAN-HPBP virgin sample. The 1134J/g net enthalpy evolved during first exothermic peak and 1086J/g during 2nd and 3rd exothermic peak is lesser than solid mixed PSAN-HPBP virgin sample.

The DSC thermogram of PSAN-HPBP propellant with 2% Cu-Co-I illustrates the endothermic and exothermic behavior of it. The heating rate is same like in other propellant samples. Two endothermic peaks with onset temperatures 121.6°C and 160.4°C are observed which are same like all other virgin propellant samples but the onset temperatures are at lower temperature side. The drastic changes have been observed in the exothermic peaks and the onset temperature exothermic reactions and the heat evolved. Four exothermic peaks are observed with onset temperatures 237°C, 266°C, 380.9°C and 486.4°C respectively whereas the virgin sample has only 3 peak with onset temperatures 220°C, 239.1°C and 416°C. This indicates that the Cu-Co-I alters the reaction mechanism of the propellant. It also reflects that the rate of reaction being altered on the addition of Cu-Co-I.

The DSC heating thermogram of PSAN-HPBP propellant with 2% Cu-Co-II illustrates the endothermic and exothermic behavior of it. Two endothermic peaks and five exothermic peaks are observed on heating it from 28°C - 800°C. The endothermic peaks are nearly similar to endothermic peaks of virgin propellant samples. The drastic changes are observed in the exothermic peaks with onset temperatures and the net enthalpy evolved. The onset temperatures and number of peaks are higher which indicates that Cu-Co-II enhances the decomposition of propellant but at different temperature region. These results reflect that the decomposition processes are spread over on a wider temperature segment and heat evolved also in steps which make the propellant hard to combust. So it can be said that enhancement at same temperature can produce higher rate of decomposition as well as maximum net enthalpy.

The DSC thermogram PSAN-HPBP propellant sample with 2%Cu-Co-III illustrates the endothermic and exothermic behavior of it. Two endothermic peaks and two exothermic peaks are observed on heating the propellant from room temperature 28°C to 600°C. It is noticed that the endothermic peaks are similar to the endothermic peaks of all virgin propellant samples. Unlike the Cu-Co-I, II’s effect of rate of decomposition of PSAN-HPBP solid propellant sample Cu-Co-III has single peak point with onset

<table>
<thead>
<tr>
<th>Table -3: DSC results of different propellants samples</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Samples</strong></td>
</tr>
<tr>
<td>Virgin PSAN-HPBP</td>
</tr>
<tr>
<td>Virgin PSAN-HPBP</td>
</tr>
<tr>
<td>Virgin PSAN-HPBP (SM)</td>
</tr>
<tr>
<td>Virgin PSAN-HPBP (00/20)</td>
</tr>
<tr>
<td>PSAN-HPBP + Cu-Co-I</td>
</tr>
<tr>
<td>PSAN-HPBP + Cu-Co-II</td>
</tr>
<tr>
<td>PSAN-HPBP + Cu-Co-III</td>
</tr>
</tbody>
</table>

The DSC thermogram of PSAN-HPBP 80/20 sample illustrates the endothermic and exothermic behavior of it. The endothermic peaks with onset temperatures 134°C and 166.3°C are nearly same as compared to the solution mixing virgin propellant samples. It is noticed that the three exothermic peaks with onset temperatures 221.4°C, 383.6°C and 413°C differ from the exothermic peaks of solution mixed PSAN-HPBP virgin samples but they are similar to the solid mixed PSAN-HPBP virgin sample. The 1134J/g net enthalpy evolved during first exothermic peak and 1086J/g during 2nd and 3rd exothermic peak is lesser than solid mixed PSAN-HPBP virgin sample.

The DSC thermogram of PSAN-HPBP propellant with 2% Cu-Co-I illustrates the endothermic and exothermic behavior of it. The heating rate is same like in other propellant samples. Two endothermic peaks with onset temperatures 121.6°C and 160.4°C are observed which are same like all other virgin propellant samples but the onset temperatures are at lower temperature side. The drastic changes have been observed in the exothermic peaks and the onset temperature exothermic reactions and the heat evolved. Four exothermic peaks are observed with onset temperatures 237°C, 266°C, 380.9°C and 486.4°C respectively whereas the virgin sample has only 3 peak with onset temperatures 220°C, 239.1°C and 416°C. This indicates that the Cu-Co-I alters the reaction mechanism of the propellant. It also reflects that the rate of reaction being altered on the addition of Cu-Co-I.

The DSC thermogram of PSAN-HPBP propellant with 2% Cu-Co-II illustrates the endothermic and exothermic behavior of it. Two endothermic peaks and five exothermic peaks are observed on heating it from 28°C - 800°C. The endothermic peaks are nearly similar to endothermic peaks of virgin propellant samples. The drastic changes are observed in the exothermic peaks with onset temperatures and the net enthalpy evolved. The onset temperatures and number of peaks are higher which indicates that Cu-Co-II enhances the decomposition of propellant but at different temperature region. These results reflect that the decomposition processes are spread over on a wider temperature segment and heat evolved also in steps which make the propellant hard to combust. So it can be said that enhancement at same temperature can produce higher rate of decomposition as well as maximum net enthalpy.

The DSC thermogram PSAN-HPBP propellant sample with 2%Cu-Co-III illustrates the endothermic and exothermic behavior of it. Two endothermic peaks and two exothermic peaks are observed on heating the propellant from room temperature 28°C to 600°C. It is noticed that the endothermic peaks are similar to the endothermic peaks of all virgin propellant samples. Unlike the Cu-Co-I, II’s effect of rate of decomposition of PSAN-HPBP solid propellant sample Cu-Co-III has single peak point with onset
temperature 229.1°C. This is similar to the endothermic as well as the exothermic peaks of solid mixed PSAN-HTPB virgin propellant. The enthalpy change during the process is about 1424 J/g. The steep rise in temperature indicates the fast reaction and higher enthalpy change comparatively per unit time.

![DSC result of propellant sample with 2% Cu-Co-III](image)

**Fig-6:** DSC result of propellant sample with 2% Cu-Co-III

Figure 7 shows the comparison of DSC thermogram which illustrates the endothermic and exothermic behavior of PSAN-HTPB propellant samples. It is noticed that the endothermic peaks of all the propellant samples are nearly same. The drastic changes in exothermic peaks have been observed. The exothermic peak of Propellant with Cu-Co-III has highest hike and its 2nd exothermic peak has broaden. The resulting higher heat generation at single temperature point makes the propellant to generate sufficient heat for the combustion process. So it can be said that the Cu-Co-III has capability to enhance the burn rate. Solid mixed PSAN-HTPB virgin sample and PSAN-HTPB 80/20 sample have comparatively same endothermic and exothermic peaks. The exothermic peaks of remaining all other samples are distributed at different temperature points that results the distribution of heat at different temperature causes lack of sufficient heat generation for the combustion to sustain.

![DSC of propellant sample](image)

**Fig-7:** DSC of propellant sample

3.2 Burn Rate Studies

The burn rate studies at various pressures were carried out using a Crawford High Pressure Strand Burner. The propellant strands were electrically ignited and the time taken to burn for 50 mm length of the strand was recorded and then burn rate was calculated by dividing length by time taken to burn the length. Nitrogen gas was used to pressurize the strand burner to simulate the operating chamber pressure conditions.

It is observed that the propellant made from PSAN solid mixing method and solution mixing method with 2% Cu-Co-III burns smoothly in this pressure range. Other propellant samples either did not burn or burnt only 2-3 cm and extinguished after that. The burning rate – pressure data for the propellant burnt completely are presented in figure 8. It is observed that the increasing the pressure increases the burn rate of propellants investigated in the present work. However the catalyzed propellant is more sensitive to pressure effect as compared to virgin propellant. Catalyzed propellant has lower burn rate compared to virgin propellant till 400 psi but above this pressure it crosses the burn rate of virgin propellant. This increase of burn rate at pressure above 400psi for catalyzed propellant shows that Cu-Co-III catalyst is more active for PSAN-HTPB propellant.

![Variation of burning rate with pressure of PSAN-HTPB propellant](image)

**Fig-8:** Variation of burning rate with pressure of PSAN-HTPB propellant with and without catalyst.

4. CONCLUSIONS

The following conclusions can be drawn from the present investigations on the combustion characteristics of PSAN-HTPB composite solid propellants with and without the copper cobalt mixed metal oxides.

1. The TGA thermograms of solid mixed virgin propellant, 80/20 virgin propellant, and propellant sample with 2% Cu-Co-III catalysts are almost similar whereas a rapid decomposition can be observed. All the thermograms show a three staged decomposition in which the rate of decomposition is highest in the 2nd stage.

2. Form the Simultaneous Thermal Analyzer it was noticed that the endothermic peaks are almost at same temperature for all the propellant samples but there is a clear difference in the exothermic peak temperatures between the virgin PSAN-HTPB sample and the propellant sample with 2% Cu-Co-III. Propellant of 80/20 PSAN-HTPB and solid mixed virgin propellant have similar endothermic and exothermic peaks like propellant sample with 2% Cu-Co-III.
3. There are four exothermic peaks for the propellant sample with 2% Cu-I catalyst, five exothermic peaks for the propellant sample with 2% Cu-II and two exothermic peaks for the propellant with 2% Cu-III. It indicates the different effects of various molar ratios of copper cobalt mixed metal oxides also indicate that the rate of heat release is maximum for propellant with 2% Cu-Co-III.

4. Results show that Cu-Co-I and Cu-Co-II either effect the combustion reaction mechanism of PSAN-HTPB propellant or they have a negative effect on the reaction mechanism. Whereas the catalyst Cu-Co-III accelerate the rate of heat release and weight loss confirming it as a good catalyst for PSAN-HTPB propellant.

5. From the phase stabilization point of view, the addition of 10% potassium nitrate to ammonium nitrate is quite effective to shift the endothermic peaks in the higher temperature side.

6. It has been observed that the addition of 2% Cu-Co-III increases the burning rate of the propellant composition studied in the present investigation. It is observed that the increasing the pressure increases the burn rate of propellant with 2% Cu-Co-III and PSAN-HTPB virgin propellant (where PSAN was prepared by solid mixing) burnt smoothly while the other propellants didn’t sustain burning. The propellant with 2% Cu-Co-III is more sensitive to pressure effect as compared to solid mixed virgin propellant but this effect comes above 400psi. This increase in burn rate with pressure for Cu-Co-III catalyzed propellant shows that the Copper cobalt mixed metal oxide catalyst with 2 molar ratios is more active for the propellants considered in the present work whereas catalyst Cu-Co-I and Cu-Co-II do not effect in the combustion of PSAN-HTPB composite propellant.

7. Although efforts of mixed metal oxide on the combustion of PSAN-HTPB based composite solid propellant and phase stabilization effect of AN have been made to explain the results obtained in the present work. However it is felt that more elaborate investigation on different aspects of combustion with catalysts is required to reach any specific conclusion.

5. FUTURE SCOPE OF WORK

In the present work an attempt has been made to synthesize copper-cobalt mixed metal oxides by citric acid complexing method. The synthesized catalysts were added to the PASN-HTPB composite formulations which were then subjected to burn rate measurement and thermal degradation studies. The ammonium nitrate was made phase stabilized by incorporating 10% potassium nitrate. Apart from these, there are several other aspects which can be investigated and studied. Some of the important areas which can be explored are

1. The copper cobalt mixed metal oxides can be synthesized with different Cu-Co molar ratios in initial reactants. The catalysts can be prepared at different calcination temperatures ranging from 400 °C to 900 °C. In the present work the CA-Cu-Co precursors were calcined only at 600 °C.

2. The copper cobalt mixed metal oxides can be synthesized by other synthesizing techniques which may improve the effect of mixed metal oxides on PSAN based composite solid propellant.

3. It is worthwhile to study the PSAN-HTPB based composite propellant where the PSAN can be prepared by adding 0.3-20% of other chemicals which might stabilize as well as increase the decomposition rate of PSAN-HTPB based composite propellant. Such study will be a good achievement to make propellant with chlorine free exhaust.

4. The use of different fine/course ratio of oxidizer along with different proportions of catalysts will be very helpful to tailor the burning rate of PSAN based composite solid propellants.

5. To increase the burning rate of PSAN based propellant the blending of fuel binder can be done in definite proportion. Such study might be useful to make ecofriendly propellant with high performance.

6. A study of PSAN based propellants by using fast burning fuel binders such as PTHF and PDMS with varying percentage of copper-cobalt mixed metal oxide catalysts will be much beneficial to have a propellant with wide range of applications.

ACKNOWLEDGEMENT

The authors would like to acknowledge the technical help and guidance of Prof. Dr. J. K. Prasad, Prof. Priyank Kumar, Mr. Satish Kumar, Mr. Sachinder, Mr. Ashok Kumar, Mr. Gangaram, Mr. Arun Mishra, Mr. Jitram, and Mr. Hareram, who have supported and helped us in our work.

REFERENCES

Basic format for books:

[1] Summerfield M.; Sutherland G. S.; Webb M. J.; Tabock H. H.; and Hall K. P.

Basic format for periodicals:

HTPB/PTHF blend binder” NDA Japan Article 24 Jan 2012.


**Basic format for reports:**


**Basic format for journals (when available online):**


[33] R. Prasad; P. Singh, “Applications and Preparation Methods of Copper Chromite Catalysts: A

**Basic format for papers presented at conferences**

*when available online:*


**BIOGRAPHIES**

**Mr. Rajan Lakra** M.E in Space Engineering & Rocketry from BIT, Mesra. Ranchi, India. After completing B.E in Aeronautical Engineering From Anna University, presently working as Assistant Prof at Institute of Aeronautical Engineering, Hyderabad

Email id: rajanlakra@gmail.com

**Mr. Cholleti Eshwar Reddy** pursuing M.Tech in Thermal Branch from PRIT affiliated to JNTU - Hyderabad, India. After completing B.Tech in Aeronautical Engineering from esteemed MLR Institute of Technology affiliated to JNTU - Hyderabad with distinction.

Email id: Eshwar.ch1808@gmail.com

**Prof. Dr Puran Chandra Joshi** Ph.D in Rocket Combustion process from BIT Mesra. Former VC, Dean and HOD at BIT Mesra.

**Dr. Narendra Yadav**. Ph.d in Solid Rocket propellant, Working as Research Scientist III at BIT, Mesra