Analysis of Composite Propellants Formulated from Maize Stalk and Pawpaw Stem

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Abstract:  
Plant biomass (Carbon) was sourced from maize stalk and paw paw stem were used as fuel to composite solid propellant. Composite solid propellant was conducted using three ratios in this order; Ammonium nitrate, Biomass (Carbon), Sulphur, Starch and Gum Arabic as binders in ratios of 80:16:2:2 (Cohen}, 65:20:12:3 (Tenny) and 85:10:4:1 (proposed ratio) by mechanical means of using mortar and pestle. The formulated composite propellants were characterized using Differential Scanning colorimeter (DSC), Scanning Electron Microscope (SEM) and Fourier Transform Infrared (FTIR). The FTIR and SEM results indicated some similarities of interaction with the individual components (the ammonium nitrate, biomass (Carbon), Sulphur, Starch and Gum Arabic (binders) displayed by the different peaks and functional groups indicating formation of composite solid propellant. From the DSC results, exothermic peaks range from 184-250°C while the endothermic peaks range from 40-168°C. thereby indicating the burning rate ability of the propellants in space.

Keywords: Composite, formulation, propellant, etc

1. Introduction  
Propellants are chemical as substances which burn at an extremely rapid rate producing large volumes of gases. Propellants are employed to initiate emergency seat ejection systems from aircrafts, used to accelerate a projectile in the bore of a gun and propel rockets and missiles along their flight paths, launch aircrafts into the air (Daintith, 2000). Propellants have existed for centuries in the form of black powder and later develop into smokeless (nitrocellulose) and then composite. These propellants exhibit different characteristics which challenge scientist to research and look into the behaviour of propellants, it is the quest for such that led to different research work on propellant characterization (Niklos, 2004).

Composite propellant, it contains mainly an oxidizer such as ammonium nitrate, ammonium perchlorate, chlorate mixed with fuel such as metal powder or carbon and also binders such as synthetic rubber, resins, polymeric binders. The composite propellant is more stable and easily kept for storage than the liquid propellant which is more cumbersome and difficult to control. It is mostly used for rocket purposes (Tenny, 1998)The major ingredients of modern military propellants are actually few. They consist of fuels, oxidizers and binders (polymers), and are fairly basic in their chemical nature and structure. The minor ingredients, used to assist or tie together the major ingredients, are more numerous and sometimes more complex. The interactions of these major and minor ingredients, when combined into a practical solid propellant, are complex. These interactions can take place at all stages of manufacture, storage, and use. The ingredients themselves represent output of the conventional chemical and explosive industries (although a few, like Nitroglycerin, are produced locally); however, the combination of these ingredients into a propellant is still thought of by some as a ‘black art.’ Making it into a science is what has justified the expenditure of so many millions of dollars over the past 50 years (Kevin, 1986). A powdered oxidizer and powdered metal fuel are intimately mixed and immobilized with a rubbery binder (that also acts as a fuel). Composite propellants are often either Ammonium Nitrate-based (ANC) or Ammonium Perchlorate-based (APCP). Ammonium Nitrate composite propellant often uses magnesium and/or Aluminium as fuel and delivers medium performance (Isp of about 210sec) whereas Ammonium Perchlorate Composite Propellant often uses aluminium fuel and delivers high performance (vacuum Isp up to 296sec with a single piece nozzle or 304sec with a high area ratio telescoping nozzle). Aluminium is used as fuel because it has a reasonable specific energy density, a high volumetric energy density, and is difficult to ignite accidentally. Composite propellants are cast, and retain their shape...
after the rubber binder, such as Hydroxyl-Terminated Polybutadiene (HTPB) Carboxyterminated Polybutadiene (CTPB) and Polybutadience Acylonitrile (PBAN).

2. Materials and Method

2.1. Sample Collection and Preparation

The maize stalk was obtained from 3 farms located along Kaduna- Zaria Road, Airport Road and Kaduna- Abuja Road while paw-paw stem was collected from 3 farms along Airport Road Kaduna, Kaduna- Abuja Road and along Kaduna- Kujama Road. The samples were collected in bags separately. The maize stalk and pawpaw stem were cut into smaller sizes, dried at room temperature and were labeled for charring labeled before charring. The gum Arabic was collected from acacia trees (acacia segalis) that produce the gummy sap using a knife which is located in Maigatari, Jigawa State while the starch flour was sourced from cassava tubers obtained from the open market. The collected cassava tubers were peeled and then washed to remove dirt. The washed cassava was then grinded into paste with a grinding machine and then kept in a bowl. The grinded cassava powder was sieved using a 20 micro size pores sieve into a clean bowl and allowed to settle for 3 hours, decanted and spread on a wide bowl to dry for 24 hours at room temperature. A fine dried powder was obtained and kept in a plastic container and labeled. The samples were coded as indicated in Table 1

2.2. Charring of Samples

The dried samples from, paw-paw stem and maize, stalks were charred to get the charcoal which constitutes mainly carbon (Biomass) out using an air tight Germany Muffle Furnace with Model Number SXL 1006 set at 400°C for one hour. The prepared samples of the plants were weighed(400g) each and inserted into the crucible having the size of the muffle furnace chamber for charring with a set temperature of 250°C for one hour. The obtained charcoal were then grinded to powdered form separately using mortar and pestle and labeled separately, kept under sealed containers.

2.3. Formation of Propellants

The formulation of the Composite propellants were done according to the ratios of Tenny (1998) (65:20:12:3), Cohen (1997)(80:16:2:2) and proposed (85:10:4:1) where exactly 32.5g ammonium nitrate, 10g of carbon, 6g of sulphur and 1.5g of starch were weighed and transferred into mortar and mixed thoroughly. The weights of the components were measured and mixed in such a way to achieve the Tenny ratio of 65:20:12:3 in 1997

The Same procedure was repeated with weights of components to achieve Cohen ratios of (80:16:2:2) and the proposed ratio of (85:10:4:1).

Exactly the same process was repeated for gum Arabic as binder.

2.4. Characterization of the Propellants

2.4.1. Differential Scanning Calorimeter (DCS)

DSC machine model DSC2 MERTNER TELEDO was used for thermal analysis of formulated composite propellants where 0.5g of the propellants were measured into crucibles and introduced into the sample holder compartment of the machine and the chamber was closed. The samples were run between 30-500°C for the analysis to obtain the transition phase, endothermic and exothermic values.

2.4.2. Scanning Electron Microscope (SEM)

Phenomenon PROX SEM machine with model number 4.5.3 was used to determined micro grain images of the formulated composite propellants. Samples were pulverized to particle size of about 0.15micron (100mesh) separately using Jaw Crusher. Exactly 0.5g of the pulverized sample was smeared on the stud housing the double adhesive carbon. The sample STUD was placed in the sample holder connected to the sample port and was introduced to the machine after which the chamber was closed. The sample was run for 5 seconds, and a prompt sound signified the sample was ready for imaging. The images were adjusted for sharpness, necessary zooming and refocusing and were then saved and printed.

2.4.3. Fourier Transform Infrared Spectrometer (FTIR)

The functional groups were identified using FTIR machine, the Agilent Carry FTIR Spectrometer (USA) with model number 630. Exactly 0.5g of the sample each was measured and compressed into a pellet in a fixed small cup like holder which was put on the crystal optical path and clicked on the software to process. The sample alignment was checked for proper sampling and the coding of the samples. The measurement was clicked and the peaks acquired over a range of 400 - 4000 cm⁻¹ using DTGS detector were selected for labeling by dragging to acquire the wave numbers as well as the transmittance. The generated and labeled peaks were saved and printed.
ated composite propellants were measured to 300g each and formed into the hallow tube
GA and CPAGA (Fig 4) indicated a DSC
ndothermic indicated a transition change in the samples which
, (2013) whose work was on ammonium perchlorate displaying similar DSC curves of first stage to be
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- transition state resulting to melting at

- curve of endothermic values of 130, 128 and 129
temperature and melting at the highest endothermic temperature before combustion at the high peak te mperature of exothermic value of 227.80,

- endothermic and th


2.5. Rocket Design
A PVC pipe was cut to the length of 300mm, with diameter of 48mm. in line with the rocket tube diameter system. The fin diameter for the rocket was obtained using aluminum sheet of 0.5mm thickness which was cut to a required measurement of 50 x 48mm.
A cone height of 50mm by 40mm was obtained by using Plaster of Paris (POP) which was molded using a mold to the required cone shape.
The propellant formulated using the charred plant biomass AMAS, AMAGA, BMAS, CMAGABPAS and CPAS and CCAS were used. The formulated composite propellants were measured to 300g each and formed into the hallow tube geometry (shaped by compressing them). This was made into 6 different geometric according to the samples ID selected. The selection was done based on the Differential Scanning Colorimeter research with the ones having higher heat values. After the compression, it was then allowed to dry at room temperature. The ignition system was designed using a cable with the two ends soldered to the filament containing an ignition charge which was made thicker and it then connected to a match composition and to a power source (battery) which allowed flow of current that ignited the propellants.

3. Result and Discussion

3.1. Physical Properties of Formulated and Charred Unformulated Samples
The unformulated charred samples obtained were solid black powder with porous white patches on their surfaces, dark brownish while the formulated samples obtained were homogenous powdered form, black greyish with white patches.

4. Chemical Characteristics
4.1. Differential Scanning Calorimeter (DSC)
The results shows that formulated propellant samples of AMAS, BMAS and CMAS (Fig 1) to have 128- 168- 170°C, 128-170 °C, and 129-170°C endothermic temperature values which indicates the phase change from the lower temperature to the highest endothermic temperature of 170 °C for the three different ratios of formulated propellants before combustion at the high peak temperature of exothermic value of 200.83, 192.37 and 197.54 °C for AMAS, BMAS and CMAS. DSC curve peaks for samples AMAGA, BMAGA and CMAGA in Fig 2 also displayed the first stage of the DSC curves of an endothermic values range of 170, 120-170 and 129-170 °C which indicates the transition state resulting to melting at the high endothermic temperature of 170 °C for the three samples respectively. The samples AMAGA, BMAGA and CMAGA displayed a DSC curve of an exothermic high peak value of 188.17, 193.35 and 209.52 °C, which indicated that decomposition has occurred resulting to combustion. All the values obtained were close to the values reported by Goncalves et al, (2013) whose work was on ammonium per chloride displaying similar DSC curves of first stage to be endothermic and the second stage to be exothermic with temperatures values of 246 °C and 365 °C respectively.
Propellants from APAS, BPAS and CPAS (Fig 3)displayed a DSC curve of endothermic of 82, 128.98 and 130- 164 °C. This temperature change in the first stage of the endothermic indicated a transition change in the samples which results in the melting before decomposing and combusting as shown in the DSC curve with exothermic value of 227.80, 223.93 and 213 °C for APAS, BPAS and CPAS respectively. Samples of APAGA, BPAGA and CPAGA (Fig 4) indicated a DSC curve of endothermic values of 130, 128 and 129-162 °C indicating the phase change gradually from the lower temperature and melting at the highest endothermic temperature before combustion at the high peak temperature of exothermic value of 208.16, 217.14 and 225.55 °C respectively for APAGA, BPAGA and CPAGA. All the values obtained were close to the values of Goncalves et al, (2013) displaying similar DSC curves of first stage to be endothermic and the second stage to be exothermic values of 246 °C and 365 °C respectively.

| S/N | Formulated Sample With Starch Binder Code (a) | Formulated Sample With Gum Arabic Binder Code (b) |
|-----|-----------------------------------------------|--------------------------------------------------|
| 1.  | AMAS                                          | AMAGA                                            |
| 2.  | BMAS                                          | BMAGA                                            |
| 3.  | CMAS                                          | CPAGA                                            |
| 4.  | APAS                                          | APAGA                                            |
| 5.  | BPAS                                          | BMAGA                                            |
| 6.  | CPAS                                          | CMAGA                                            |

Table 1: Sample Codes and Their Ratios

Key
A - Ratio - 65:20:12:3 (Tenny, 1998)
B - Ratio - 80:16:2:2 (Cohen, 1997)
C - Ratio - 85:10:4:1 (Proposed)
MA - Maize Stalk
S - Starch Binder
PA - Paw-Paw Stem
GA - Gum Arabic Binder
Figure 1: Showing the DSC Curve of AMAS, BMAS and CMAS
KEY: Blue: CMAS, Red: BMAS, Black: AMAS

Figure 2: Showing the DSC Curve of AMAGA, BMAGA and CMAGA
KEY: Blue: CMAGA, Red: BMAGA, Black: AMAGA

Figure 3: Showing the DSC Curve of APAS, BPAS and CPAS
KEY: Black: APAS, Blue: CPAS, Red: BPAS
4.2. Scanning Electron Microscope (SEM)

The SEM results of composite propellants formulated from maize stalk sample with codes (AMAS, BMAS, CMAS, AMAGA, BMAGA and CMAGA) revealed morphology of rhombic crater type, white uneven spherical, molten magma lump, flaky with trace like stripes and flaky uneven size particles as indicated in Fig 5 and Fig 6. The traces of white substance are shown to be embedded in the cluster indicating that the biomass had interacted with the sulphur and starch. The formulated sample dimension ranges from 65.4μm – 3.77μm. These images and dimensions observed were within the ranges of the study on analysis of low explosives in forensic investigation by Bonder (1998).

The formulated composite propellants from pawpaw stem with codes (APAS, BPAS, CPAS, APAGA, BPAGA and CPAGA) (Fig 8 and Fig 7) exhibited morphological characteristics of scraggy, honey comb shape, flake cake like, crater like with uneven holes and cracks and some white spheroidal substances and which appeared to be same for all the samples due to the same plant biomass.

The formulated sample dimensions ranges from 62.3 mm – 5.3 mm, which were observed to be similar to the range of values reported by Rodrigo et al (2005) in his study of characterization of a Hydroxy Terminated Rocket (HTPE) propellants.
Figure 6: SEM Image, Morphology of Samples AMAGA, BMAGA and CMAGA

Figure 7: SEM Image, Morphology of Samples APAGA, BPAGA and CPAGA
4.3. Fourier Transform Infrared (FTIR)

The results of the FTIR for maize stalk with codes AMAS, BMAS and CMAS (Fig 9) which represent different ratios A:B:C and AMAGA, BMAGA and CMAGA (Fig 10) did vibrational stretchings in the ranges of 3855.9 cm$^{-1}$ - 3609.9 cm$^{-1}$, 3205.5 cm$^{-1}$ - 2115.8 cm$^{-1}$, 1578.5 cm$^{-1}$ - 1299.0 cm$^{-1}$, 2107.8 cm$^{-1}$ - 2115.3 cm$^{-1}$. These values observed were as indicated in Fig 9. These spectra indicated the presence of alcohol, Esters, Nitro group, Amines, Isothiocyanate which showed the presence of AN, Carbon, Sulphur and binders used in homogenous mixture. This study agrees with the result of Anniyappan et al (2006) who detected Isothiocyanate at 1778.0 cm$^{-1}$, alcohol at 3384.2 cm$^{-1}$, Nitro group at 3354.2 cm$^{-1}$, C=O at 1882.5 cm$^{-1}$.

The pawpaw stem with codes APAS, BPAS and CPAS (Fig 11 representing different ratios of A;B;C and APAGA, BPAGA are CPAGA Fig 12) exhibited O-H, N-H, C=O, N=O, N=C=S and C≡C vibrational stretchings in the range of 3738.5 cm$^{-1}$ - 3216.7 cm$^{-1}$, 3213.0 cm$^{-1}$ - 2117.1 cm$^{-1}$, 1871.1 cm$^{-1}$, 1041.8 cm$^{-1}$, 1578.8 cm$^{-1}$ - 1300.8 cm$^{-1}$, 2120.9 cm$^{-1}$ - 1902.8 cm$^{-1}$ and 1757.3 cm$^{-1}$ - 1753.7 cm$^{-1}$. These spectra show the presence of Ester, Alcohol, Nitro group, Amines and Isothiocyanate. The Spectra values reported were within the range of spectra values observed by Cohen (1997) who detected Esters and Alcohol using tetrahydrobotane as binder.
Figure 9: Spectral of Composite Samples of AMAS, BMAS and CMAS

Figure 10: Spectral of Composite Samples of AMAGA, BMAGA and CMAGA
Figure 11: FTIR Spectral of Composite Sample of APAGA, BPAGA and CPAGA

Figure 12: Spectral of Composite Sample of APAS BPAS and CPAS
4.3.1. Rocket Testing

The designed rockets were tested at Research and Development Centre Field, Defence Industries Corporation of Nigeria (DICON) in order to evaluate the performances of the propellants. The revealed that propellants AMAS, CMAGA and CPAS did not lift off but burnt on the ground covering distances between 11m-60 m. However, propellants CPAS and BPAS projected distances of 280 m and 220 m respectively. High exothermic values and porosity of the surface area of the propellants giving good combustion accounted for this good performance.

5. Conclusion

The physical characterization of unformulated charred propellants revealed solid black powder with porous white patches, dark brownish and soft while the formulated were black grayish, homogenous powder. The FTIR result of the study revealed that all formulated propellants exhibited different vibrational stretchings such as N-H, O-H, C=O, N-O and C-H which indicated presence of Alcohols, Esters, Isothiocynate, Amines, Nitro groups, Alkenes among others. These vibrations were indicative of different functional groups thus confirming their presence in the composite propellants. SEM study of the formulated propellants shows morphology being predominantly crater, honey comb, scraggy, flaky, rhombic shape and fiber like stripe which were either large, small, dense spheroidal on the grain surface indicating composite formation. From the DSC results, the endothermic peaks range from 91 – 164 °C and exothermic range from 209 – 230 °C which indicates a transition change from lower temperature and melting at the highest temperature and thus indicating the burning ability of the propellants in space. The rocket testing revealed that propellants AMAS, CMAGA and CPAS did not lift off but burnt on the ground covering distances between 11m-60 m. However, propellants CPAS and BPAS projected distances of 280 m and 220 m respectively. The result therefore indicated that the propellants were effective in projecting the rockets.

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