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Comparison of New and Legacy TATBs[†]

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<u>Abstract</u>

Two newly synthesized versions of the insensitive high explosive (IHE) 1,3,5-triamino-2,4,6-trinitrobenzenes (TATBs) were compared to two legacy explosives currently used by the Department of Energy. Except for thermal analysis, small scale safety tests could not distinguish between the different synthetic routes. Morphologies of new TATBs were less faceted and more spherical. The particle size distribution of one new material was similar to legacy TATBs, but the other was very fine. Densities and submicron structure of the new TATBs were also significantly different from the legacy explosives. Pressed pellets of the new explosives were less dense. Recrystallization from sulfolane improved the density and thermal stability of both new TATBs, though the morphology of the recrystallized TATB was nearly hexagonal platelets.

Introduction

Modern Department of Energy (DOE) weapons systems gain enhanced safety from plastic bonded explosives (PBXs) which contain the insensitive high explosive (IHE) 1,3,5-triamino-2,4,6-trinitrobenzene (TATB) bonded together with

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Kel-F 800, a copolymer of vinylidene fluoride and chlorotrifluoroethylene.

Currently LLNL has limited wet-aminated TATB reserves for formulation.

Although TATB is being synthesized and may soon be commercially available, the synthesis processes have changed and the explosive must be re-evaluated. Legacy wet and dry-aminated TATB have not been synthesized in the US in any significant quantity since about 1985. Furthermore, significant quantities of the PBXs based on TATB/Kel-F (LX-17 and PBX 9502) have not been formulated since about 1990.¹ Over the last few years as part of a DOD MANTECH, Thiokol² and Holston Army Ammunition Plant (HAAP)³ have produced moderate quantities of TATB (~5 kg batches) with plans to scale up for DOD applications. Thiokol TATB is polycrystalline with an average particle size of about 40 μm (similar to WA TATB) but HAAP TATB is only 5-6 μm (similar to ultrafine). Small quantities of these materials were obtained from the Naval Air Warfare Center at China Lake. CA⁴ for evaluation.

Characterization of the four TATBs: the new BAE Holston and ATK
Thiokol and legacy wet (WA) and dry (DA) aminated TATB which are no longer
manufactured in the US is the subject of this paper. Small scale safety data on
these TATBs are unable to distinguish between them except that newer TATBs
decompose thermally about 10°C below the decomposition temperature of dry
aminated TATB which is about 5°C below that of wet aminated. The morphology
of these TATBs was characterized by scanning electron microscopy. The density
of compression molded disks approached the average density of the TATB

crystals with increasing pressure. Ultra small angle x-ray scattering (USAX) measurements on pressed pellets showed different interfacial characteristics for ATK TATB compared to the others. Density distributions of new TATB crystals were substantially less than old crystals as measured using a gradient column. It may be possible to remove some of the impurities and improve the crystal structure of these TATBs by recrystallization. To that end small scale recyrstallization of BAE and ATK TATB from sulfolane produced crystals with density and thermal decomposition temperatures comparable to wet aminated TATB.

Experimental:

Legacy wet and dry aminated TATB used in these experiments was originally synthesized by Aerojet Corporation, sold to Holston Army Ammunition Plant and given lot numbers 12-03-82-0324-273 and 13-03-85-0307-538, respectively. Sample quantities of these TATB's were obtained by LLNL and given the lot numbers C-090 and C-084. Five hundred grams each of three different lots of Thiokol (ATK) and Holston (BAE) TATB were graciously provided by the Naval Surface Warfare Center at China Lake, CA. All of the tests described in this paper used ATK lot TATB001 and BAE lot BAE6E295-001, hereafter referred to as C-562 or C-559, respectively.

All explosives used at LLNL are subjected to small-scale safety tests including impact sensitivity, friction, spark and two thermal stability tests known as

chemical reactivity (CRT) and differential scanning calorimetry (DSC) tests prior to other evaluation or use in prototype devices. A thorough critique of various small-scale tests is given elsewhere. ^{5,6}

Morphological comparisons of the various TATBs were made using a LEO 438 VP (variable pressure) scanning electron microscopy (SEM). The variable pressure SEM reduces surface charging of an organic sample by allowing a low-pressure atmosphere to pass over the sample to bleed off excess charge. Particle size distributions were determined using the Saturn DigiSizer 5200 laser light scattering apparatus manufactured by Micromeritics Instrument Corporation. Samples of approximately 15 mg of TATB in 2-ml DI water were dispersed with Coulter 1A surfactant and 3 minutes of sonication in an external sonicator. The morphology of sulfolane recrystallized TATBs was evaluated using a Zeiss 40-A polarizing microscope under crossed polars with a first order red plate. Optical micrographs were calibrated against photographs of a stage micrometer at the same magnification with Axiovision software. Further description of the equipment used in the morphological studies can be found in the manufacturer instruction manuals.^{7,8,9}

A Techne model DC-2 density gradient column was used for all crystal density measurements. The column was immersed in a water bath with temperature controlled to 23 ± 0.05 °C. The manufacture claims density can be estimated to 0.0001 g/cc with this equipment. The gradient was achieved using slow mixing

of a high density aqueous ZnBr₂ solution (approximately 2.00 g/cc) into a low density solution initially about 1.84 g/cc which was fed into the gradient column. Calibrated glass density beads were added to the column and the height of the center of each bead was measured to within 0.1 cm. A least squares calibration curve for the column was generated from the bead position and its density. Approximately 0.001 gram of TATB was immersed in the low density liquid to wet out the crystals and added to the top of the column with an eye dropper. The crystals were allowed to sink overnight to reach equilibrium density prior to counting. TATB particles within each gradation on the column were counted and the density calculated from the least squares fit of the calibration beads. Several thousand particles were counted in the gradient column. Further description of this apparatus and its application to explosive crystals is given elsewhere. ^{11, 12}

Approximately 0.19 grams of TATB was compression molded using a Black
Hawk press into 1.27 cm diameter disks approximately 0.08 cm high at pressures
between 35 – 200 MPa (5,000-30,000 psi) pressure at ambient for 1 minute and
released. Pellet densities were calculated from dimensional measurements and
weight. The surface of the pressed pellet was examined under epi illumination
with differential interference contrast (DIC) inserts in the Zeiss 40-A polarizing
microscope. ⁹

Results and Discussion

Small scale safety data

Small scale safety data on the four different types of TATB are given in Table 1. A discussion of these safety tests has been presented elsewhere. 5, 6 As can be seen from the data, all of these explosives produce no reaction in impact, spark and friction tests. In Figure 1, the DSC traces of the new TATBs decompose at 10-15°C lower temperatures than the older TATBs. Wet aminated TATB has the highest decomposition peak temperature (386.9°C), with an onset of about 353°C. This synthetic route produces high purity TATB. 13 Dry aminated TATB decomposed about 5-8°C lower (378.3°C peak temperature). It is known that DA TATB contains about 0.5% residual NH₄Cl from the synthesis process. 14-15 Scanning electron microscopy reveals micron size voids where this salt has been extracted from the crystal during washing, but it is currently unclear whether this or smaller size or impurities such as residual NH₄Cl is responsible for the reduction in thermal decomposition temperature. The ATK sample had a second peak at about 352.7°C and the main decomposition peak at 368.4°C. This is significantly lower than their reported peak temperature of 378.1°C.5 Thiokol reports about 96% purity for their early TATB production which has been improved to 98% in more recent syntheses. BAE TATB reports purity of 98.5-99.1% and decomposition peak temperatures of 364-368°C, consistent with results given here.⁶ They also report 99.9% pure TATB synthesized at their Bridgewater plant in the UK had decomposition peaks between 389-381°C, again consistent with our wet aminated result. We have found that ultrafine TATB produced by colloidal milling wet aminated TATB shows a reduction of about 5-7°C in the decomposition peak (around 379-380°C), not enough to account for

the 20°C reduction in the new BAE TATB. Although several kinetic studies of TATB with DSC¹⁶⁻¹⁸ have tried to estimate activation energies for thermal decomposition, they contain sparse information on the effects of particle size or impurities. It would appear that there is a relationship between purity, particle size and decomposition temperature, but further study would be required to determine what it is. The effect of particle size on thermal decomposition has also been seen in recent one dimensional time to explosion (ODTX) tests. ¹⁹

Morphologies of TATBs

TATB morphologies were evaluated by scanning electron microscopy (SEM). As can be seen in Figure 2, SEM's of the various TATB's have very different morphologies. Both new TATBs in Fig. 2b and d are spherical aggregates. The ATK TATB large aggregates in Fig 2b averaged about 77 \pm 16 μ m and were made up of smaller spherical particles about 28 \pm 8 μ m which are in turn made up of smaller crystalline aggregates of about 3 \pm 3 μ m. BAE TATB in Fig 2d was much finer, but also composed of spherical aggregates about 5 μ m across composed of submicron particles of about 0.7 \pm 0.3 μ m. In Fig. 2a, WA TATB shows its characteristic polycrystalline structure, with crystallite sizes near 30-60 μ m, along with fines on the order of 5-10 μ m. Facets are often observed and steps on the surface have been seen in WA TATB. The classic DA TATB morphology is shown in Figure 2c. ^{20,21} Facets of the triclinic unit cell can be seen peppered with holes where the ammonium chloride has been extracted during the last washing cycle.

Density distributions of TATBs

Densities of TATB were measured using the density gradient technique. 22-24 Theoretical maximum density (TMD) of TATB is 1.937 g/cc.²⁵ Density distributions of dry and wet aminated TATB in Figure 3 were almost identical with the mean density of wet aminated being 1.9173 g/cc or about 98.98% TMD and dry aminated mean density of 1.9181 g/cc or about 99.02% TMD. It is known¹ that dry aminated TATB contains approximately 0.5% ammonium chloride from its synthesis route compared to about 0.1% for wet. Since the density of ammonium chloride is about 1.53, the void volume of dry aminated TATB may be higher than the simplistic estimate given here. Looking at the SEM micrograph of dry aminated TATB in Figure 2c would seem to indicate that the 1-2 μm holes which cover the surface of the dry aminated crystals can be wetted and penetrated by the density gradient fluid. Assuming all the voids to be air would imply a void content of about 1%. The Thiokol TATB density distribution maintained the shape of the wet and dry aminated distributions, but was shifted to lower density. This may imply that the mechanism of void formation is similar in these 3 samples. The average density (1.8355 g/cc or only 94.75% TMD) was much lower than the legacy TATBs. It is known that ATK and BAE TATBs contain on the order of 2% impurities.^{2, 3} Although the density of these impurities is not known, it is expected to be comparable or higher than ammonium chloride. Again the estimate of percent voids would probably be higher than the simple difference approximation (5.3%). The very fine BAE particle size made the density distribution difficult to measure. As can be seen in Figure 3, a very broad

distribution was found for this TATB with a mean density of 1.8010 g/cc. Near 1.91 g/cc density a group of large particles was observed, above this density until about 1.78 g/cc fine particles increased in number and below 1.78 g/cc a cloud of very fine particles was observed up to nearly the top of the column (1.765 g/cc). The number of particles in the cloud was estimated by counting a region and assuming similar regions contained similar number of particles.

Particle size distributions

Light scattering measurements of particle size distribution showed the new TATB's have more fines than WA or DA as shown in Figure 4. The samples of BAE TATB were finer than UF by almost a factor of 2, cf, 2.2 μ m compared to 3.8 μ m average diameter for ultrafine TATB ground in a Sturtevant micronizer. ^{26, 27} Perhaps the spherical aggregates of ATK break down during sonication to produce the extra fines, but the mean particle size was 56 μ m, only slightly smaller than B-090 dry aminated TATB (70 μ m). Wet aminated TATB had an average particle size of approximately 48 μ m. So although the morphology is drastically different, ATK is a reasonable particle size match for current stockpile TATB's. BAE claims to have the ability to produce multiple sizes², similar to the classes of HMX and RDX but the sample obtained from NAWS at China Lake was very fine and might light better than UF. The relative frequencies of the distributions for ultrafine and BAE TATBs in Figure 5 clearly show the bimodal nature of these materials.

Characterization of sub-micron structure:

Ultra small angle x-ray scattering (USAXS) probes structures from ~10 nm to ~2 microns. ²⁸ Figure 6 presents USAXS from wet-aminated (WA), dry aminated (DA), Holsten (BAE) and Thiokol (ATK) TATB from small disks pressed at 30,000 psi. Although pressed samples presented here and, to a greater extent, the powders showed multiple scattering, qualitatively the data shows a number of interesting characteristics. Scattering occurs due to electron density differences, and thus cannot distinguish between scattering from particles or voids. In these cases, the calculated volume contributing to the scattering calculated using the invariant²⁹ is about 5% and 4% for wet and dry aminated, 13% for the Holsten and 4% for the Thiokol TATB. With the exception of the Thiokol, these are much higher than the void volume calculated by density, so we must assume the observed scattering occurs from both voids and particles.

The Holsten and Thiokol samples have a much larger population of small scatterers (less than ~ 50 nm) than the wet- and dry- aminated as seen by the more intense scattering intensity at high scattering angle (larger than about 10⁻² / Å). For the Holsten TATB, this is consistent with the smaller crystallites seen in the SEM micrographs of Figure 2d. The surface-area-to-volume ratio calculated from the USAXS is also about one order of magnitude larger in both the Holsten and Thiokol compared to the wet- and dry-aminated TATB.

The clearest scattering difference between the Thiokol and the other TATB samples is the power-law exponent annotated in Fig. 6. For an ideal, two phase

system (i.e. TATB and air) with three-dimensional scatterers, the exponent approaches -4.^{29, 30} Wet-aminated (-3.89 to -3.94) dry-aminated (-3.85 to -3.90) and even the Holsten (-3.97) are very close to this value, indicating an abrupt interface between TATB and air, consistent with crystalline facets at the interface. Thiokol TATB, however, has an exponent of -3.42. Although different power-law exponents are associated with reduced dimensionality, the exponent seen in the Thiokol case most likely arises from a diffuse or rough fractal-like surface interface.²⁹

Pressing characteristics

The density, ρ , of ~0.2 g disks of TATB was evaluated as a function of pressing pressure, P, between 35 < P < 210 MPa (5,000 to 30,000 psi) for a single, one minute dwell at ambient. For zero pressure the bulk density of the sample was used. As molding pressure is increased, the part density should approach the theoretical maximum for the density, ρ_T , of the TATB crystal (1.937 g/cc). The percentage of ρ_T is:

% TMD =
$$100\rho/\rho_T$$
 (1)

Assuming only air is trapped internally and its contribution to the density is negligible, the void volume fraction, $\phi(v)$, would be 1- ρ/ρ_T . A schematic diagram of the process is shown as an insert in Figure 7. The part density as a function of ram pressure for a single 1 minute dwell was highest at 200 MPa (30 ksi) in all

cases. WA and DA TATB could be pressed to nearly theoretical maximum density (TMD), 99.3 and 99.7%, respectively. ATK TATB could be pressed only to 94.9% TMD and the fine powder BAE TATB would only press to 90.9%. This may be due to the spherical structure of these TATBs and the smaller particle size of BAE. Ultrafine TATB (ground) also does not press to very high density. ²⁴ Table 2 lists the densities and void volume fraction of the different TATBs.

Each pressed pellet surface was examined in reflection using Differential Interference Contrast (DIC) microscopy. In some instances it is possible to distinguish inter-crystalline interfaces due to differences in refractive indices between crystals. ³¹⁻³³ Photomicrographs, shown in Figure 8, were taken at the highest magnification (~1000x) and the 20 μm bar in the lower left micrograph is representative for all photos. At the highest pressures, usually TATB crystals are no longer identifiable due to surface damage and flow. At lower pressures an attempt was made to measure crystal dimensions where such structures were observable.

In Figure 7, the density dependence of wet aminated TATB with increasing molding pressure was sigmoidal. The DIC micrographs of the pressed pellet at low pressure, first row first column in Figure 8, show a rough interface. Since the depth of field is low, out of focus regions at low pressure represent depressions or high spots on the surface. The area of the out of focus parts of the first micrograph in Figure 8 (35 MPa) under WA consists of about 12%. This is not

very close to the 24% voids estimated from the density difference at this pressure. This would seem to imply that the defects are mainly interstitial crystal packing problems and are fairly large at low mold pressures. At higher pressures plastic deformation allows the polycrystalline aggregates in WA TATB to deform and fill the interstitial voids. The WA crystals on the surface begin to coalesce at about 140 MPa (15 ksi). Prior to coalescence, particle size on the surface does not seem to change significantly running between 15 < dia. < 35 μ m. This is about 25% smaller that the average particle size of the unpressed WA TATB consistent with crystalline fracture and particle size reduction observed in compression molded PBXs. ³⁴

In Figure 7, the density dependence of dry aminated (DA) TATB with molding pressure increased asymptotically toward TMD. Judging by the density, the larger crystals of DA TATB packed better than WA TATB at lower pressure although in the high pressure limit the densities were indistinguishable (1.933 and 1.924 for DA and WA TATB respectively). These DA crystals withstood the pressure better than WA based on nearly constant crystal dimensions with increasing pressure (compare Fig 8, column 1 with column 2). Some of the DIC images at low molding pressure showed voids on crystal faces parallel to the surface which were on the order of 1-3 μ m similar in size to the voids observed in the DA TATB SEM in Figure 2c. The out of focus surface area of DA TATB pressed at 35 MPa, 12.6 %, compared well with 16 % void volume based on density measurements.

The BAE TATB has very low bulk density (0.472 g/cc) compared to all of the other TATBs (0.8 to 0.9 g/cc). The pressing characteristics are shown in Figure 7. Low pressure increased the density by 3x but with increasing pressure only 91% TMD was obtained, the lowest of all pressed pellets. The small particles of the original BAE TATB sample are only observed in DIC micrographs of the disk surface at the lowest pressure (see Figure 8 column 3). Above 35 MPa (5 ksi) only the marks of the ram surface and large aggregates of unknown origin were observed. These aggregates are not seen at all pressures, but could be found in different regions of the surface above 35 MPa (5 ksi). Comparison of the surface to volume void content for BAE TATB was not possible because of the small size of the TATB particles. This material cannot be the so called "dirty binder", 35, 36 since no binder was used to prepare these samples and the idea of rubbelized particles being generated seems less likely with these fine particles. With pressure, as expected, these small primary particles seemed to compact well, compared to substantial plastic flow in the TATBs with larger crystals at higher pressures. Densities, unfortunately, do not improve beyond 91% at maximum pressure.

Pressing characteristics of ATK TATB are similar to WA as shown in Figure 7.

The spherical particles seen in the SEM in Figure 2c are also evident on the pellet surface at low to moderate molding pressure in Figure 8. At about 20 ksi, the marks of the ram surface are seen in the pellet. The internal structures of the

spherical particles appear as the pressure increased. Estimating the out of focus area gave 17.3% compared to density void volume of 17.0%, the best comparison of all samples.

Improved thermal stability by recrystallization.

The thermal stability and the density distribution of these new TATBs can be improved significantly by rapid recrystallization in sulfolane. Previous attempts at improving crystal quality have used a variety of solvents including dimethyl sulfoxide, diphenyl ether, solfolane, ^{37, 38} nitrobenzene²⁵, 1,2 dichlorobenzene, benzonitrile, dimethyl formamide and chlorobenzene.³⁹ Recently in an attempt to make fine TATB particles superacids methane and ethane sulfonic acid and hexamethylphosportriamide were found to dissolve TATB. 40 Sulfolane was selected because of its relatively rapid dissolution of TATB at 190°C and sharp and rapid precipitation characteristics on lowering the temperature to 175°C. Also a high degree of recovery of TATB was observed for crystallization under these conditions. The new TATB was suspended in sulfolane (1.2% w/v) and rapidly heated with stirring for 4 min when dissolution occurred (~190 °C). The heat source was removed and the resulting solution cooled and stirred with crystallization commencing at ~ 175°C. The crystalline suspension was stirred (30 min) and then filtered (~ 30 °C). The recrystallized TATB was washed with

acetone, water and acetone and dried under vacuum. Yellow, crystalline TATB recoveries varying from 89 to 100% have been obtained.

Polarized optical micrographs of the sulfolane recrystallized TATB from the BAE sample are shown in Figure 9. These large, triclinic crystals averaged 83 ± 17 μm normal to the [001] axis and many contained what appear to be etch pits or inclusions about 3.4 ± 0.6 μm across. It would seem that one of the BAE impurities from their synthesis route tends to act as a nucleating agent and remain included in the crystal. The peak decomposition temperature in the DSC for these crystals in Figure 11 increased by 13°C compared to the original BAE TATB in Figure 2. The bimodal character of the exotherm is often observed in TATB 17 and has been attributed to an autocatalytic first step in the decomposition of TATB, but may reflect the influence of impurities on the thermal decomposition.

The ATK recrystallized crystals in Figure 10 averaged $120 \pm 40 \, \mu m$ with lower levels of inclusions than were found in the BAE crystals and what appear to be dislocations in nicely formed triclinic crystals. Again the DSC decomposition temperature increased after recrystallization. In this case, the decomposition peak was 0.6° C higher than the legacy WA sample, also shown in the figure. Further optimization of a short residence time recrystallization procedure would seem to be of interest since it seems to lend itself to scale up, but was not

pursued here. Average density of the recrystallized ATK TATB increased from 1.835 g/cc to 1.9140 g/cc, comparable to wet and dry aminated densities.

Conclusions

Significant differences between new and legacy TATBs were found in this study. Although the particle size distribution as measured by light scattering for WA, DA and ATK TATBs was similar, the morphology, thermal stability, density, and submicron structure were significantly different. BAE TATB had much smaller particles than the legacy TATBs. How this will affect the formulation and performance of DOE plastic bonded explosives (PBXs) such as LX-17 and PBX 9502 prepared from new TATBs compared to legacy material is currently under study, but results are not expected to be similar to the extensive data on current PBXs. Different morphologies and surface characteristics are expected to alter processing characteristics in the Holston slurry coating reactor. Molding powder size, bulk density, and granularity may be affected. Pressing characteristics and mechanical properties are expected to change. Lower densities of the new TATBs are expected to alter initiation characteristics, probably making the PBXs easier to initiate. Reduced thermal stability may alter cook off characteristics. Experiments are ongoing or planned to address some of these questions.

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<u>Tables</u>

Table 1. Small scale safety test results show all synthetic routes produce insensitive HE.

TATB	DH 50	Friction	Spark	DSC (Ton)	Tpk
WA	>177.4 cm	>36.0 Kg	NS	353	386.9
DA	>177.4 cm	>36.0 Kg	NS	348	378.3
ATK	>177.4 cm	>36.0 Kg	NS	285	352.7; 368.4
BAE	>177.4 cm	>36.0 Kg	NS	310	367.3

Table 2. Density, ρ and void volume, $\phi(v)$, of 0.2-g disks of TATB as a function of pressing pressure, P.

P\TATB	ρ(DA)	φ(v)	ρ(WA)	φ(v)	ρ(ATK)	φ(v)	ρ(ΒΑΕ)	φ(v)
0 MPa	0.973	50	0.773	60	0.968	50	0.472	76
35.2	1.629	16	1.473	24	1.617	17	1.475	24
66.9	1.799	7.1	1.524	21	1.662	14	1.634	16
106.2	1.886	2.6	1.772	8.5	1.793	7.4	1.706	12
137.2	1.888	2.5	1.824	5.8	1.832	5.4	1.743	10
208.2	1.933	0.21	1.924	0.67	1.842	5.1	1.761	9.1

Figures

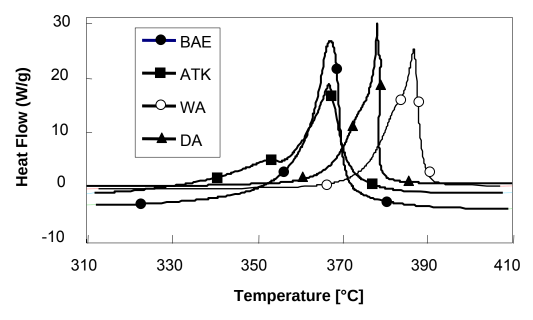


Figure 1. Differential scanning calorimetry (DSC) traces from the four synthetic routes show lower decomposition temperatures for the newer TATBs apparently due to impurities.

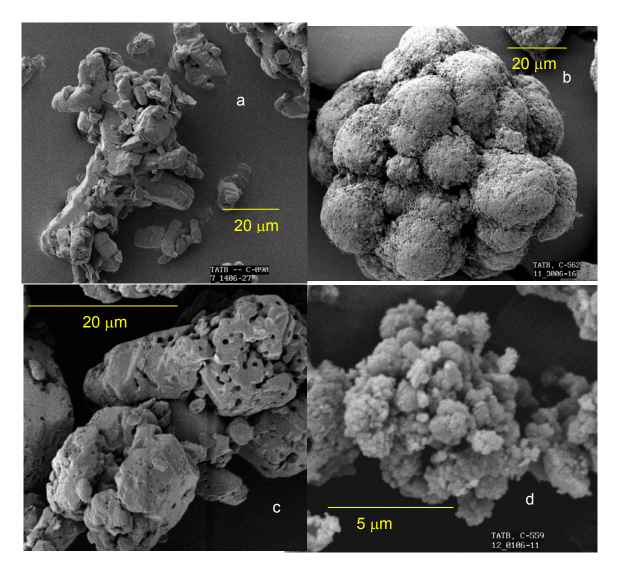


Figure 2. SEMs of (a)Wet aminated TATB; (b) ATK Thiokol TATB (c) Dry aminated TATB and (d) BAE Holston TATB.

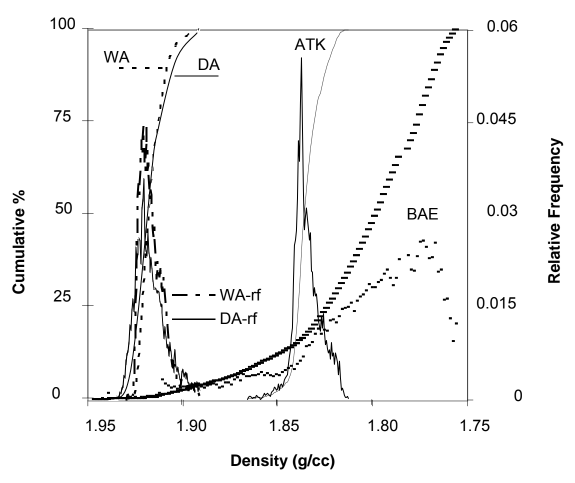


Figure 3. Density distributions in new TATB was much lower than in original wet and dry aminated TATB, probably due to impurities and voids.

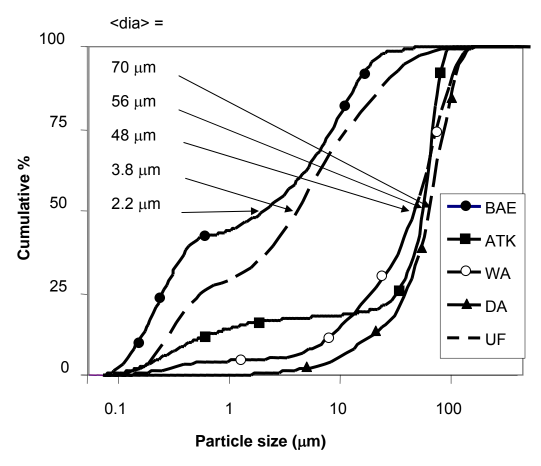


Figure 4. ATK, wet and dry aminated TATB particle size distribution were fairly similar, but BAE TATB was finer than ultrafine TATB made in a Sturtevant mill.

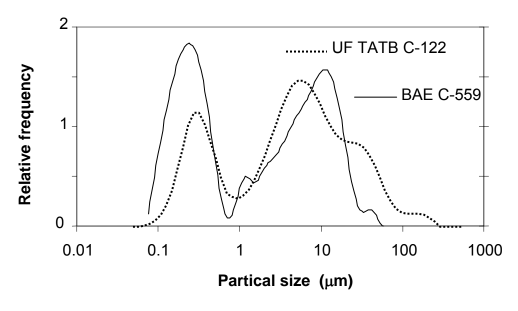


Figure 5. Relative frequency of particles of BAE and ultra fine TATB show the bimodal characteristics of these materials.

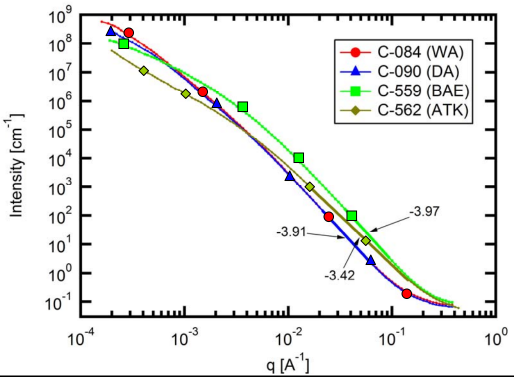


Figure 6. Comparison of ultra small angle x-ray scattering of WA (red), DA (blue), BAE (green) and ATK (yellow) TATBs from pressed disks shows substantial differences between manufacturers.

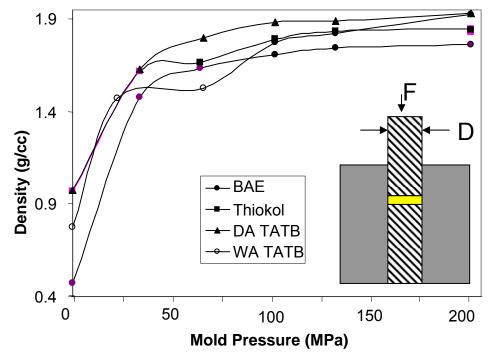


Figure 7. Density as a function of pressure for single dwell ambient pellet of TATB increased slowly as the pressure increased above 5000 psi.

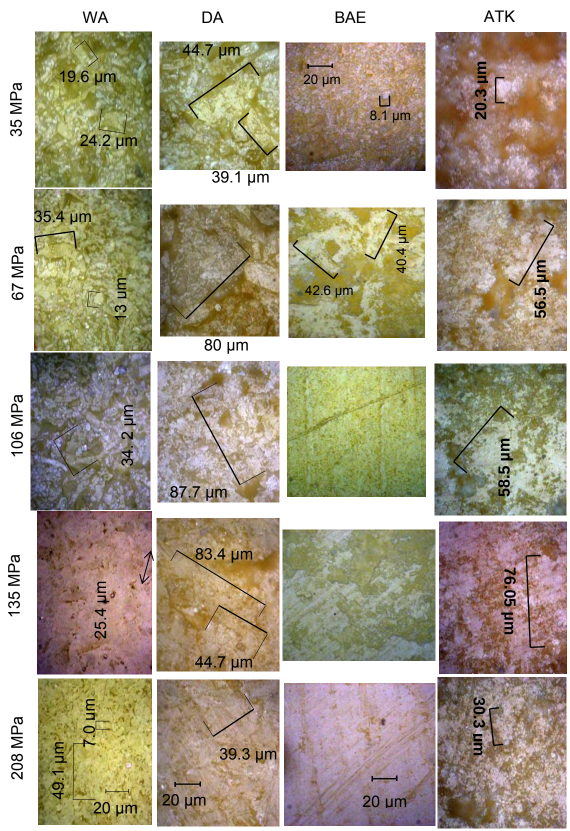


Figure 8. Differential interference contrast micrographs of the surfaces of the various TATBs show increasing plastic deformation with pressure.

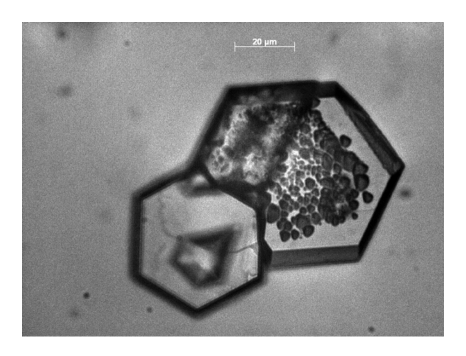


Figure 9. BAE TATB recrystallized in sulfolane showed etch pits or inclusions in many crystals.

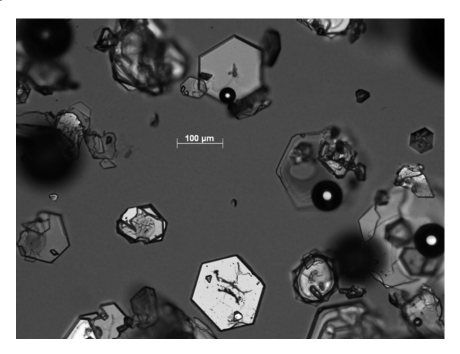


Figure 10. ATK TATB recrystallized from sulfolane produced these large platelets.

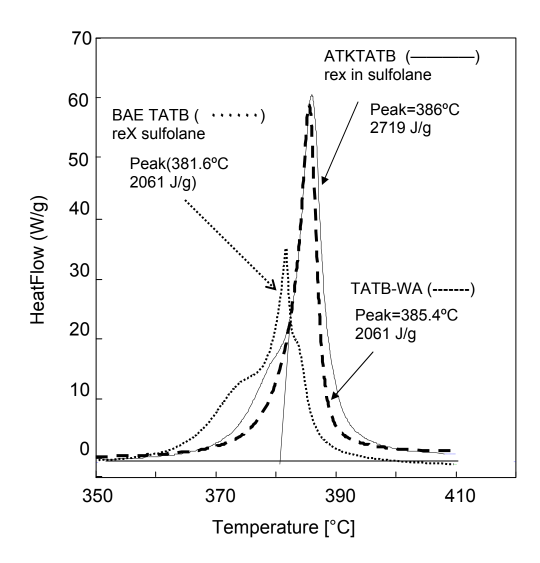


Figure 11. Thermal decomposition of BAE (dotted line) and ATK (solid line) TATBs recrystallized from sulfolane compared to wet aminated TATB (dashed line) showed increased peak temperature compared to original samples.