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Determination of polymer content in energetic materials by FT-IR

Abstract: A new methodology was developed to characterize and to quantify the polymer content in PBX (HMX/Viton) by Fourier Transform Infrared Spectroscopy (FT-IR), using the Thermogravimetric analysis (TG) as reference techniques for the quantitative method. The quantification methodology, proposed by us, using the Fourier transform infrared-attenuated total reflectance (FT-IR/ATR) showed excellent results, being faster than the usual methodologies and can eliminate the generation of chemical residues.

Keywords: Explosives, HMX, FT-IR, TG, ATR, Viton quantification.

INTRODUCTION

The development of space programs, deep oil well drillings, etc., has projected a need for “heat resistant” or “thermally stable explosives”. The main objective for using such explosives or explosive formulations is to support systems or applications, which must be reliable and secure at high temperatures (Agrawal, 2005).

An appropriate military explosive needs to be secure and of easy manipulation. It can be stored over long periods of time in different climates and difficult to detonation, at least under specific circumstances (Mathieu, 2004). The development of plastic bonded explosives (PBX) is a way to get more energy in less volume (Hayden, 2005).

PBX is an acronym for “Plastic Bonded Explosive”. A term applied for a variety of explosive mixtures, which is characterized by high mechanic resistance, good explosive properties excellent chemical stability, low sensitivity to the shock and manipulation and low sensitivity to thermal initiation (Federoff, 1966). A lot of the PBX’s have been developed since the middle of the last century, in general, resulting in an engineering material for specific or large scale applications (Hayden, 2005; Kim, 1999). These explosive mixtures have a large content of secondary explosives such as cyclotrimethylene trinitramine (RDX), cyclotetramethylene tetranitramine (HMX), hexanitrostilbene (HNS) or pentaerythritol-tetranitrate (PETN) in a composition with a polymeric matrix such as polyester, polyurethane, nylon, polystyrene, some rubbers, nitrocellulose and fluor polymers. The

compatibility and the miscibility with other explosives, polymers, and additives must be evaluated before pressing, molding, extruding, etc., for manipulation safety and for storing the products (Mathieu, 2005; Hayden, 2005).

As the weapons can be placed in aggressive thermal and mechanical environments, it is important to characterize the properties of the PBX in order to know its physico-chemical behavior (Thompson, 2005; Kasprzyk, 1999).

A method to obtain an explosive load is by pressing the explosive covered by hydraulic press, which represents the most important process for high-performance explosive loading manufacture (Wanninger, 1996). As the explosive is obtained in crystal form, it is necessary to recover them with polymers, performed by a covering process.

Different methods employing several techniques have been developed to determine HMX contents (Mattos-a, 2004). High performance liquid chromatography (HPLC) is used to investigate explosives to a large extent (Mattos, 2004). Although the HPLC method presents good results, there are several steps related to the sample preparation or measurements that are quite time consuming. It is possible to apply the Thermogravimetric Analysis for the quantitative determination of different energetic compounds in explosive compositions, as reported by Silva et al. (2008). FT-IR techniques can be applied for the characterization of the polymeric composition in PBX (Mattos, 2008). In a previous publication (Mattos, 2008 and Mattos, 2009), a sample of HMX covered with Viton in different concentrations was analyzed by ATR technique to obtain good quality spectra. In that specific case, the sample was kept in perfect physical contact

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with the crystal surface (Mattos-b, 2004). Several steps are eliminated and although the data of ATR methods represent the average of five analyses, the time spent in the analysis is less than the time spent in the HPLC method, as demonstrated in another publication (Mattos, 2009).

The covering process and the characterization of energy materials in the crystal form or after getting covered by different polymers are important research studies performed in the Institute of Aeronautics and Space (IAE) laboratories (Mattos-b, 2004; Mattos, 2002; Mattos, 2003). Thus, the aim of this work is to show a new quantitative method for the determination of the HMX/Viton ratio based on FT-IR/ATR, using the TG data like reference.

The advantages of the methods here presented are the few steps involved and the fact they do not require complex sample preparation.

EXPERIMENTAL

Materials

Fluoroelastomers are generically referred to as FKM polymers (nomenclature per ASTM D1418). Viton is a trademark of a fluoroelastomers series manufactured by DuPont, which is available in several formulations (copolymer, terpolymer) and forms (slab, stick, pellet) (Mattos, 2008; Hohmann, 2000). Fluoroelastomers are used in the covering of energetic materials. Viton shows excellent chemical stability, being a widely used polymer for energetic materials coating of propellants. The relationship between Viton and the oxidant compounds has significant influence in the oxidation time, as cited in the literature (Hohmann, 2000).

HMX is also known as 1,3,5,7-tetranitro-1,3,5,7-tetrazacyclo-octane, cycloteramethylene tetranitramine ($C_4H_8N_8O_8$) or octogen. It is an important ingredient in modern solid propellants due to its desirable properties, such as absence of smoke, high specific impulse and thermal stability (Tang, 1999). It is used in certain propellants and explosives (Kohno, 1994). Propellants based on HMX can be found in armament applications and in solid rocket propulsion systems (Tang, 1999).

The specific gravity of the crystals is 1.90 g/cm^3 and the melting point is around 280°C . HMX exists in 4 polymorphic forms (α HMX, β HMX, δ HMX, γ HMX). The most common modification, stable at room temperature, is β HMX (United States-TM 9, 1979).

The HMX has been used in metal conformation, energy transference and in the composition of melting and pressing plastic bonded explosives (Calzia, 1969; Urbanski, 1984).

The plastic covering of explosive grains for “slurry” in water is characterized by the migration of the explosive, where the transfer of the explosive occurs from one liquid phase to another, in a liquid system with two immiscible phases (James, 1965; Benziger, 1973; Kneisl, 2003).

In the covering of HMX crystals by polymers, the explosive is dispersed in water (inorganic liquid phase) and the polymer dissolved in an organic solvent (organic liquid phase), the organic phase is usually called “lacquer”, which is a plastic dissolved in a low boiling point organic solvent (James, 1965; Benziger, 1973; Kneisl, 2003).

The covering process starts with the dispersion of the energetic material in water (system with moderate agitation and heating). The hot lacquer is added to the system, due to the low solubility of the lacquer in the system, the agglutination of the plastic plates occurs in the explosive crystals (James, 1965; Benziger, 1973; Kneisl, 2003).

The organic solvent is removed by means of the distillation process and then the formation of “pellets” occurs, due to the agglutination between the polymer and the explosive crystals (James, 1965; Benziger, 1973; Kneisl, 2003). It is important to mention that the inorganic liquid phase (water) is very important for the security process, during the warming step of the system with explosive not yet covered and in the distillation step. It enables good temperature homogenization, preventing the system’s overheating (Kim, 1999; James, 1965; Benziger, 1973; Kneisl, 2003).

Measuring methods

Thermogravimetric analyses were carried out in a TG/SDTA Mettler model (Model TGA/SDTA851e), which was calibrated in a heat ratio of $10^\circ\text{C}/\text{min}$, with aluminum and indium, under nitrogen gas flow ($40 \text{ mL}/\text{min}$). About 2.0 mg of the sample was used and the heating range was between 50 and 700°C .

The FT-IR spectra were recorded using a Spectrum One PerkinElmer. The conditions were: resolution 4 cm^{-1} ; gain 1; range 4000 to 700 cm^{-1} , 40 scans, germanium crystal as reference material and incidence angle of 45° . The samples were analyzed by ATR germanium techniques. The sample was placed on both sides of the crystal to obtain a better spectrum.

RESULTS AND DISCUSSION

PBX samples with Viton and HMX was prepared (HMX/Viton systems) with the following Viton content: 2, 5, 8, 10, 13 and 15%, in order to develop a quantitative methodology FT-IR.

TG Analysis of HMX

HMX (class 3) was used to attain the polymeric coverings, obtained in the SNPE (Société Nationale des Poudres et Explosifs), which shows the following thermal behavior in Fig. 1.

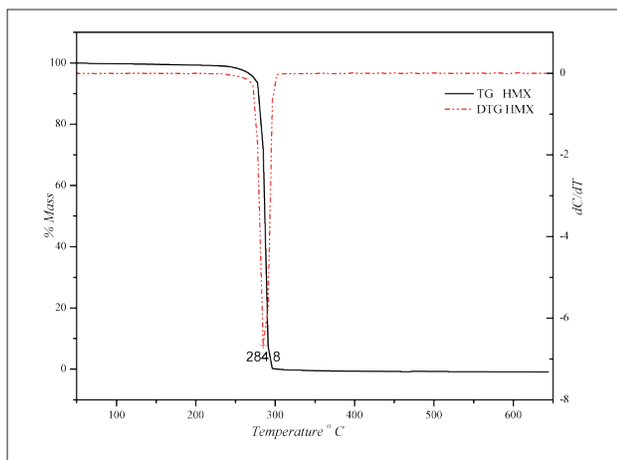


Figure 1: TG and DTG curves of HMX sample in atmosphere of N₂

It can be observed that the HMX undergoes a sudden thermal decomposition and maximum mass loss occurs at around 285°C in relation to the DTG.

TG Analysis of VITON

To analyze the thermal behavior of Viton B, TG analyses were conducted. The thermal curve in Fig. 2 illustrates the thermal behavior of this polymer.

Observing Fig. 2 in comparison with Fig. 1, it is evident that the Viton decomposition takes place at a very high temperature of the HMX, allowing for the quantification of both materials, when these are in the PBX. The Viton mass loss in relation to the DTG can be observed at the temperature of 487.8 °C.

PBX characterization of PBX (HMX/Viton) by TG analyses

The samples cited in Table 1 for PBX (HMX/Viton) were analyzed by TG and Table 1 shows the results obtained. However the covering process with higher contents was not so efficient, according to Table 1, for samples 06/06 and 02/06.

Figure 3 shows the thermal decomposition curves of PBX (HMX/Viton).

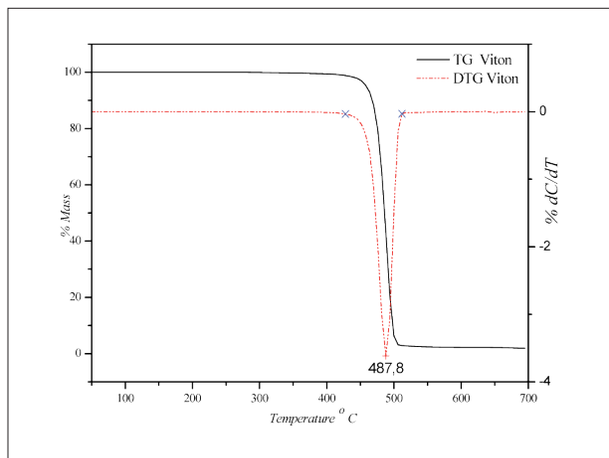


Figure 2: TG and DTG curves of Viton B sample in atmosphere of N₂

Table 1: HMX contents in the HMX/Viton samples obtained by TG

Code	Theoretical HMX/Viton content (%)	HMX contents by TG (%)	Viton contents by TG (%)	Standard deviation (%)
01/06	98/2	97.87	2.13	0.68
01/95	95/5	95.24	4.76	0.25
05/06	92/8	93.67	6.33	0.93
04/06	90/10	91.10	8.90	1.06
06/06	87/13	89.88	10.12	0.79
02/06	85/15	89.98	10.02	1.06

The Figure 3 clearly shows that the decomposition stages of the PBX constituent are quite different and do not influence the decomposition process. The contents

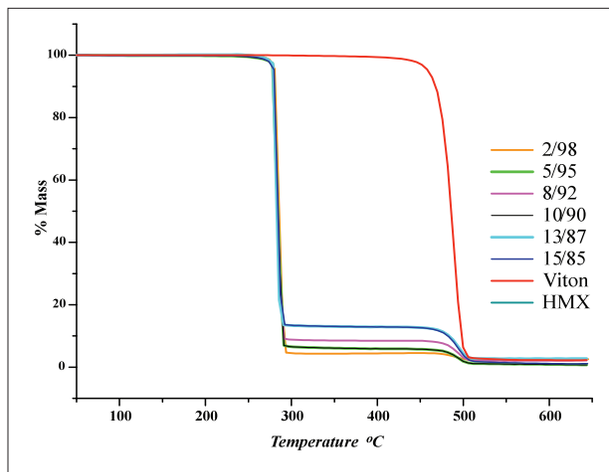


Figure 3: TG Curves of the system HMX/Viton sample in atmosphere of N₂

of the constituents can then be determined by means of TG, when associating the nominal content with the percentage decomposition of each stage of mass loss observed.

FT-IR/ATR of HMX sample

The main absorptions observed in the HMX spectrum FT-IR (Fig. 4), associates a probable attribution (Smith, 1979; Litch, 1970; Hummel, 1981; Bedard, 1962) are at 3035 cm^{-1} ($\nu\text{ CH}_2$), 1564 cm^{-1} ($\nu_a\text{ NO}_2$), 1462 , 1432 , 1396 and 1347 ($\delta_s\text{ CH}_2$), 1279 and 1202 cm^{-1} ($\nu_s\text{ NO}_2 + \nu\text{ N-N}$), 1145 , 1087 and 964 cm^{-1} ($\nu\text{ N-N} + \nu\text{ ring}$), 946 cm^{-1} (ring stretching), 830 and 761 cm^{-1} (δ and $\gamma\text{ NO}_2$). The observed bands are characteristic of β HMX, according to the literature (Smith, 1979; Campbell, 2000; Achuthan, 1990).

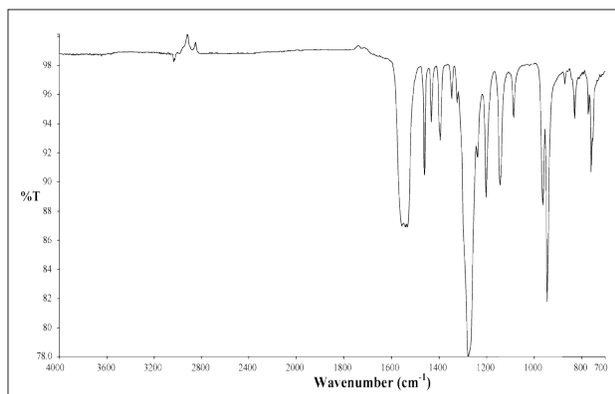


Figure 4: FT-IR/ATR spectrum in crystal germanium of HMX

FT-IR/ATR of Viton sample

The presence of C-F groups is detected in Mid-Infrared (MIR) (Mattos, 2008; Urbanski, 1977) on the basis of the intense absorptions in $1397 - 1074\text{ cm}^{-1}$ ($\nu\text{C-F}$) (Mattos, 2008; Smith, 1979). The characteristic bands of linking C-F_2 are found in 1273 , 1191 , 1134 and 1111 cm^{-1} ($\nu\text{C-F}_2$) and 890 and 820 cm^{-1} ($\nu\text{C-F}_3$) (Mattos, 2008; Silverstein, 1981), of average intensity.

The ATR/FT-IR technique can be conducted with two different crystals (germanium and KRS-5). The ATR technique with germanium crystal (sample in both sides of the crystal - 45°) shows good results, where it is possible to see the changes in the characteristic bands of the polymer ($\nu\text{C-F}$, $\nu\text{C-F}_2$ and $\nu\text{C-F}_3$) as a function of the polymer contents in the sample and good definition in the base line.

Figure 5 illustrates the characteristic spectrum for the Viton B sample.

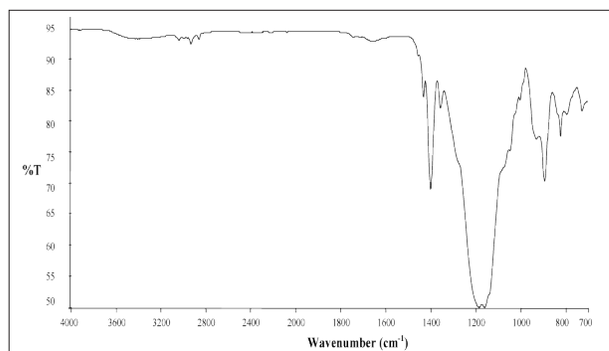


Figure 5: FT-IR/ATR spectrum in crystal germanium of Viton B

ATR analysis of the HMX/Viton

The combination of the infrared spectroscopy with the reflection theories produced advances in the surface analysis. The FT-IR/ATR technique combines the power of IR spectroscopy with the optics of the attenuated total reflection. The concept of internal reflection spectroscopy originates from the fact that radiation propagating in an optically dense medium of refractive index undergoes total internal reflection at an interface of an adjacent medium of lower optical density. This wave is termed evanescent (Nogueira, 2000; Kwan 1998; PerkinElmer 2005).

This evanescent wave penetrates only a few microns ($0,5\mu - 5\mu$) beyond the crystal surface and into the sample. Consequently, there must be good contact between the sample and the surface of the crystal. In the infrared spectrum regions, where the sample absorbs the energy, the evanescent wave will be attenuated or altered. The attenuated energy from each evanescent wave is passed back to the IR beam, which presents the opposite end of the crystal and is passed to the detector in the IR spectrometer. The system then generates an infrared spectrum (PerkinElmer, 2005).

By ATR technique with germanium crystal, using sample in both sides of the crystal, characteristic spectra were obtained for the Viton and the PBX, as can be observed in the sets of spectra in Fig. 6.

The samples of the HMX/Viton system were analyzed using different FT-IR techniques. However, it was observed that using the FT-IR/ATR technique with Germanium crystal at 45° showed spectra with better resolution (Mattos, 2009).

As observed in the spectra of Figure 6, the spectrum of the HMX/Viton (85/15%) sample presents characteristic bands of Viton in 1169 cm^{-1} and 890 cm^{-1} . In this way, the best analytical band for the Viton is chosen.

All the covering samples (HMX/Viton) were analyzed by ATR with Germanium crystal at 45° . Therefore, with

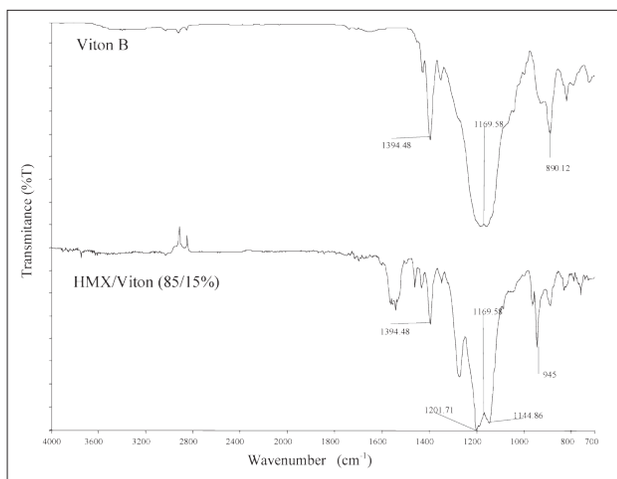


Figure 6: FT-IR/ATR spectra using germanium crystal: Viton B and Cob. 02/06-85/15%

this technique better results were obtained for indicating the analytical band and quantitative determination, as represented in Fig. 7.

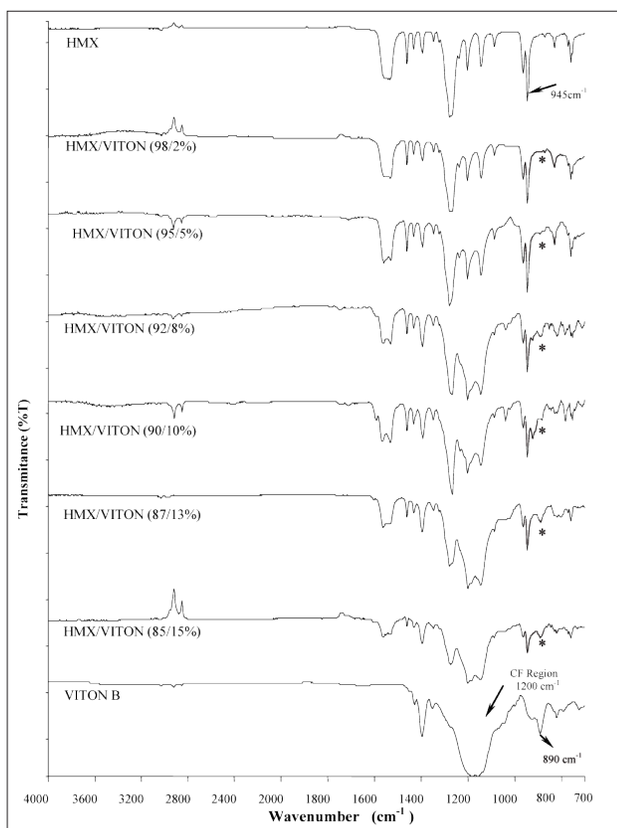


Figure 7: FT-IR/ATR spectra for HMX/Viton using germanium crystal with sample in both sides of the crystal at 45°

According to the set of absorption spectra FT-IR/ATR in Fig. 7, characteristic bands for the HMX and Viton can be observed. The band at approximately 1200 cm⁻¹ is the characteristic groupings C-F existing in the Viton. It can be

observed that while the Viton concentration increases in the mixture, there is a widening in this region, indicating the presence of polymer in the energetic material composition of HMX/Viton. The band at 945 cm⁻¹ characteristic for the HMX is also evidenced in this set of spectra. And while the Viton content increases, it is observed the reduction in the intensity of HMX band due the presence of the polymer.

Other characteristic Viton bands were chosen at 890 cm⁻¹ and 1170 cm⁻¹. For the HMX, the bands in 945 cm⁻¹ and 1145 cm⁻¹ can be observed. Since there are Viton bands that increase the intensity with the polymer concentration in the system, it was possible to initiate the quantification of these samples for FT-IR.

TG was used as reference for IR methodology in this system, (Table 1). Next, analytical IR bands for determining the Viton percentage in the HMX/Viton system were analyzed. The base line chosen for the bands at 945 and 890 cm⁻¹ was: 986 at 853 cm⁻¹, and for the bands of 1145 and 1170 cm⁻¹: 1476 at 984 cm⁻¹.

The absorbance values represent the median (μ) of five analysis. According to Hórak (1978), it is recommended to work with median numbers when working with a low number of experimental values. It may happen that the values of parameters $\hat{\mu}$ and $\hat{\sigma}$ thus determined are subject to larger errors. These errors are difficult to determine due to a non-uniform distribution of random errors in the set. Therefore, a difference assessment is made. The standard deviation, $\hat{\sigma}_{\mu}$ of the median absorbance is calculated as follows [Hórak, 1978]: equation

$$\hat{\sigma}_{\mu} = \frac{\hat{\sigma}}{\sqrt{n}} \tag{1}$$

where $\hat{\sigma}$ is the assessed standard deviation of the basic set and is a quantitative measure of the precision for each individual measurement; n is the number of experiments.

$$\hat{\sigma} = K_R \cdot R \tag{2}$$

where K_R is the coefficient for the calculation of the average standard deviation from the variation range (for five experiments, $K_R = 0.430$) (Hórak, 1978); and R is the difference between the largest and the smallest element ($X_n - X_1$). $\hat{\sigma}_{\mu}$ is an evaluation of the precision of this median, that is, an evaluation of the result obtained from the treatment of a finite set of measurements, which are repeated under completely identical conditions. The relative error for each sample was determined as follows:

$$relative\ error\ (\%) = \left(\frac{\hat{\sigma}_{\mu}}{\mu} \right) \times 100 \tag{3}$$

Table 2 shows the median absorbance values for analytical bands available.

Table 2: Median absorbance values for analytical bands of the HMX/Viton system

HMX/Viton (%)	Median A_{945} HMX	Median A_{1145} HMX	Median A_{890} Viton	Median A_{1170} Viton
98/2	0.060	0.032	0.002	0.008
95/5	0.041	0.032	0.005	0.017
92/8	0.027	0.042	0.006	0.035
90/10	0.034	0.040	0.007	0.032
87/13	0.030	0.049	0.010	0.044
85/15	0.031	0.064	0.014	0.059

Analyzing Table 2, it is observed that the values of the bands at 890 and 1170 cm^{-1} must obey a linear relation. With the data from the termogravimetric analyses (reference method) and relative to the FT-IR analysis for Viton characteristic bands at 890 cm^{-1} and 1170 cm^{-1} , it is possible to have the following linear relations for the content determination of Viton in agreement with the HMX/Viton, using relative bands method.

Viton content by TG/FT-IR using relative band

It is known that the relative bands method (relative absorbance) can be used for thickness correction of the specimens (Gedeon, 1985). Thus, for quantitative FT-IR analysis in HMX/Viton samples, the absorbance values at 945 cm^{-1} (HMX) were related to those corresponding to the absorption at 890 cm^{-1} (Viton), in an attempt to evaluate whether there would be an improvement in the precision of the methodology developed. With regards to the points of the base line at 986 cm^{-1} , the 853 cm^{-1} was established for the calculations of absorbance values for the vibration absorption of the ring 945 cm^{-1} and C-F band at 890 cm^{-1} . To apply the Lambert-Beer law in this analysis, the relative absorbance data of A_{945}/A_{890} was placed as a function of explosive/plastic composition (percentage in weight), as presented in Table 3.

Figure 8 shows the relative graph of the medium values of A_{945}/A_{890} as a function of the relation of HMX/Viton concentration (%/m); a good linear relation is obtained ($R=0.993$). From the analytical curve obtained by FT-IR/TG analysis (Table 3), the following relation is proposed;

$$y = -3.95 + 0.7x \tag{4}$$

Table 3: FT-IR data (relative band A_{945}/A_{890}) PBX samples of (HMX/Viton)-reference: data TG

HMX/Viton Content by TG	Relative Band A_{945}/A_{890} (median)	Standard deviation ($\hat{\sigma}$)	Relative error (%)
45.95	28.50	6.50	22.81
20.01	9.60	1.11	11.56
14.72	4.28	0.15	3.50
10.24	4.62	0.16	3.46
8.88	3.00	0.12	4.00
8.98	2.21	0.07	3.17

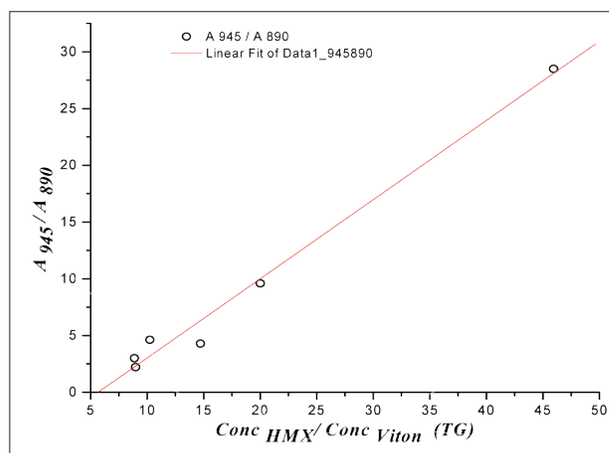


Figure 8: Medians values of the relative absorbance A_{945}/A_{890} as a function of HMX/Viton content obtained by TG.

Table 4: FT-IR data (relative band A_{1145}/A_{1170}) of the PBX samples (HMX/Viton)-reference: data TG

HMX/Viton Content by TG	Relative Band A_{1145}/A_{1170} (median)	Standard deviation ($\hat{\sigma}$)	Relative error (%)
45.95	4.000	0.500	12.50
20.01	1.947	0.036	1.85
14.72	1.206	0.005	0.41
10.24	1.289	0.016	1.24
8.88	1.120	0.007	0.62
8.98	1.071	0.005	0.47

where y is the median value of the relative absorbance (A_{945}/A_{890}) as a function of the relative concentration of HMX/Viton.

Figure 9 shows the relative graph for the median values of A_{1145}/A_{1170} as a function of the relation of HMX/Viton concentration (%m/m); a good linear relation is obtained ($R=0.991$). From the analytical curve obtained by FT-IR/TG analysis (Table 4), the following relation is proposed;

$$y = 0.344 + 0.0789x \quad (5)$$

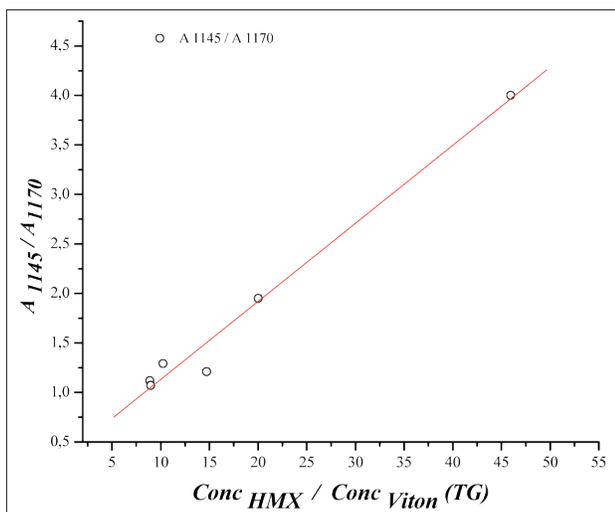


Figure 9: Medians values of the relative absorbance A_{1145}/A_{1170} as a function of HMX/Viton content obtained by TG.

where y is the median value of the relative absorbance (A_{1145}/A_{1170}) as a function of the relative concentration of HMX/Viton.

With the analytical curves obtained (IR/TG) for the different IR bands used (relative band) the concentrations of Viton were calculated. The values of the linear correlation coefficients (R) were evaluated and are presented in Table 5.

Table 5: Viton contents by TG and FT-IR technique (relative bands at 1145/ 1170 cm^{-1} and 945/ 890 cm^{-1})

Viton content By TG analysis (%)	Viton content IR relative band 1145/1170 cm^{-1} TG reference (%)	Viton content IR relative band 945/890 cm^{-1} TG reference (%)
2.13	2.11	2.10
4.76	4.68	4.90
6.33	8.36	7.82
8.90	7.71	7.53
10.12	9.24	9.13
10.02	9.82	10.18
Linear Correlation	$R = 0.991$	$R = 0.993$

The data in Table 5 showed that the use of the relative band was the best adjusted for Viton content determination by FT-IR for PBX (HMX/Viton), presenting excellent results, that is, better coefficients of linear correlation in relation to the obtained values of Table 5. The best relative band was of 945/890 cm^{-1} using the TG technique as a reference technique, with a linear correlation of 0.993, and furthermore it's possible to better visualize the bands in the spectra in Fig.7.

CONCLUSIONS

ATR technique with germanium crystal where the sample is placed in both sides of the crystal (45°) showed a better and more detailed spectra where it is possible to see the changes in the characteristic bands of polymer as a function of the content in the explosive mixture and a more defined base line;

All the covering samples of the HMX/Viton system obtained in laboratory were analyzed by ATR technique with Germanium crystal (45°), therefore they show better results for indicating the analytical band and quantitative determination;

The increase in intensity of the Viton bands with the increase in the polymer content in the system, enabled to initiate the quantification of these samples by FT-IR;

The use of a relative band showed to be adjusted for determining the Viton content by FT-IR for PBX (HMX/Viton), presenting excellent results. The best relative band was 945/890 cm^{-1} , using the TG technique as a reference technique;

The use of the new FT-IR methodology for polymer quantification in PBX showed to be faster without generating chemical residues.

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