DEVELOPMENT OF THERMALLY STABLE POLYBENZIMIDAZOLE (PBI) FIBER

KENNETH R. SIDMAN

JOHN B. GREGORY

TECHNICAL REPORT ASD-TR-72-50



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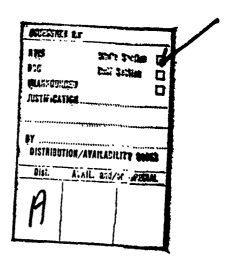
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FOREWORD

This report was prepared by Dynatech R/D Company under Contract No. F33657-70-C-1156 to the Aeronautical Systems Division, Life Support System Program Office, Endeavor Number AO117, Wright-Patterson Air Force Base, Ohio. The work was administered by the Nonmetallic Materials Division, Air Force Materials Laboratory, Directorate of Laboratories. Mr. Robert M. Stanton and Mr. Stanley Schulman were project engineers.

This report covers work conducted from June 8, 1970 to November 16, 1971. Dynatech Report Number: 1022.

The program was directed by Mr. Ralph L. Wentworth. Technical supervisor of the program was Mr. Kenneth R. Sidman. Physical testing and polymer evaluation was supervised by Mr. John B. Gregory. Assisting in the evaluation of candidate reagent systems and the set-up of the fabric treatment facility were Mr. Glenn K. Armstrong, Dr. Shafik E. Sadek, and Mr. William H. Crandell. Consultants to the project team were Prof. P. L. T. Brian and Prof. J. Baldwin.

Publication of this report does not constitute Air Force approval of the report's findings or conclusions. It is published only for the exchange and simulation of ideas.

JACK H. ROSS, Chief Fibrous Materials Branch Nonmetallic Materials Division ALBERT P. LOVELADY, Colonel, USAF System Program Director Life Support System Program Office Deputy for Subsystems UNCLASSIFIED

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garment in more intimate contact with the wes	arer. Elimin	ation of the	tendency to shrink
would reduce the rate of heat transfer to a we	arer, prolong	ing the peri	iod of protection in a fire
exposure situation. The work reported here	vas undertake	n to develo	p a treatment for PBI
which stabilizes it with respect to thermal shi			
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basic polymer chains of PBI was found to stat			
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5% chlorosulfonic acid in phosphorous oxychlo	ride followed	by heating	and neutralization was
found to be satisfactory. Fabric so treated is	resistant to	thermal shi	rinkage yet unchanged in
strength and nonflammability. The treatment			
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adequacy of this fabric, a pilot processing lin			
treated on this pilot line was converted into fa			as constructed. 1 D. to
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age of PBI fiber has been developed. It is re-			
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properties of treated material.			

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ABSTRACT

Textiles composed of polybenzimidazole (PBI) fibers combine superior qualities of nonflammability and comfort with functional utility. These characteristics qualify PBI fabric especially well for service in Air Force garments such as flyer's suits and coveralls. A further improvement desired in the qualities of PBI fabric is reduction of its tendency to shrink when exposed to flame and high temperature.

Such shrinkage may bring the fabric of a garment in more intimate contact with the wearer, thus increasing the rate of heat transfer to the individual from a flame source. Elimination of the tendency to shrink would reduce the rate of heat transfer to a wearer, prolonging the period of protection in a fire exposure situation. The work reported here was undertaken to develop a treatment for PBI which stabilizes it with respect to thermal shrinkage.

Chemical treatment of PBI fiber or fabric which introduces chemical bonds between the basic polymer chains of PBI was found to stabilize PBI against thermal shrinkage. The effectiveness of a number of such treating agents and the coincident effects on other fiber properties were determined. A treatment involving immersion of PBI in a liquid mixture of 5% chlorosulfonic acid in phosphorous oxychloride followed by heating and neutralization was found to be satisfactory. Fabric so treated is resistant to thermal shrinkage yet unchanged in strength and nonflammability. The treatment is stable to laundering.

Woven PBI fabric was treated by the process developed. Following demonstration of the adequacy of this fabric, a pilot processing line for treating PBI tow was constructed. PBI tow treated on this pilot line was converted into fabric for evaluation.

A functionally acceptable process for reducing the thermal shrinkage of PBI fiber has been developed. It is recommended that further investigation of the tow treatment process be conducted in order to attain improved balance of the color and fabric properties of treated material.

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Section 1

SUMMARY

Polybenzimidazole (PBI) fibrous structures exhibit stability to high temprature oxidation, excellent textile physical properties, and high moisture regain. However, when subjected to flames PBI fabrics tend to shrink. Work at Dynatech R/D Company under contract for NASA-MSC improved both the thermal and oxidative resistance and dimensional stability of PBI fabric. An optimization of the vapor treatment was undertaken for the Aeronautical Systems Division, Life Support System Program Office ASWL, Wright-Patterson Air Force Base.

Reagent systems other than the vapor phase treatment of PBI with phosphoryl chloride proved to be superior, and treatment of fabric with 5% chlorosulfonic acid in phosphorus oxychloride proved to be outstanding. Seven yards of fabric were treated by this process and submitted to the Air Force Material Laboratory for evaluation. This fabric met the target requirements: exhibiting less than 10% thermal shrinkage in a test of exposure for seven seconds to a controlled Meker burner flame, less than 10% shrinkage on treatment, no reduction in tensile strength, and flammability essentially the same as the untreated fabric. The treatment was stable to prolonged rinsing in alkaline solutions and washing in alkaline soap solutions.

In Phase II of the program the chlorosulfonic acid treatment was converted to a continuous process to treat PBI tow. A pilot processing line capable of treating approximately one pound per hour of PBI tow was constructed. Process variables were adjusted on this line unit a combination of tow properties including low thermal shrinkage, good color, and high tensile strength was achieved. Thirty pounds of treated tow were prepared and converted into fabric for evaluation.

It is recommended that development work on the treatment of tow by this process be extended to include the investigation of other parameters which could not be examined under the scope of this program. It is anticipated that the material and specifications developed in such an investigation will enable the attainment of the balance of color and fabric properties necessary for Air Force textile material.

Section 2

INTRODUCTION

2.1 Background

As a result of the interest of the Department of the Air Force in developing a chemical treatment of polybenzimidazole (PBI - specifically, 2, 2' (m-phenylene) -5, - (6, 6' -bibenzimidazole)) fibrous structures to effect a superior nonflammable, thermally stable, and nonshrinking material, Contract No. F33657-70-C-1156 was awarded to Dynatech R/D Company. The primary emphasis of this work was to be the optimization of vapor phase treatment of PBI with phosphoryl chloride to effect a thermal stabilization of the fiber. A two phase investigation was proposed - the first phase consisting of the research and development of the treatment method and the second concerned with scale-up of the process to treat quantities of material.

2.1.1 Scope

As the investigation progressed, it became apparent that the phosphoryl chloride treatment was not exceptionally effective in stabilizing fibrous structures of the polymer. Consequently, the study was broadened to include the investigation and optimization of other reagent systems that provide thermally stable crosslinks between polymer chains. The crosslinks were intended to restrict the polymer chain mobility so that the highly strained, anisotropic chain configuration produced by the fiber drawing would not relax when heated.

2.1.2 Technical Discussion

Polybenzimidazole fibers have tremendous potential value in the manufacture of nonflammable clothing – they exhibit the flexibility, drape, surface characteristics, moisture regain and physical properties known to determine comfort. In addition, they are self-extinguishing in environments containing less than 35% by volume oxygen (for fabric conditioned at 70°F and 65% relative humidity). The present PBI fibers, however, exhibit thermal shrinkage in the brief exposure, high intensity heat of the open pit flame test. This shrinkage is apparently due to a relaxation of the polymer chains.

Two methods present themselves for minimizing thermal shrinkage. One involves preshrinking the fabric through chemical or heat treatment so that the resultant cloth exhibits a very low thermal shrinkage. The other involves reducing the mobility of the polymer chains.

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The PBI fibers may be preshrunk through the use of chemical agents able to swell and plasticize the polymer. In this case the swelling agent enhances chain mobility and permits the polymer molecules to reorient themselves to relieve the strain. The same relaxation may be achieved by heating the fibers. In either case, a decrease in tensile strength occurs because the anisotropic, strained configuration contributing to the fiber strength is lost.

To reduce chain mobility, crosslinks may be introduced along the polymer chain in sufficient number so that these bonds will connect the individual polymer molecules into a three-dimensional network. Such a network greatly restricts the fiber shrinkage. In this approach the tensile strength of the fiber is retained.

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Crosslinks may be introduced into the fiber thermally or chemically. Thermal crosslinking of certain polybenzimidazoles was described by Plummer and Marvel (Ref. 1). They found that heating the polymers under vacuum at temperatures exceeding 300°C tended to change the yellowish compounds to a brownish-black form that was insoluble in all solvents. The conversion of the aromatic rings to diphenyl groups was discussed as a possible crosslinking mechanism. In oxygen-containing atmospheres, temperature above 350°C have been reported to induce crosslinking. In these cases also, a reduction in polymer solubility was taken as evidence of crosslinking behavior. Shulman and Lochte reported (Ref. 2) that at pyrolysis temperatures between 450°C and 500°C crosslinking of PBI occurs via amide interchange or bisphenyl coupling.

Chemically, crosslinks may be introduced through the incorporation of any moiety on the polymer chain that will bridge across chains. The 1- and 3-positions of the benzimidazole ring and the aromatic moieties are particularly susceptible to attack. Consequently, multifunctional agents selected on the basis of their reactivity with these sites, offer promise as potential crosslinking agents.

The site-numbering system is as follows:

From a review of the chemistry of benzimidazoles (Ref. 3), it is possible to identify a number of candidate reactions. The imidazole ring, for example, may be attacked by phosphorylating, silanating, chlorinating, and methylolating agents.

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These agents may crosslink through imidazole rings. For example,

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Oxidizing, sulfonating, nitrating, acylating, and free-radical generating agents have been shown to attack aromatic nuclei. For example,

Free-Radical Generation

Crosslinking may then occur either between aromatic rings

or between an aromatic and an imidazole ring.

The effectiveness of the crosslinking is measured with respect to improvements in fabric dimensional stability while at the same time maintaining desirable textile properties.

2.2 Method of Approach

In Phase I of the program two experimental approaches were undertaken. The first involved an extension of the POCl₃-oxygen, vapor-phase treatment developed for NASA-MSC under Contract NAS 9-8179. The second approach consisted of a search for and optimization of crosslinking agents which dimensionally stabilize PBI fibrous structures.

Treatments were evaluated first for effectiveness in reducing the thermal shrinkage of fabric samples. If shrinkage behavior was found to be improved then the durability of the treatments to laundering was evaluated. If they passed these two examinations, the treated fabrics were examined for color, hand and drape, flammability, and moisture regain.

The goal of this first phase of the contract was to produce a treated fabric with less than 20% thermal shrinkage by the end-use application test of the Air Force Ma' rials Laboratory, less than 30% fabric shrinkage on treatment, less than 25% reduction in tensile strength, and flammability essentially the same as the untreated fabric.

In Phase II of the program the feasibility of the process developed in Phase I was evaluated for treating PBI in the form of fabric, yarn, staple, fiber, and spinning dope. The advantages and disadvantages of treatment in the various forms are summarized in Table I. On the basis of this analysis, it was decided to treat PBI tow.

A tow treatment facility was established on which the process variables were adjusted to give the best combination of low thermal shrinkage, good color, and high tensile strength.

Thirty pounds of PBI were then processed to provide a sufficient quantity of treated cloth for end-item tests in full scale open-pit flame trials.

Table I

The Advantages and Disadvantages of Treating PBI in Various Forms

ine Adv	antages and Disadvantages of Treating	PBI in various Forms
PBI Form Treated	Advantages	Disadvantages
	- Uniform treatment of PBI ensured regardless of form of end item and end item precursors.	- Changes in PBI reaction conditions, spinning conditions, and drawing conditions would be required
Spun and drawn fiber	- Uniform treatment of PBI ensured-independent of end item.	- Changes in fiber handling and processing equipment may be required to adjust for changes
	- No change required in present PBI processing techniques.	in surface characteristics and denier of fiber
	- Applicable to proposed 1-2 million lb/yr facility producing fiber and tow.	
Staple	- Fiber may be processed through to staple with no change	 Not applicable to continuous filament processing.
	in equipment settings or equipment. Batch processing possible.	 Changes in staple handling and processing equipment may be required.
Yarn	- PBI may be processed through to staple with no change in equipment setting or	- Changes in present yarn handling and processing equip-ment may be required.
	equipment.	 Changes in chlorosulfonic acid process reaction conditions will be required depending on weight of yarn.
Fabric	- Fiber may be processed into end-item-no dislocation of any PBI processing step is necessary.	 Chlorosulfonic acid process would have to be modified for various weaves, weights, and widths of fabric.

Section 3

PROCEDURES AND RESULTS

3.1 Phase I - Experimental

The experimental fabric treatments discussed in this report were conducted in four types of reactors - one constructed of stainless steel, two of glass, and one of stainless steel and glass. The 316 stainless steel reactor, a Benco Model 150-E equipped with a gas sparge tube, dual heating mantles, condenser, and pressure gauge, is shown in Figures 1 and 2. The Benco stainless steel reactor was used to evaluate phosphorus Reactor #1, consisting of a 4000 ml beaker heated by a hot plate and covered by a 2000ml round flask to serve as an air condenser, was used to evaluate treatments involving fabric immersion. Glass Reaction #2, consisting of a 2000mf resin reaction kettle, Scientific Glass Company Catalogue Number J4401, equipped with an electric heating mantle, oxygen sparge tube, and a condenser, was used to conduct reactions in the vapor phase under reflux conditions. A reactor constructed from 310 stainless steel and Pyrex glass was used to process continuously the seven square yards of fabric submitted to the Air Force Materials Laboratory in fulfillment of one of the contract requirements of Phase I. This continuous fabric treatment reactor, which was the prototype for the tow treatment facility in Phase II, described in detail below.

3.1.1 Continuous Reactor

A pilot scale treatment facility was constructed to evaluate the critical elements in the continuous process. A seven inch 316 stainless steel pipe was used as the reactor. Heat lamps were set up before and after the reactor to dry the fabric and vaporize excess phosphorus oxychloride, respectively. A motorized tall—up roll with variable speed control was used to vary the residence time of the fabric in the reactor. A Schematic diagram of the pilot facility is given on page 13. Following treatment of a web of fabric in the pilot facility, the roll of fabric was unwound and draped in an air circulating oven at 300° C for varying periods in order to complete the treatment.

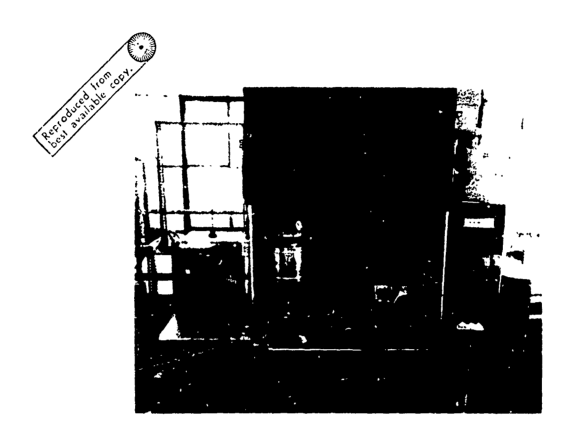


Figure 1.Photograph of POCl₃ Treatment Apparatus

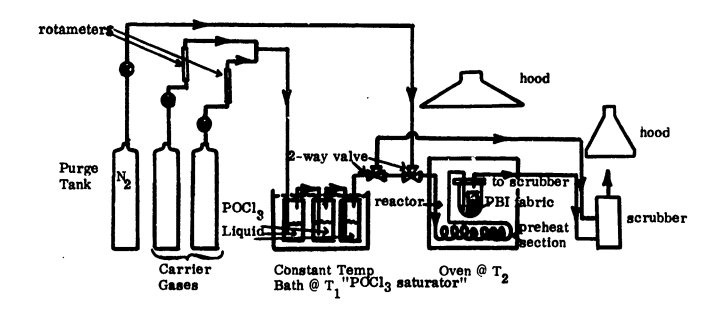
From left to right: Gas Cylinders Control Panel

Constant Temperature Oil Bath

Bubblers (removed from bath for photograph)

Reactor Oven

Scrubber (not shown)



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FIGURE 2 Schematic Arrangement of $POCl_3$ Treatment Apparatus

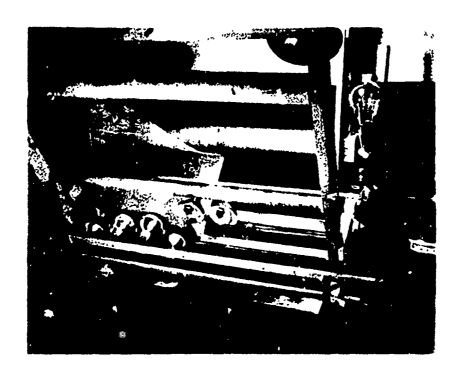


FIGURE 3

Continuous Process Line Set Up to Show

Path of Fabric

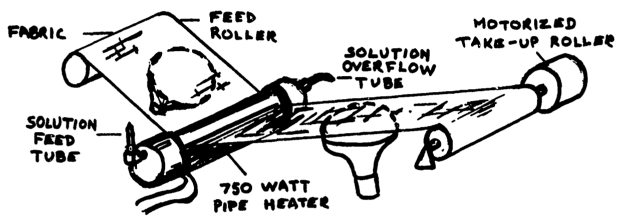


Figure 4. Schematic Diagram of Pilot Scale Treatment Facility

The pilot reactor was scaled up to treat a 46 inch wide fabric swatch. From the pilot work it was found that contact between layers of the fabric on the take-up roll caused discolorations and stiffening in the contact areas. To remedy this, polyethylene film was inserted between each layer. The full scale unit is shown schematically below and is pictured in Figure 3.

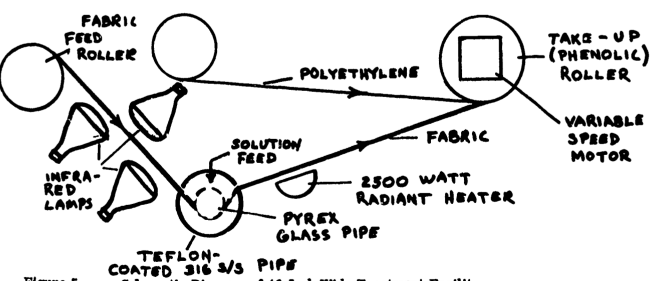


Figure 5. Schematic Diagram of 46-Inch Wide Treatment Facility

3.1.2 Chemicals

Reagent grade chemicals were used in the experiments except where otherwise noted.

3.1.3 Polybenzimidazole Fabric

The physical properties of the fabrics used in Phase I are given in Table II. The method for testing for fabric shrinkage during flame impingement is given in Appendix A.

3.1.4 Study of Effectiveness of Various Treatments

The study of treatments of the reduction of thermal shrinkage of polybenzimidazole fabric is summarized in Tables IV and XI.

3.1.5 Preparation of 7 Yards of Fabric Using Optimum Conditions

Table XII gives the details of the fabric treatment procedure and the physical properties of the fabric before and after treatment.

3.2 Phase II - Experimental

3.2.1 Apparatus

The pilot facility for the treatment of PBI tow is shown in Figure 4.

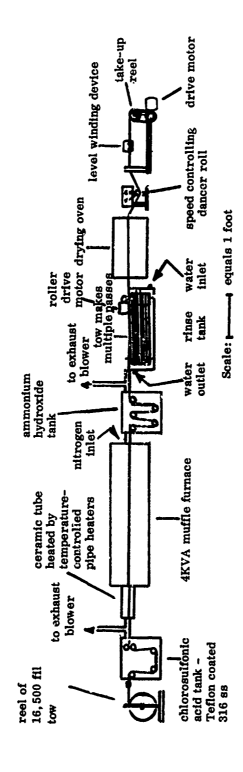
3.2.2 PBI Tow

In the process evaluation experiments 550 fil (1.5 denier/fil) fiber was plied to 16,500 fil tow. To prevent the tow from expanding as it passed over rollers in the process line, a twist of 1.5 turns/inch was incorporated into the tow.

In the production of 30 lbs of treated fiber, 5000 fil tow, plied to 15,000 fil and twisted 1.5 turns/inch was utilized.

The tow was treated in the as received condition - the lubricating oils were not removed prior to processing.

The physical properties or the fiber used in the process evaluation experiments are given in Table III.



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Chlorosulfonic Acid Stabilization Treatment to Processing Facility for the Application of the PBI Tow Figure 6.

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3.2.3 Chemicals

Technical grades of chlorosulfonic acid (duPont) and phosphorus oxychloride (Hooker) were used in Phase II processing.

Table II
Physical Properties of Polyberzimidazole Fabric

Physical Properties of	Polyberzim	idazole Fabi	<u> 10 </u>	
Sample Number	$6-1^{(1)}$	$6-2^{(1)}$	73-1 ⁽⁷⁾	118-1 ⁽⁸⁾
Thread Count - threads/in.	68 x 68	90 x 90	-	-
Weave	2/1 Twill	3/3 Twill	2/1 Twill	2/1 Twill
Weight - oz/yd ²				
As received	4.49	4.77	-	-
After Dry Cleaning (3)	4.51	4.88	-	-
After Dry Cleaning (3) and Washing(4	4.64	4.77	-	-
Tensile Strength $^{(5)}$ - lbs/in.				
' As received	98	97	94	-
After Dry Cleaning	95	95	-	-
After Dry Cleaning and Washing	91	93	98	100
Elongation at Break ⁽⁵⁾ - %				
As received	25	20	23	-
After Dry Cleaning	21	10	-	_
Af After Dry Cleaning and Washing	20	13	22	17
Shrinkage Druing Flame Impinge-				
ment(6) - %	57.5	61.9	62	65
FOI(3)	-	-	9. 36	0.34

Notes

- (1) Samples received from Air Force Materials Lab (AFML) on 6 July 1970, approx. 3 yds² of 6-1 and 4 yds² of 6-2.
- (2) Fed. Test Method Std. 191, Method 5040, full width short specimen method.
- (3) Dry cleaned for 45 minute cycle in a coin operated dry cleaning machine.
- (4) Washed in 8 lb. commercial coin operated washer with reciprocating agitator for full cycle through spin dry, using hot water and 38 grams of standard neutral step chips obtained from AATCC Research, Triangle Park, N.C. 27709, and 38 grams of technical sodium meta silicate, Will Scientific Co., Cambridge, Mass., and dried for 30 minutes in a commercial coin operated dryer.
- (5) Fed. Test Method Std. 191, Method 5104 three specimens warp direction only. Test performed on fabric after washing and dry cleaning.

Notes continued on next page.

(6) The shrinkage test used consisted of exposure of fabric to 1300° F temperature for 1 minute (see Appendix A). This thermal exposure greatly exceeds the thermal input anticipated in personnel-survivable fires. This test was used as a research tool to screen candidate stabilization treatments and the shrinkage results obtained greatly exceed the shrinkages obtained in end-item application tests such as the open-pit flame test.

(7) Approx. 20 yds² received from AFML on 21 August 1970.

(8) 3 yds² received from AFML on 11 January 1971.

(9) LOI (limiting oxygen index) is the minimum percent exygen by volume in a mixture of oxygen and nitrogen in which the sample will burn.

Table III Physical Properties of PBI Fiber (550 fil)¹

Denier 1.5
Tensile Strength² Shrinkage on Thermal
Exposure³ 60-70%

Notes

- (1) Received from the Air Force Materials Laboratory March 1971
- (2) See Appendix B
- (3) See Appendix C

TABLE IV

Study of Treatments for Reduction of Thermal Shrinkage of Polybenzimidazole Fabric*

steel 140 washed with H ₂ O steel 150 washed with H ₂ O steel 150 washed with H ₂ O steel 140 sods ash + H ₂ O steel 90 water wash steel 240 water wash steel 60 water wash steel 60 water wash	150 150 150 150 150 150 150 150 150 150	15	11 15 15 15 15 15 15 15 15 15 15 15 15 1	150 140 150 150 150 150 150 150 150 150 150 15	120 33 3 3 3 5 5 8 8 8 8 8 8 8 8 8 8 8 8 8	140 140 140 140 140 140 140 140 140 140					waahed with Hgo waahed with Hgo waahed with Hgo eods ash + Hgo water wash wat	waahed with HgO waahed with HgO waahed with HgO sods ash + HgO water wash NaOH + HgO NaCO3 + HgO NaHCO3 + HgO	washed with H ₂ O washed with H ₂ O washed with H ₂ O soda ash + H ₂ O water wash water wash water wash water wash water wash NaOH + H ₂ O NaH + M	anbed with H ₂ O anbed	washed with H ₂ O washed with H ₂ O sods ash + H ₂ O washed with H ₂ O sods ash + H ₂ O water wash water wash water wash water wash water wash water wash NaOH + H ₂ O NaH + M ₂
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anhydrous anhydrous anhydrous anhydrous bubbler anhydrous	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux wet with	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux under reflux toff H3PO ₄	ith 5 H ₃ PO ₄	ith 5 H ₃ PO ₄	418 A 3 PO.		8. j	84, al	8 ⁷ , a	S, ā	Š, ž F	Š, ž į	Š, ž ť
					anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux under reflux wet with 10% H ₃ PO ₄	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux inder reflux inder reflux	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux under reflux under reflux fabric wet with 10% H ₃ PO ₄	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux 20% H ₃ PO ₄ painted on wet with POCI ₃ , no reflux	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux under reflux under reflux under reflux under reflux under reflux reflux tog H ₃ PO ₄ fabric wet with 10% H ₃ PO ₄ 20% H ₃ PO ₄ painted on wet with POCi ₃ , no reflux reflux, no POCi ₃	anhydrous anhydrous anhydrous anhydrous anhydrous under reflux reflux 10% H ₃ PO ₄ iabric wet with 10% H ₃ PO ₄ 20% H ₃ PO ₄ painted on wet with POCl ₃ , no reflux reflux, no POCl ₃ no reflux, no POCl ₃	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux reflux 20% H ₃ PO ₄ painted on wet with POCI ₃ , no reflux reflux, no POCI ₃ no reflux reflux, no POCI ₃ reflux sample heated 700°F, 3-1/2 hrs.	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux reflux anhydrous 20% H ₃ PO ₄ painted on wet with POCl ₃ , no reflux reflux, no POCl ₃ no reflux, no POCl ₃ no reflux reflux annyle heated 700°F, 3-1/2 hrs.	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux reflux 20% H ₃ PO ₄ painted on wet with POCl ₃ , no reflux reflux, no POCl ₃ no reflux, no POCl ₃ no reflux reflux sample heated 700°F, 3-1/2 hrs. heated 700°F, 3-1/2 hrs.	anhydrous anhydrous anhydrous anhydrous anhydrous anhydrous under reflux reflux reflux, no POCi3 no reflux, no POCi3 reflux aample heated 700°F, 3-1/2 hrs. heated 700°F, 3-1/2 hrs. POCi3 wet out POCi3 wet out POCi3 wet out POCi3 wet out
															
Ť	2 2 8		2 2 8 2 2 3 5 2 3 5 3 5 5 5 5 5 5 5 5 5 5 5 5	228	238 238	2 2 2 2 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9		2 228	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	228 228 228 228 228 228	222 228 0 0 0 2 228 228 228 228 228 228	2 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	1 228 6 6 2 228 1	2	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
240	300	240 300 218-228	240 300 218 - 228 218 - 228	m 300)mm 218-228)mm 218-229 m 225-235	n 240 nn 300 0mm 218-228 0mm 225-228 n 225-235	m 300 mm 218-228 218-228 mm 225-235 m 225-235 240	240 228 240 240 240 240 240 240	240 218 - 228 218 - 228 225 - 235 240 240 240 - 240	240 7 218 - 228 7 218 - 228 7 225 - 235 7 240 240 240 240 240 240 240 240	240 218 - 228 218 - 228 218 - 229 240 240 240 240 240 240 240 240	240 218 - 228 218 - 228 218 - 229 219 - 229 240 240 240 240 240 240 240 240	240 300 300 300 300 318-228 325-235 340 240 240 240 240 340 340 340	240 300 300 300 300 318-228 318-228 340 240 240 240 240 240 340 340 340 340 340 340 340 3	240 300 300 300 300 300 318-228 340 240 240 240 240 240 240 240	240 218 - 228 218 - 228 240 240 240 240 240 240 240 240
8	on Turn Turn Turn Turn Turn Turn Turn Tur	10 mm 10 mm 10 760 r	0 mm 01 mm 0	40 mm 40 mm 1. 760 i 5. 760 i	. 440 mm (gas, 760) (gas, 760) 440 mm ? POC! ₃	s, 440mm s, 440mm l/gas, 7601 l/gas, 7801 ls, 440mm r, POC1 R	as, 440mm as, 440mm as, 440mm tq/grs, 7601 tq/grs, 7601 dq/grs, 7601 of: POCl ₃ of: POCl ₃	as, 440mm as, 440mm kg/gra, 7401 kg/gra, 7401 jus, 440mm joř, POCl ₃	10, 440 mm 10, 440 mm 10, 420, 760; 11, 410 mm 10, 70C; 10, 7	10, 440 mm 10, 440 mm 10, 624, 760; 10, 440 mm 17, POC! 17, POC! 18, 440 mm 19,624, 760; 10, 440 mm	a. 440 mm a. 440 mm b. 440 mm b. 62a, 760; c. 440 mm c. 70ct ₃ 7 POCt ₃ 7 POCt ₃ 7 POCt ₃ 8, 440 mm f/gas, 760; f/gas, 760; f/gas, 760; f/gas, 760;	1. 440 mm 1. 440 mm 1. 440 mm 1. 440 mm 1. 400 mm	440 mm 440 mm 440 mm POC13 POC13 POC13 POC13 POC15 POC15 POC15 POC15 POC15 POC15 POC16 POC16 POC17 POC	0 mm 0 m	
	POCIs pre. 44														
	300 Og, 0. 8SCFH	300 O ₂ , 0.85CFH 218-228 O ₂ , 0.48CFH	200 O ₂ , 0.8SCFH 218-228 O ₂ , 0.4SCFH 218-229 O ₂ , 0.4SCFH	m 300 O ₂ , 0.8SCFH Dmm 218-228 O ₂ , 0.4SCFH Dmm 225-228 O ₂ , 1.9SCFH	m 300 O ₂ , 0.8SCFH Dmm 218-228 O ₂ , 0.4SCFH T 225-235 O ₂ , 1.9SCFH 240	m 300 O ₂ , 0.6 SCFH Dmm 218-226 O ₂ , 0.4 SCFH T 225-235 O ₂ , 1.9 SCFH 240	m 300 O ₂ , 0.8 SCFH Dmm 218-228 O ₂ , 0.4 SCFH m 225-235 O ₂ , 1.9 SCFH 240 240	Dmm 218-228 O ₂ , 0.4 SCFH 218-229 O ₂ , 0.4 SCFH 225-235 O ₂ , 1.9 SCFH 240 240 240 240	Dmm 218-228 O ₂ , 0.4 SCFH Dmm 218-229 O ₂ , 0.4 SCFH T 225-235 O ₂ , 1.9 SCFH 240	m 300 O ₂ , 0.8SCFH Dmm 218-228 O ₂ , 0.4SCFH 218-225 O ₂ , 0.4SCFH 240	218-228 O ₂ , 0.4 SCFH 218-229 O ₂ , 0.4 SCFH 240	m 300 Og. 0.8SCFH Dmm 218-228 Og. 0.4SCFH Dmm 218-229 Og. 0.4SCFH m 225-235 Og. 1.9SCFH 240 240 240 nm 218-228 Og. 1.9SCFH nmm 218-228 Og. 1.9SCFH nmm 218-228 Og. 1.9SCFH nmm 218-228 Og. 1.9SCFH nmm 218-228 Og. 1.9SCFH	780 02, 0.85CFH 782 226 02, 0.48CFH 782 225 225 02, 1.95CFH 782 240 786 24	mm 218-228 O ₂ , 0.48CFH mm 218-228 O ₂ , 0.48CFH m 225-235 O ₂ , 1.98CFH 240 2	m 300 O ₂ , 0.8 SCFH 0mm 218-228 O ₂ , 0.4 SCFH 0mm 218-229 O ₂ , 0.4 SCFH 1 225-235 O ₂ , 1.9 SCFH 240 240 240 240 340 340 340 340 340 340 340 340 340 340 340 340 340 340 340 340 340

See Table II for Fabric Identity

ANTINE OF THE SERVICE OF A LONG TOWNS AND THE SERVICE OF THE SERVI

S è	Fabric Type	Reagent	Reagent Concentration	Reaction Temp. F	Carrier Gas	Spectal Conditions	Reactor Type	Reaction " ,mc min.	Neutralization Procedure	Wt. '' Addition	Shrinkuge	Tensile Strength Ibs/in
11	3	POCI3	liquid	230			Kluss	09	water rinse		45.1	:
:	7	POC13	Mguld	33	:	•	glass	30	water rinse	11.7	44.7	:
:	6-2	Poci	13quid	150	!	:	glass	9,	water rinse	10.4	43.5	i
20	ï	POCI3	Itquid	130	:	!	gl.158	۰,۰	water rinse	10.4	53.5	;
21	ï	POC13	liquid	2	:	1	glass	30	water rinke	9.0	53.2	
22	6-2	POC13	liquid	2	:	•	gjass	30	Na,CO,+H,O	10.3	8.4.8	1
ន	6-2 -4	Poci	Liquid	ş	;		glass	81	Na2CO3	!	58.1	1
7	7-9	POC13	Bquid	2	1	:	glase	s	water rinse	4.9	58.1	ļ
ដ	ï	Poci3	liquid	8	•	1	glass	30	water rinse	9.6	58.1	1
2	6-2	POC13	Ilquid	٤	1	1	glans	2	water rinse	0.9	61.9	i
27	6-2	POCI3	liquid	39	;	•	glass	s	water rinse	2.5	61.6	;
28	-7	Poci	Uquid	93		ł	glass	ဗ္ဂ	water ringe	3.1	59.4	
2	6-2	POCI3	liquid	8	!	*	Klzss	83	water rinse	1:9	58.0	
8	6-2	POCI3	Isquid	၉	!	1	glass	30	water rinse	4.2	2.3	1
E	6-2	20C13	11qu/d	5	1	:	glass	9	N2,CO3 + H2O	8:	59.0	1
Ħ	ĩ	POC13	liquid	130	1	1	glass	ဗ	3,a,co, + H,o	10.3	46.4	ł
8	6-2	POC13	liquid	150	1	!	gluss	99	Na,CO, + H2O	9.0	50.1	ł
ಕ	6-2	POCI3	Ilquid	130	1	-	Slass	ເວ	Na2CO, . H,O	6.6	53.1	i
<u>.</u> ع	6-2	POCI	liq gas, 750 mm	218 - 228	O2. 1.9SCFH	H, PO, sprend on frbric	steel	8	Na2CO3 + H2O	!	23.0	;
				*****		waste POCI3 used						
ş	6-2	H3PO4	Nguid	360	!		glass	ಜ	Na2CO3 + H2O	-	30.3	;
98	2-9	нзРо4	Manid	268	!	-	glass	ಜ	V.1,2CO, + H2O	-	23.0	ì
š	6-2	H3 PO	liquid	268	:	:	glass	ê	Na2CO3 + H2O	!	30.8	}
ž	2-9	#3.00 1	Mand	243	!	;	g1388	13	N.12CO + H2O	;	30.3	i
36	6-2	н3№	liquid	288			KLISS	30	N.12CO3 + 112O		31.2	;

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TABLE V-1

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

			į	i const	200	Ringing		2	3				ě	S String	. 1
₹ %	Fabric Type(1)	Respent	T. J.	Temp Time			Tenp fime		Jemp Time	Aiklitional Treatment	Remarks	Character	Treatmen	Treatment Over-all Only Shriskage	Thermal Exposured)
5	2.	200 %					·		•						
		lomi Br ₂		:							Fabric amelia of Br2		•	,	ĸ
8	-	10% L h, Cr207	212				,		,	•					
		to of 1 to HCC	212	17	0 ⁷ H	E	_	380	12	ı	Color unchanged	ė	•	•	2
23	7	10g/L K2Cr20,													
		to PR of 1 & MC	212	17	Н20	RT	_	2	2	•	Color unchanged	non- burning	,	'	8
2	7	10g/L K2Cr201											•		}
		to pill of 1 th HCE	212	n	H ₂ 0	ž,	_	380	so	1	Color unchanged	•	•	1	35
-	:	100g/LK2Cr207													
		to par of 1 . HCP	312	·s	H ₂ 0	Ħ	_	380	2	i	Color unchanged	•	,	1	3
2	7	Set Solu K ₂ Cr ₂ O ₇													
		to per of 1 & HCZ	212	٠,	H ₂ O	TH.	_	85	55	ı	Color unchanged		,	•	3
***	;	10g/L K2Cr2O7			_										
		to per of 1 & HC!	312	10	н20	RT	-	965	17	ı	•	•	•	•	*
*	:	10g/L K2Cr207													
_		to pill of 1 & HCt	33	12	H ₂ 0	RT	_	9	22	ļ	,			,	3
ส์	7	100g/LKgCr207	212	'n	0,7 E,0	RT	_	980	\$	•	1	•	,	ı	\$
_		STE SE TO HOS								•					
ř	†		Same a	same as above			<u> </u>	98	SS SS	•	•	•	,	•	43
e e	十		same a	same as above			†	980	120	•	Darker after beating	•	•	,	*
-			₹ 	omitted			t	ş	210	•	,	•	•	•	5
<u> </u>	:	SS/LK,CrgO,	<u> </u>	2					2	leach in (NaPO ₄)6 + HCt 5 min rinse	Color br black after	slight	•	•	#
		S PE OF S RC	2	8	H ₂ O	Þ		390	02	leach(NaPO,16+ NaOH, 4 rinsos NaOH at 212 dry	lat day very slight green at end	E Duca			

des last page of Table for notes. O - with

CARLE STATE CONTRACT CONTRACT

在在原始的时候,我们也是有一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人, 第一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就是一个人,我们就

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC TABLE V-2

a a ser i de de la companya del la companya de la companya del la companya de la companya de la companya del la companya de la companya de la companya del la c

									a de la companya de l	à		Ł
	Fabric	SIE	Conditions Temp Time	Kineing Comp Time			Additional Treatment	Remarks		Treatment	Over-all Shrinkage	Thermal Exposure (3)
2 3	10g/LK_Cr.O.	1 2	2	H,O RT 1	_			Durkened slightly	Slight			
				•		1	•		Dorning			;
	pice acid											
_		8.1me 1	sume as above		8	15		Soft darkened	prime	· —		6£
		#2TH	same as above		8	105	•	Less soft than				
<u> </u>								S8, and darker	1			23
7-5 VR	10g L KildaO	213		H2O NT 1	·		•					æ
36B 6-2		212		H,O RT 1	•		•	Very dark	,	<u>.</u>	,	#
	10mf glantal											
	acetic actd			E 3			1	-				•
**	<u>. </u>	212	- P					Very dark			•	÷
-	100 m Checks											
	10e/L 13440, •	32	1 0	H,O RT 1	2	<u>.</u>		•	•		'	Ş
				•						~~~		
- - -		8	-	•			•	•	•	<u>.</u>		8
	various doses							_				
	up to 12 Mrade								,			
Control 73-1		omitted	_		1		1	Soluble in boiling	LOI =0.36(2)			•
		_						NMP				
M 6-2	2 4 mi /L 70%	212	2	runeling H ₂ O	410	s	•	Insoluble in boiling	LO?=0, 35(2)	-		22
	NO.			1				NMP (purple color)				
		- 4	same as above		3	-	•	Color unchanged	LOI=0.35(2)	- 12	,	38
13	4m1 /L 70%	212	2 - 2	ruseling H ₂ O	*	-	•	Some brown spots	,		'	37
				•	\$	s	,	•	•	•	•	
¥	İ	- 3	eagne as above		166	-	,	•		· 	•	;
1		_			2	_		•				32
ž		Same E	same as above		₽	<u>د</u>	Sample dyed with Maxillon	Old not turn purple			'	'
<u>.</u>							Blue before drying					

See last page of Table for Notes.

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TABLE V-3

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

			3			å	Ĭ			Remine	en Æ	S Shrinkage	ă
19	7	Fabric Region 7		14	Rissing	Temp Time	1	Additional Treatment	Remarks	_	Treatment Only	Over-all Shrinkage	Thermal Exposure (3)
\$	8-9 Yet	40 mt /Libraics	BT	•		8	2		,		•		,
\$	<u>:</u>	2 200 mf /Libranic acid RT	E	•	Ĭ	200	2	,	•		22		,
\$	1 2 2	2 400 mt /Lfermicacid RT	E	8	reming H ₂ O	*	=	•			42		•
\$	<u>;</u>	3 600 mt /L formic acid RT	Ł	•	Ĭ	*	2	•	•		3		•
\$	7 20		H	•	Ĭ	*	2	•	•		ı		
1	<u>:</u>		2	'n	remaine M.O	•:	•	,	deep purple color		,		11
£	<u>:</u>		212	•	reside H O plus	•	٠	•			,		
					1g/L NeOR room temp 10 min & ruening water								
7	7	2 M 2 M 3 PO	Ħ	•	NaOH pH12,212"F, 15	\$	22	•	soluble in boiling NMP	<u> </u>	32		,
		•			mis reming water								
ฮ	CEL	2 400 m/ /L 5.20%	=	=	•	2	• —	•	cloth stiff and color				*
		Chlera							slightly darker				
ช 2	CEL PER		Ë	2	remaing bot water	9	"						2
2	_	Chlere											
៩	770	2 400 mt /L 5.28%	22		•	410	••	,	cloth stiff and color		,		25
		Chlores plus 4 gm/L							slightly darker				
			_	_									
ប	CLF F.		=======================================	s	resaling hot water	e Ç	*	•	•		,		33
\$		2 TOS HNO.	Ħ	~ ያ	ringe and immerse in	9	51	Rinsed and immersed in KNO, & HCr	deep purple color in-		•		2
	_		_		Sact, sol's & HC!			after SnCf , treatment and before	soluble in boiling NMP	<u>.</u>			
	_				•		_	drying	non-dy sable at boil at				
				_					5% formic acid con-				
									taining 1.5% of various	9			
									dyes				
\$	-	TAS HNO.	Pa :	=	rince and immerse in Sect 2, sol'n & HCt	8		Rinsed and immersed is POC' 3 at 76°F, after SuCf., treatment then	purple color		,		22
					1			rinse in Alcosox sol'a, riaved in H ₂ O before drying					
	\dashv		_	1									

See hat page of table for setse.

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TABLE V-4

COMMERCE DESCRIPTION OF THE OWNER, THE PROPERTY OF THE OWNER, THE

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STUDY OF TREATMENTS FOR MEDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

Treetment Cyer-all The Shrinkage Exp			Conditions		Daying			Burning	æ	y Saringage	Ã.
Same as above Same as abov		Reserved	Temp Time		Temp Time	Additional Treatment	Remarks	Charac-	Treatment Only		Thermal Exposure(3)
1-2 4 m L TOT HNO ₂ 69 5 running water 100 15 rinse - color unchanged color color color unchanged color color color unchanged color color color unchanged color col	1-		rim	l		same as above except dyed with	color green	ŀ	 -		22
1.2 4 m 1.707 HNO, 100 15 11 11 11 11 11 1			38.0	't goot'n & HCf		Maxillon blue GRC before final					
1-2 4 m L 707 H100 30 5 Transing water 300 15 Transing water 300 1						rinse - color green					
1.2 34 ms / 1.76 HNO; 10 15 10 10		4 m L 707 HNO,	2	running water		,	color unchanged	•	,	,	ន
Same as above Same as abov	25					•	purple stains	•	•	'	3
1-3 34 mt/L 195 MMO 30 5 Turning water 240 15 15 15 15 15 15 15 1	. E		same as above			•	color unchanged	•	•	,	28
1-3 24 m/L 175 Mod be 5 stand as above same same as above same same as above same same same same same same same sam							very brittle after				
10 15 15 15 15 15 15 15							shrinkage test				
1-3 34 mt / L 10\$ HNO ₂ 36 5 5 5 5 5 5 5 5 5	¥		same as above			•	purple stain		,	•	38
Same as above Same as abov	_	24 mt /L 10% HNO,	2	running water		•	color unchanged	•	•	* 	ಕ
Same as above Same as above 240 6 - - - - - - -	<u> </u>					,	color unchanged	•	•	•	ಸ
Same as above Same as above 410 S* Carrier purple Carrier purp	=======================================		same as above			•	elightly purple	•	•	•	*
Same as above Same as above Cart purple Cart p	ž		same as above			•	elightly purple	_	1	•	*
143 mi /L 70% HNO3 90 5 running water 240 16 -	¥		same as above			•	dark purple		,	•	23
143 mi /L 10% HNO ₃ 96 5 running water 240 16 - brittle after shrinkage test - brittle after same as above same as above same as above shifted test - continued test - contin			same as above			•	dark purple	<u>.</u>	•		99
### same as above		143 mi /L 70% HNO,				•	brittle after	_			
### ### ### ### ### ### ### ### ### ##							shrinkage teet	•	ı		8
Same as above Same as abov	lo,		same as above			1	brittle after .				
Same as above Same as abov	-						shrinkage test		•		8
######################################			same as above			•	very brittle after				
### ### ### ### ### ### ### ### ### ##	•						shrinkage test	_	•	•	8
L-2 40° Be conc HNO ₃ 93 5 Tunning water 300 5 - alightly purple (edges)	lo l		same as above			•	purple color	_	ı	•	¥
1-2 40'Be conc HNO ₃ 93 5 running water 300 5 - alightly purple (adges)	10,					1	•		ı	•	23
•	411, 6-2	40. Be come HNO.		running water		•	slightly purple	_	•		19
	•	•					(edges)				
		-						_			

See last page of table for notes.

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TABLE V-5

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

	Remarks purple streaks
	Durk purple streaks
Purple at	And the Market of the Park
Provan	Very Might Garagin.
purple	Color brown
purple	Calor brown
purple at	Tolor with seek
Turned purple	
Turned purple	
2 Deep blue color and out turn purple at 400°F	Turned purple
7 2 Deep blue color and	
7 2 - 15 42 42 42 42 42 42 42 42 42 42 42 42 42	•
15 56 Deep blue color and	•
72	
25 63 54 54 54 54 54 54 54 54 54 54 54 54 54	
Deep blue color and	'
Deep blue color and	,
Deep blue color and root turn purple at 400° F	
Deep blue color and not turn purple at 400° F	
-	
	not turn purple at

See last page of table for notes.

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TABLE V-6

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

			Conditions			2	;			Rumine	:~I	Shrinkage	Š
Ş	Fabric		a die	Time		:1=	<u>.</u>				Treatment	Over-all	Thermal
Š	Jane C	Resgent	4	Mins.	Rinaing	•	Mis.	Additional Treatment	Remarks	- 1	Chily	Shrinkage	Exposure(3)
458		7	ine is to	A except.	same is 15A except after treatment fibric dipped into PCC 3 at RI for	out boddip	LKC 3.	t R1 for	Olive Inage) green	٠			¥
			neutralis	Led with N.	2: neutralized with NagCO 3 solu ringe and dr.	ŧ	1		Color did not turn				
Ş	1-7:	*0. Be cone HNO,	77	n	SH,OH 12 plus	00±	2	Ringe in 30 -Htf sat with Snd ,	Purple in oven at 400°F		ı	•	66
		Mass?H230,11			ruming water			for 5 at RT wash II,O, ringe	•	_			
		lo vol						NH OH wash II to immerse					
								in liquid FOG gneutralize in	Forest blue color - did				
								Na CO solu, wash ringe NH OH	400°F				
47		<u> </u>	Asime as above	940		٠	•	same as above except repeated	Same as above			•	67
!				- 				POC 3 dip and wash					
‡	75-1	40° Be conc HNC,	-30	v	NH OHIPH 12	8	13	1	Coler unchanged before		25	88	\$
	dried 15'	dried 15' plus 96't H ₂ 50 ₄			plue running				shrinkage test				
	at 200°C	at 200°C 1 1 by vol.			water								
\$	12-1	40° Be conc	۰	•0	NH,OHFH 12	904	15	96%H,SO_added to HNO, after	Non-uniform purple		- 23	25	19.5
	dried 15" HNO3	HNO			plus ruming			S' temp rose to 12°C	color developed				
	at 200°C		•	50	water								
494	13-1	Pod 3	T.	so.	NH OHPH 12	00 †	13	Dip sample in cold ethylene	•	•	•	\$	•
					plus rumung			glycol after immersion in					
					water			POC/3					
4			same as above	ove			-	Dip sample in cold methanol	•	<u>.</u>	,	,	•
								after immersion in POCt 3				\$	
¥	13-1	POG 3	212	8	NH4OHPH 12	400	15	Dip sample in boiling ethylene	•	•		ដ	•
		,			plus running			glycol for 20 mins and rinse					
					nater			Dip sample in boiling methanol				39	15.7
								for 15 mins and rinse					
8	- 5. - 1. - 1.	40. Be conc	ę	15	NH OHDH 12	00 	15	Purple only in crease	Purpled only in crease	•	· —-	•	•
	13. 21	HNO3			plus running				but tensile strength 51				
	200°C				water				lbe/in vs. 87 for control				
		County for many											

See last page of table for notes.

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TABLE V-7

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

Reader Conditional Richard Free Free Free Free Free Free Free Fr													
Racgon Or Mine Reconc Or Mine Reconc Or Mine	í			Hillone		Tall Day	Time			Burning Charac-	By Treatment	Over-all	The Tail
10 Be conc 5 13 NH, OH pill 2 plus 100 13 NH, OH pill 2 plus 100 13 NH, OH pill 2 plus 100 15 NH, OH pill 2 plus 100			ò	Mins.	T T	٥	Mins	Additional Treatment	Remarks	teristics	(iuc)	Shrinkage	Exposure (3)
HNO3	dried	ŀ	37	13		3	2	•	Nurpled only in creases.				15.7
157 HNO3, 110 13 1311, OH pill 2 plus OH pill	*	HNO			running w.iter			-	but tensile strength 51				
### 1767 H.W.Q. ### 12 13 NII, OH PHI12 plus	ن د	n			1				lb/in vs. 87 for control.	•	•		
967 H.SO. ³ 967 H.SO. ³ 967 H.SO. ³ 17 Truming water 18 12 100 18 20 4 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	, _	70° HNO.	21-	13	NH, CH pH12 plus	2	::	•	Fabric yellowish brown				•
90°C H.N.O. 3 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 73 100 19 74 10 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 10 19 75 10 10 10 19 75 10 10 10 19 75 10 10 10 19 75 10 10 10 19 75 10 19 75 10		98- H SO			running water				large discolored area				
73 73 100 9 yol 10 sec 834 10 0 13 20 0 13 20 0 13 20 0 13 20 0 13 20 0 13 30 0 13 30 0 13 30 0 13 30 0 13 30 0 13 30 0 13 30 0 13 30 0 14 400 15 30 0 15 30 0 15 30 0 15 30 0 10 3		10. H. XO							appears to have yellow				
by voil 9 youl 9 of une 9 ht 0.3 1 same as above		75 75 100							precipitate on fabric.			_	
## 200° for 10 sec Same as above 10 sec Same as above 10 sec 854		· ·	_										
Soc 86 St Soak Hamber H		o, vo.	-10	10 sec		200	S	•	•	•	•		
HyPO4. Sog ures		Sec 854	; ** •	1		400	2	•			28	÷	36
1970, 1960		300	, 					_				_	_
1980cc H ₂ O 1980		3,00											
Sog urea pH 0.3 same as above plus same as above plus same as above plus heating water same as above plus heating ater and dry same as above plus heating water same as above plus water same as above plus heating water same as above plus as above plus water same as above plus as above plus water same as above plus water		160cc H ₂ O											
### 8.00 15 Cloth darkened and political washing cycle(4) 400 15 Cloth darkened and political washing cycle(4) 400 15 Cloth darkened and place on bot plate and political washing cycle(4) 400 15 Cloth darkened and political washing cycle(4) 400 15 C		Sog ures				_							
Same as above plus heating water 100 15 15 15 15 15 15 15		PH 0.3	_								;	;	;
Same as above plus beating water running water running water running water running water		t pre sure	Par	1	NH OH PH12 plus	\$	15	•	,		2	8	si —
State as above plus beating same as above plus 100 15 15 15 15 15 15 1				•	runding water								
2 min place on bot plate 2 min place on bot plate 4 and POC 3 added, 4 and POC 3 added, 4 dropper 6 cone H ₃ PO ₄ 862 5 10t H ₃ SO ₄ at 6 for 10° ringe in 15t 6 2 min place on bot plate 4 and POC 3 added, 4 dropper 6 and POC 3 added, 4 dropper 6 and POC 3 added, 4 dropper 7 fabric attrank and 8 turned yellow 4 at 400°F. Fabric turned yellow 4 at 400°F. Fabric turned yellow 6 at 400°F. Fabric attrank and 7 fabric attrank and 7 fabric attank and 7 fabric attrank and 8 fabric attran	82 H	s as above plus h	Mille		same as above plus	\$	15	•	Cloth darkened	•	•		\$
Diluge HNO 3 PHO. 3 for - Fabric turned yellow 2 min place on bot plate when POG 3 added, 4 add POG 3 with eye dropper add not turn purple come H3PO, - - Fabric shrank and e62 5 - - Fabric shrank and 10t H3SO, at 322° for 15° wash heat at 395° C - - After H2SO, color plue 85t H3PO, at 570 for 10° rinee in 15t NH4OH, rinse in water atter H3PO, greenish brown atter H3PO, greenish brown	21 70				washing cy cle(4)								;
2 min place on hot plate 2 min place on hot plate 2 min place on hot plate 4 and POG 3 added, did not turn purple 3 at 400°F. come H ₃ PO ₄ 85°F 10°F H ₃ SO ₄ at 10°F H ₃ SO ₄ at 10°F H ₃ SO ₄ at for 10° rine in 15°F plus 85°F H ₃ PO ₄ at 139°C Fabric shrank and turned black & brittle. After H ₃ SO ₄ color yellowish brown. at 230°F for 10° and dry and dry brown	_	Dilute HNO, pl	10, 3 %		•	4 0	15	•	Fabric turned yellow		•		g
# add POCT 3 with eye dod not turn purple		2 min place on	hot pla						when POG 3 added.				
dropper as 400°F. come H ₃ PQ ₄ Fabric shrank and 85% 662 5 10°E H ₂ SQ ₄ at turned black £ 10°E H ₂ SQ ₄ at hard H ₂ SQ ₄ color 10°E H ₂ SQ ₄ at for 10° rinse in 15% plus 85% H ₂ PQ ₄ NH ₄ OH, rinse in water at 230°F for 10° and dry		and Pod .	rith eye						did not turn purple	•		•	
cone C H ₂ PO ₄ 85°F 10°E H ₂ SO ₄ at cornor rines in 15°F plus 85°F H ₃ PO ₄ at 230°F for 10° and dry Fabric shrank and		dromer	_						at 400 F.				
10°E H ₂ SO ₄ at 10°E H ₂ SO	_	conc H. Po.			•		1	•	Fabric shrank and	•	*	,	
wash heat at 395°C After H ₂ SO ₄ color for 10° rinse in 15°C at lower H ₃ PO ₄ greenish and dry		\$ C 258	662	47					turned black &			_	
wash heat at 395°C After H ₂ SO ₄ color for 10° rine in 15°C attentian brown, attent H ₃ PO ₄ greenish brown and dry		Š	_	,					brittle.				
for 10' rinse in 15% NH ₄ OH, rinse in water and dry		10% H 30 at	_		wash heat at 395°C	•	•	•	After H, SO, color	•	\$		22
MH ₄ OH, rinse in water and dry		100 % for 15			for 10' ringe in 158				yellowish brown.				
Alp pur		Of the page of the	_		NH OH whee in water				atter HaPO-greenish				
שים פול		pine 63't m3 re	-						Process 5			_	
		at 230 F for 10	è		and dry				ma Old				

See last page of table for notes.

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TABLE V-8

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

													2 Shrinkage	i
N. S.	Fabric Type (1)	t Reagent	Conditions Temp Time F Mins.		Rineing Fine Time		Dring Temp Time		Additional Treatment	Remarks	Burning Charac- teristics	By Treatment Only	By Treatment Over-all Only Shrinkage	by Thermal Exposure(3)
37.4	_	10. H. SU. at			•			•		Fabric shrank			\$*	•
		plus Lupersol												
		133 (4) at 437°F											•	
		for 3:								:			!	
37B	73-1	Epichlorobydria	•		•				_	No stiffening or fabric			÷	•
		brushed on fabric	¥.						_					
		dried 500°F 5'				_								
\$7C	<u></u>	10% H ₂ CO_plus	•		•		•	•	_	Treated areas did	•	•		
		drying at 700°F				_			-	did not dye with				
		for 10°				_				combination of				
										Geigy Maxillon				
									-	Blue and Polyamide				
돛 27	73-1	Same Y-431067 392		_	•		•	•		Fast Blue		•	,	19
25	2-52	E 'me A 1100 (6) 352	h 392 15	_	•	_	•			•	•	•	,	3
8	Ę	250 cc POCt 3 19g	ž.							•				
		terephthalic						-		•				
		scid, 25 cc PCr \ 212	§212 15	_	•					•	,	,	•	3
		10g At Cit 3			i									
3	73-1	Same as above	212		Water & NH OH	_	428 10		_	•		•		\$
		Substituting		* water	ta es	_								
		terestere												
		seld												
61		Similar to above using	_			_								
	Chlere	chlerendic anhydride	same as above	DOV 6					_	•	•		1	20
613	•	same as above	-	me as above	same as above except rinsed after drying	after	Irying			•	•		•	31
\$	£	10% HySO tested at 428°F for Water and	d at 428°F fo.	r Water	pu#	-	428 10			•	,	•	,	ន
		10' plus 85'E H3PO, at 230°F	PO at 230°5	HO HN	52									
		for 15'				_								
62B		same as above	except dried I	before neu	same as above except dried before neutralizing in NH, OH and rinsing	#5 Full	rinsing				,	•		27
				_		-								

a anterior <mark>de establicad</mark>a vasta Caretz de Bantana a Territa man de Carista mantes Mande de La Sala Lague e Aste i

TABLE V-9

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

No. Type 11 Reagent Type Time	l				Condi	tions	Rinsing	Drying			Burning	ğ.	3 Shrinkage	BA BA
63A 73-1 30 cc chlorosulfonic 212 3 157 63B 4	ĽŽ	9 6	Fabric Type (1)	Reagent	Temp	Time Mine.	Jemp Time F Mins.	Temp Time F Mins.	Additional Treatment	Remarks	Charac- teristics	Treatment Only	Charac- Treatment Over-all teristics Only Shrinkage	Thermal Exposure(3)
63B 4 PCC: + 2g Al Cl ₃ 63B 4 actid plus 270 cc 63C 73-1 30 cc chlorosulfonic 212 3 no actid plus 270 cc POCl ₃ 63D 73-1 30 cc chlorosulfonic 212 3 157 actid plus 270 cc PCCl ₃ 63E 4 same 23 63C plus treatment in 107 H ₃ PO 63F 13-1 10g 1 ₂ in 400 cc 85F H ₃ PO ₄ and	65	*	73-1	30 ce chlorosulfonic	212	ş	1ST NH OH	5.36 .3	Repeated treatment	•	•	•	53	
63B 4 8 9 8 9 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9				acid plus 270 cc POC: + 2s M Cr			•							
63C 73-1 30 cc chloroeulfonic 212 5 no actic plus 270 cc POQ 3 actic plus 270 cc POQ 3 actid plus 270 cc POG 3 actid plus 270 cc POG 3 actid plus 270 cc POG 3 actid plus 270 cc Aloroeulfonic 212 3 15% POG 3 actid plus 270 cc Aloroeulfonic 212 3 15% POG 3 actid plus 270 cc Aloroeulfonic 212 3 15% POG 3 actid plus 270 plus treatment in 10% H ₃ PO 63F 3 and 22 in 400 cc 300 30 15% NH ₂ OH 300 30 15% And 300 30 30 30 30 30 30 30 30 30 30 30 30	3	<u>.</u>	1	6			 same as above except	heated 30 mms at	212	•	•	•	SS	,
63D 73-1 30 cc chlorosulfonic 212 3 15°7 NH ₂ OH 536 acid plus 270 cc POCC ₃ 63E 4 same as 63E except risead in 10°7 H ₃ PO ₄ and drying 63F 73-1 10g I ₂ in 400 cc 83F H ₃ PO ₄ and rinse	3	ŭ	73-1	30 cc chlorosulfonic	212		no rinse	336 5	Repeated treatement	•	•	81	ដ	91
63D 73-1 30 cc chlorosulfonic 212 3 157 NH ₂ OH 536 acid plus 270 cc POCf ₃ 63E 4 same as 63E except risated in 107 H ₃ PO ₄ and drying 63F 73-1 10gT ₂ in 400 cc 83F H ₃ PO ₄ and rinse and rinse				actú plus 270 cc									·	
63D 73-1 30 cc chlorosulfonic 212 3 157 NH_OH 536 acid plus 270 cc POCf ₃ 63E 4 earne 22 53C plus treatment in 107 H ₃ PO ₄ and drying 63F 23-1 10gT ₂ in 400 cc 83F 73-1 10gT ₂ in 400 cc 83F H ₃ PO ₄ and rinse				P00 3										
63E 4 earne az 53C plus treatment in 10°T H ₃ PO ₂ and drying 63F same as 63E except rinsed in 15°T NH ₂ OH and dried 64 73-1 10g 1 ₂ in 400 cc 300 30 15°T NH ₂ OH and 428 65°T H ₃ PO ₄ and dried 65°T H ₃ PO ₄ 65	3	<u>e</u>	73-1	30 cc chlorosulfonic	212	"		536 5	Repeated treatment	•	•	8 2	8	*
63E 4 earne az 53C plus treatment in 10°T M ₃ PO ₄ and drying 63F same as 63E except rinsed in 15°T M ₄ OH and dried 64 73-1 10g 1 ₂ in 400 cc 300 30 15°T M ₄ OH and 428 65°T M ₅ PO ₄ and dried				acid plus 270 cc										
63E 4 same as 63E except risated in 19°F H ₃ PO ₄ and drying 63F 64 73-1 10g 1 ₂ in 400 cc 300 30 15°F NH ₂ OH and dried 64 85°F H ₃ PO ₄ and or 10°F NH ₂ OH 428				POC.										
63F same as 63E except rissed in 157 NH ₂ OH and dried 64 73-1 10g 1 ₂ in 400 cc 300 30 157 NH ₂ OH 428 655 H ₃ PO ₄ and rinse	2	Ä	1	same as 590 plus t	reatmen	t in 10%	H, PO, and drying		1	Fabric somewhat stiffer	•	13	ន	13
64 73-1 10g 1, in 400 cc 300 30 15f NH OH 428 8SF H3PO4 and rinse	3	<u> </u>		same as 63E excep-	rineed	in 15% N	TH, OH and dried			than before treatment	1	18	*	2
ber 1,046,	J		1-57	10g 1, in 400 cc	8	8	15°F NH, OH			Treated fabric lighter				
	28			854 H3 PO.		-	and rinse			than original	•	•	39	•

NOTES

(1) See Table I for fabric description. All fabric was laundered as specified in Table I before treatement.

(2) LOI (limiting oxygen index) is the minimum percent oxygen by volume in a mixture of oxygen and nitrogen in which the sample will burn.

(3) See Appendix A for procedure.

(4) 90% 2.5 dimethyl 2.5 di (t butylperoxy) hexane Lucidol Div.- Wallace and Tiernan

(5) Alcohol solution of proprietary silane-Union Carbide Co.
(6) gamma aminopropyl triethoxy silane - Union Carbide Co.

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC⁽¹⁾ TABLE VI-1

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		Conditions	900	Ringing		Dry	2			ă		
# 2. # 2.	Rengent	Time Time Semp F (Mins.)	Time (Mins.)		Time (Mins.)	Time Temp 'F (Mins.)	Time (Mins.)	Additional Treatment ⁽²⁾	Remarks	Treatment Only	Over-all Shrinkage	By Thermal Exposure(3)
1	100 ml L 96% H ₂ SO ₄	100	15	running water	01	100	15		Soft, flex de char after shrinkage test	,	,	91
3	50-2 100 ml/L 964 H ₂ 50 ₄	100	15	ruming water	2	300	22	Washed once in standard soap solution	Soft, flexible than after shrinkage test	1	•	2
3	59-3 100 ml/L 967 H2804	100	51	ruming water	2	8	2		Soft, florible char after shrishage test		•	2
ĭ	100 ml/L 18% H250	100	82	rundag water	10	200	51	Washed once in standard soap	Soft, Rexible char after shrinkage test	,	•	et .
2	100 ml/L 96% H_2O4	81	15	runding water	2	88	52	Washed twice in standard scap solution; pH of solution after wash ~6	Soft, flexible char after shrinkage test	1	•	£1
Ĭ	100 ml/L 96% H_50	100	51	runalog water	<u>s</u>	200	52	Washed three times as above: pH of solution after wash ~11	Soft, flexible char after shrinkage test	ı	•	a
7	100 ml/L 96% H ₂ 80 ₄	8	ŭ	runding water	2	200	55	After 4th wash as above, pH~11.5	Soft, flexible char after shrinkage test	,	,	‡
į	50 ml/L 96% H ₂ 80 ₄	98	5	runding water	2	300	55		Soft, flexible char after shrinkage test	1		••
?	56 m/L 98% H ₂ 804	100	51	runding water	2	900	15	Washed once in standard sosp solution	Soft, flexible char after shrinkage test		,	•••
?		861	ដ	running water	<u> </u>	200	22		Soff, flexible char after shrinkage test	ı	•	22
Į	50 ml/L 96% H ₂ 50 ₄	100	21	runaing water	2	800	22	Washed once in standard soap solution	Soft, flexible char after shrinkage test	1	1	2
\$	66-5 50 ml/L 96% H ₂ 80 ₄	100	22	conc. NH4OH plus running water	2	200	5		Hard, brittle char after shrinkage test	1	•	\$
==	61-1 20 ml/L 96% H ₂ BO ₄	100	22	running water	2	300	22			,	,	12
2-19	20 ml/L 96% H ₂ 80 ₄	100	25	running water	9	200	25			·	1	٠

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TABLE VI-2

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC⁽¹⁾

			,	i		•					9 Shrinksge	
Z. No.	Reagent	Temp F (Mins.)	Time (Mins.)	Medium	Time (Mins.)	Temp F (Mins.)	Time (Mins.)	Additional Treatment ⁽²⁾	Remarks	By Treatment Only	Over-all Shrinkage	By Thermal Exposure(3)
61-3	20 ml/L 98% H ₂ SO ₄	8	15	running water	95	200	22	Washed once in standard sosp	Soft, flexible char after shrankage test	•	-	a
7-19	20 ml/L 98% H ₂ 80 ₄	8	22	conc. NH4OH plus ruming water	0.	200	52		Hard, brittle char after shrinkage test		ı	
62- 1	++ ml/L 98% H ₂ SO ₄	E	v	ruming water	9	§ 	2	Immerse 10 gm/L urea in water after first rinse and rinse again; pH of urea solution after treat- ment = 2	Soft, flexible char after shrink go test	•	,	2
62-2	44 ml/L 969 H ₂ 80 ₄	¥	2	ruming water	9	220	9	Immerse 10 gm/L urea in water after first rinse and rinse again; pH of urea solution after treatment = 2	Soft, flexible char- after ahrinkage tost	•	ı	8
₹ 30	44 ml/L 96% H ₂ 80 ₄	R T	ĸ	ruming water	9	8	2	immerse 10 gm/L ures in water after first rinse and rinse again; pif of ures solution after trest- ment = 2	Soft, flexible char after ahrinkage test	•	•	92
2	44 ml/L 98% H ₂ 804	#	ဟ	ruming water	2	400	01	immerse 10 gm/L urea in water after first ringe and ringe again; pH of urea solution after treatment = 2		1	١	•
64-2		RT	ď	running water	2	00+	2	Washed once; pH cf wash = 6		•	,	22
2.3	44 ml L 98% 112504	H	20	runbing water	2	9	2	Washed twice		,	,	.
2.2	44 ml L98% H2804	T#	'n	running water	2	6	01	Washed twice; pH of wash = 9.9		,	1	93
65-1	20 ml'L glacial acetic	TH	22	cuou	•	91	2			•	•	62
65-2	20 ml/L glacial acetic	Ħ	2	none	•	007	8					62
65-3	20 ml/L come. HCl	RT	7	none	<u>'</u>	160	21				•	20
65-4	20 ml/L conc. HCl	RT	2	none	•	9	81	,		,	•	99
8 -1		BT	2								,	
	then into 120 ml/L 98% H ₂ SO ₄	RT	ç			00+	01				'	11
66-2	20 gm ures + 50 ml 407 HCHO + 100 ml water then into 120 ml L 989 H, 50,	RT	10			400	0.	Washed once; pH of soap was 3.8 after washing		•	ı	23
For	For notes see end of tables.											

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC⁽¹⁾ TABLE VI-3

TO THE PARTY OF THE PROPERTY O

		Conditions	980	Breeze			j				9. Brinkage	
2 & B	Reagent	Time Temp ° F (Mine.)	Time (Mine.)	Medium	Time dium (Mins.)	Time Temp F (Mins.)	Time (Mins.)	Additional Treatment (2)	Remarks	By Treatment Only	Over-all Shrinkage	By Thermal
1-1	Into 50% H3PO4 5' then 26% NH4OH for 1' then in 50 gm/L Cn 1 ₂ for 5'			running water	<u> </u>	400	r:					30 to 40
67-2				running water	01	400	'n			,		8
1-43						92	\$		Fabric turned brown	,	ı	\$
68-2	None	····				700	096		Fabric very dark brown, flexible, atrong char after shrinkage	ı	•	8
6-3						700	960	Rinse conc. NH4OH 5 mins., dried at 400°F	Flexible, strong char after af		1	88
3 3		TH	ب	running water	2	240	15	Washed once; pH of soap = 9 after washing	Brittle char after shrinkage	,	•	*
; 1	20 ml/L cone. HNO ₃	H.	v	running water	70	240	. 15	Washed twicu; pH of soap = 10 after washing	Brittle char after shrinkage	·	•	15
*	20 ml/L conc. HNO ₃	æ	6	running water	<u> </u>	48	w	Washed once; pli of soap = 9 after wabsing	Fabric turned purple after drying	ı	,	ង
1 2								Washed twice; pH of soap = 10		,	•	8
\$ - 8								Washed thrice:pff of soap = 10		ı	,	ដ
9-69								Washed four times; pH of soap = 10		,	•	ផ
\$	·				·			Washed 69-5 in conc. NH4OH for 5' at RT; dried at 400° F		ı		8
16-1	30 gm/L KyCr2O, plus 20 ml/L 98% H2O,	T.	vs.	ruming water	ន	ş	က	Washed once			,	80
71-1	None					92	120		Fabric brown - will not dye	•	ı	20
71-2	None					700	240		Fabric brown - will not dye	,	,	*
71-3						619	1400		Fabric brown - will not dye	ý	,	88
72-1	30 gm/L KgCr2O7 plus 20 ml/L 98% H2SO4	R.	350					Washed once	Fabric very dark		'	ន
For n	For notes see end of table.											

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TABLE VI-4

THE STATE OF THE S

SIUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC⁽¹⁾

NOTES

- See Table I in the second status report on this project dated July 16, 1970 for a description of fabric 6-2 used in these tests. £
- The washing procedure used was to boil the fabric for 15 minutes in 800 ml of soap solution containing 1 gram of neutral chip soap and 1 gram of sodium metasilicate. શ
- See Appendix A for shrinkage test procedure. The % shrinkage is based on the area of the treated sample. ල

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TABLE VII-1

THE STATE OF THE PROPERTY OF T

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

											Shrinkse.	
1	7	Condit	Conditions	Riseing	100	Drying	T. Pare			Treatment .	Orer-ell .	By Thermal
2	Reagent Tem	, de	Temp OF (Mins.)	Medium (Temp of (Mine.)	(Mine.)	Additional Treatment	Romarks	Only	Shrinktye	Exposure (1)
3	40°Bé HNO3		-#-	Hot H2O	6	280	8		•		٠	15
63 -2		ق. ا	Pame as above	~ 			1	Dip in POC' 3 5' wash hot H ₂ O 5'		'	,	:
								ringe in conc NH4OH and dry				
£-3		<u>:</u> 	same as above					Sat with SnCr 2 5' at RT, wash		1	,	z
								hot M ₂ O, neutralise in NH ₄ OH and dry		-		
3		_=	same as above				1	Same as above plus dip again in	•	,	•	*1
								POCY, rines hot H ₂ O,3; dip conc NH ₄ OH, wash 2' and dry 10' at 200°F				
3		-,,				-						:
	H2804 50 50 by vol -		ž	Hot H ₂ O 5'		280	2	•	•		•	2
			ā.	plus NH OH ringe	3						,	;
Z -3		<u>=</u> 1	eame as above				Î	Dip in Poct 3 5' wash hot H2O	•		2	ř;
33	,		asme as above				1	ringe NH OH Dia in cone RC sat with SaC	•	•	,	*
		\			_ 			5' ringe meutralise in NH OH				
3		<u> </u>	same as 64-1				ſ		,	,	,	#
8.78			- - 64-3 pe	same as 64-3 plus dio in POC. 5'	15		1	ŧ	•	,	•	я
•			seutralize wash and dry	and dry	, n		•					
65-1	same as above30		ž	Hot H ₂ O 5 plus	. —	280	15	1		•	•	2
			z-	NH OH rings								
65-2	same as above -10	<u> </u>	6	same as above	ove			1	•	•		#
5 3	Boiling POCC 31' bot plate to drive off POCC 3 dip in methe	- 장	to in methano	and dry on bot plate - neutralize NH, OH	late - neuti	ralize NH	но	1	ı	•	23	2
1	Dry fabric into POCt , at boil dry on bot plate	plate		•		,	,	ı		,	,	\$
6-3				,	-		•	•	1		1	s,
		1						The state of the s				

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TABLE VII-2

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

							•		1	4 Shrinkage	
Run No. Reagent	Temp of	Conditions Time Time	Medium	Time (Mins.)	Temp of	Drying Time 1p of (Mins.)	Additional Treatment	Remarks -	By Treatment Only	Over-all Shrinkage	By Thermal Exposure(2)
\$-50			same as above except for damper fabric	e except fo	daniper (abric	•	-	-		ş
65-7 ethylese glycol beat to 300°F	,	•	,			_	•	,	•	•	\$
63-5 ethylene gly col dry then POCf	•	•	,	•	,	,		•	•	•	5 2
									-		
65-: like 66-5 except negralized in	•	,				,		•	'	•	58
66-1 POCC at RT16 box box plate	,								,		
			,	ı	·	ı 	•	•	,	•	9
66-2 same us above except methanol dip plus wash in cold H,O before drying	dip plus wash i	in cold H,O	before drying			1	•	•	•	01	6
66-3 Dup in POCI plus cone HCf at 150°F.2'. Pad on methanol, dry, neutralize with NH OH, wash with hot H2O	150°F.2'. Pad	on methano	Ldry, neutralize	with NH 1	H, wash wif	th hot H ₂ O	•	•	,	•	\$
66-4 Dip is mixture of POCf ₃ and methano; at 150°F for 2', dry	theno: at 150°F	for 2. dry.	cwash in meth mol	5			•	,	•	•	2
67-1 1% gly cod 85% H3 PO4	212	•	'	•	•	•	,	•	'	•	3
67-2	- same as above except SF	except 5% i	801'n		- - -		•	•	1	3-1	s
67-3	. same as above except 10% sol'n	except 10%	n'los			- ,	•	1	,	2	19.5
Like 67-1 except after treating with 70% wrea solution a	with 70% area	solution at	it boil, rinse with dil NH4OH, wash, and dry	DT NH IP	H, wash, a	ad dry		•		•	\$
Like 67-2 smoopt after treating with 70% ures solution	with 70% ures		it boil, ringe with dil NH4OH, wash, and dry	OF NH (FD	H, wash, a	and dray	•	ı	•	•	9
	with 70% area		it boil, ringe with dil NH OH, wash, and dry	OF HN IP	H, wash, a	ad dry	•	,	•	•	35
67-7 5% by vol 85% H ₃ PO ₄	212	'n	•		35	5 days	•	Sample turned brown	1	•	\$
67-8 1% by vol 85% H ₃ PO	212	vo	•	,	392	5 days	Boil in 24 Alconox and	•	,	•	*
- CS H 25 anian annia as an	_ *						đ,				
				_				1	•	•	;
67-10 same as above using 10% H ₃ PO, sol'n 4	u, 104 *							•	1	•	a
66-1 PCr ₃	212	5	•	•	382	009	•	•	,	•	÷
68-2 Equal parts POCt 3 and PCf 3	212	•		•	192	90	1	•	'	•	2
60-3 PC! 3	212	ر. د	methanol plus H20	H ₂ 0		,	•	,	,	•	53
68-4 A Cr. + PCr. then	212	9	rinse methanol plus				Het POC! for 1' after				
		· ·	H ₂ O rinse				treatment with AIC(3		. =		
							etc. and before dip in				
							methanol and ringe				

See end of table for notes.

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TABLE VII-3

Windle-Statement of the Estatement

THE PARTY OF THE PROPERTY OF T

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FARRIC (1)

Part											, -	T Shrinkage	
Hot 3 Histaph of rinse 10 10 10 10 10 10 10 1	2 8	Reagent	Condi Temp ^o F	Time (Mins.)	Medium	Tine Office	Temp ^o F	line (Aline,)	Valitional Treatment	Remarks	By Treatment Only	Over-all Shrinkage	By Thermal Exposure (2)
Repeat of 69-1 Repe		H_3004	071	,	Rinae pH o	rinse	392	10				Ŀ	ន
Seperal of G9-1 except neutralize with MH_OH io PH 9.5 Cold 1007; chlorous/phonic sole Same as above except neutralize in case NH_OH Cold 1007; chlorous/phonic sole Same as above except neutralize in case NH_OH Cold 1007	8 -7		Repeat of 69-1										
Cold 1007 chlorowalphonic scients Cold 1007 chlorowalphonic scients Same as above except neutralize in case NH ₄ OH	69-3		Repeat of 69-1 exce	pt neutralize	with NH OH	to pH 9.5							
Same as slove except neutralize in case NH ₂ O 2	Į		Cold 100', chloroeul	Iphonic acid						Fabric dissolves	<u></u>		
Same as above except neutralize in case Nijoli 300 15 185 5 300 15 Britile char 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	\$-5	100% chloroculphonic acid plu PCC 3 1 9		*	How H ₂ O	64		•	1	•	•		5
Same as 60-5 except	Ĭ		Same as above exce	pt nentralize	in case NH	- = -	1	ı	,	•	•	,	22
185 5 300 15 Britite char 2	6		Same as 69-5 except				1	,	,	•	•		z
185 5 -	1-11	20mf L 65% H ₃ PO4	185	s,	ı	•	300	35	ı	Brittle char	•	•	÷
### 185 5 - 1 100 15	11-2	Some / L 85% H ₃ PO ₄	185	'n	•	•	900	15	,	Brittle :her	•		2
offer 15 - Flexible char - offer - 300 15 - - offer - 300 15 - - (3) - 500 15 - - (3) - - - - (4) - - - - (5) - - - - (4) - - - - (4) - - - - (5) - - - - (5) - - - - (6) - - - - (7) - - - - (2) - - - - (2) - - - - (2) - - - - (2) - - - - <td>11-3</td> <td>100 m/L 85% H₃PO₄</td> <td>185</td> <td>v</td> <td>,</td> <td>,</td> <td>300</td> <td>15</td> <td>,</td> <td>Flexible char</td> <td>•</td> <td>8</td> <td>11</td>	11-3	100 m/L 85% H ₃ PO ₄	185	v	,	,	300	15	,	Flexible char	•	8	11
treatment 1-2 washed after 300 15 300 15 treatment - 300 15 300 15	7	same as 71-1 washed after treatment	,	ı	•	•	8	13		Flexible char	•	•	ŧ
treatment of the except best treated after	71-5	same as 71-2 washed after treatment	•	•			006	15	,	•	•	,	3
Asses as 71-4 except beat Treated before washing (3) Same as 71-5 except beat Treated before washing (3) Same as 71-6 except beat Treated before washing (3) Same as 71-6 except beat Treated before washing (3) Same as 71-6 except two	4-F	same as 71-3 washed after treatment	•	•		,	88	15	,	•	•	•	ţ
State as 71-5 except best treated before washing (3) asme as 71-6 except best treated before washing (3) same as 71-6 except two washes (3) asme as 71-6 except two asme as 71-6 except two asme as 71-6 except two	7-11		,	,			Š.	15	,	•	,	•	88
atme as 71-6 encupt beat treated before washing (3) same as 71-4 encupt two washes (3) same as 71-6 encupt two same as 71-6 encupt two	#-E	same as 71-5 except heat treated before washing (3)	,				8	15		•	•		23
	8-H	same as 71-6 except heat treated before washing (3)	•	,	•		200	15	,	•	•	,	01
	71-10	same as 71-4 except two washes(3)	•	•				•	,	•	,	•	ŧ
	11-11	same as 71-6 except two	•	•		•		•	1	•	,	٠	50

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TABLE VII-4

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

										ĺ	% Shrinkage	
3 %	Resgrat	Conditions Time Temp ^o F (Mins.)	fime (Mine.)	Medium	Fine (Mire.)	Temp ^o F (Mins.)	ing Linie (Mina.)	Abilitional Treatment.	Remarks	By Treatment Only	Over-all Shrinkage	By Thermal Exposure(2)
i	71-12 Same as 71-7 except two washes			•			-	•	•	,		20
7-22	72-1 Soak in 507 ures - 857 H ₃ PO ₄ sol'n	91	2	,	•	017	01	•	•	•	•	8
12-27	Same as above	···				410	92		•	•	•	3
ţ	72-3 Sat solution of diammonium phosphate	,	•	•		014	25	•	,	•		\$
Ë	73-4 seek in 10°; H ₂ 80 ₄ ½ hour 70°C wash & then in solution 10 mi H ₂ 0.	•	•	•	•		•	•	•	•		3
	10g area, 10g 377 HCHO											
1-52	73-11 83% H ₃ PO ₄	220	13	337 ures solution at 150 F	-	220	120	•	•	ı	S	*
36	99 73-31 50 m: 1.65% HyPO4	921	13	33% ures solution at 150 F	-	220	8	•	•	•	2	\$
74-1	74-1 120md 'L 85% H ₃ PO ₄	176	v	•		ş		•	•		a	•
72	1							neutralite in NH ₄ OH at bed	ı	•	•	(P
P -2	4 same as above but washed in NH OH a second time	M. HO	econd time									8
74	10% H ₂ 504	241	"	Conc NH ₄ OH two washes	¥			,	Flexible char	,	•	61
2 2	74-6 POCI at 312°F for 10° plus 10°? R,90 at 178°F for 5°				,	8	92	dip in POCi 3 at 212°F for 2' .	Florible char			= :
												l

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TABLE VII-5

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

			į	č						T. Shrinkage	
% % 9.	Reagent	Time Time Temp ^o F (Mins.)	Time (Mins.)	Medium (Mins.)	-	Tenip ^o F (Mins.)	Viditional Treatment	Remarks	By Treatment Only	Over-all Shrinkage	By Thermal Exposured)
1-1-1		same as 74-6	, e			Î	dip in POCI 3 for $\frac{1}{2}$ hr	Flexible char	•		11
¥-1;	74-5 chlorosulfonic acid in POCI ₃	110	22	neutralize in II,0	977	9	•	•	ı	•	•
14-9		same as al	bon e except	same as above except no neutralization					-		_
73-1	73-1 107 H ₂ SO ₄	167	17	25m/ 1. conc NH ₄ OH for 15' just before shrinkage test	# 7	· ·	10', II, IV), S' at 185°F after drying and then 'rving repeated'	•	•		• a
13-2	73-2 107 H ₂ 80 ₄	191	so.	same as above	•		30° in POCI ₃ st.23E ⁰ F after drying and then drying repeated	,	ı		8
រុំ 37	73-3 POCE 3	212	ဓ္က	same as above	ng:	ı;	10. H ₂ KO ₂ fluid dring plus POCL ₃ at 212 ⁰ F plus dr.ing	•	•		£
75-4		same as above -	900			1					5
2-51 S-51	75-5 10" chlorosulphonic acid in POCI3	212	6	same as above	284	v.	167 H ₂ SO ₄ plus drying	•	•		8
,; ,;	-	same as above	****				10': H ₃ PO ₄ plus drying	•	,		\$
7-52	73-7 107 H ₃ PO ₄	336	0.	same as above	284	17	10', 11 ₃ PO ₄ plus dip into 50'; urea solution & dry		•	•	\$
75-8		same 24 above	• • • • • • • • • • • • • • • • • • •				POCI ₃ at 212 ⁰ F for 15 min plus 10% H ₃ PO ₄ plus dip into 507 urea	•	•		;
75-9	73-9 POCI 3	212	8	eame as above	584	'n	107 H ₃ PO ₄ plus dry plus H ₃ PO ₄ dry plus 507 urea dry & dip	•	•		‡
75-10		same as above	XOVe			7					

See end of table for notes.

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TABLE VII-6

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

										i	2 Shrinkage	
		Conditions	lone	Rittsing		Drying				By	By	
9 9	Reacent	Time Temp F. (Mins.)	Tine (Mins.)	Medium	(Mins.)	(Mins.) femp	(Nine	Additional Treatment	Remarks	Only	Shrinkage	Shrinkage Exposure(2)
-	10% chlorosulfonic acid	. 611	17	unh d ethuol	rinse	316	20	•	Moderately Regible	-	•	8
76-2		- H. Wash befor	e chrinkage	age test	,	,			•	•		\$
76-3		200	1	nahyd ethanol	rinse	236	ន	•	Moderately flexible char	1	•	96
792	same as 76-3 above except NH OH wash before shrinkage test	H wash befor	e shrinkage				-	•		•	•	t
76-5	same as 76-1							plus 5' in 10° chloro- sulfonic acid in pyridine at 200°F plus 30° beat in at \$20°F	Fairly flexible char		•	8
76.4	same 2276-5 above except NH ₄ OH wash before shrinka	H wash befor	shrinkage					•		•	•	\$
7-91	8 mi chlorosulfonic acid 8 mi yyridise 2 mi POC 3	200	in.	anlayd	rinse	836	2	wash in NH ₄ OH before shrinkage test		,	•	ន
- 2	5 mf chlorosulfonic acid in 100 mf POCt 3	200	8	NH, OH wash	after beating	527	8			•	•	\$
38												

NOTES

1. See Table I for a description of fabric 6-2 used in these trets.

2. See Appendix A for shrinkage test procedure. The % shrinkage is based on the area of the treated sample.

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYPENZIMIDAZOLE FARRIC (1) TABLE VIII-1

		County		Binetin		j	9			- [S. Shrinkage	
22	Reagner	Temp ⁰ F (Mins.)	Time (Mine.)	Medium	Time Mins. 1	Temp ^o F (M	Time (Mins.)	Additional Treatment	Remarks	By Treatment Only	Over-all	By Thermal Exposure
*	3% solution of P.O. in 85% H.3PO.	2	51	water. Ne.CO.		905	9			11		2
**	same as above	2	8		same as above	.pove				•	•	~
2-32	Same as above	8	8		same as ahove	Phove		•	•	8	•	
\$	same as above	18	'n		same as above	apove.		•	•	•	•	ş
3	same as above	100	15		same as above	bove		•	•	2	•	8
3	same as above	8	ጸ		same as above	above		•		8	•	\$
1-15	20 mt/L 855 H3PO, plus	2	1/2	cold water	15	•		•	fairly resistant to burning	,	,	z
<u>:</u>		500	2		,	200	ç	treated for 10' in boiling KHSO ₃ solutior and dried again 10' at 300°F	no significant effect on tensile strength	,	,	*
3	66-1 56g/L K ₂ Cr ₂ O ₇ is 65% H ₃ PO ₄	700	-	Na ₂ CO ₃ plus		500	2					
				K ₂ 80 ₃				ı	tensile strength 67 lbs/in.	•	•	=
<u>.</u>	865 H ₃ PO ₄	267	8	water. Na ₂ CO ₃ sol'n, water		200	2	•	fabric practically destroyed, tensile strength 9 lbs/in.	,	,	
70-1	same as above	210	e,	same as above	,	200	្ព	•	bearily stained	•	,	91
76-3	same as above	210	\$	same as above	•	200	2	•	unerven staining	•	•	#
Ť	State as above	325	g	same as above	,	200	2	•	•	•	,	22
1-12	same as above	225	8	same as above	,	200	2	•	fabric not stained			*
ţ.	73-2 4g K ₂ Cr ₂ O ₇ in 400 mt 86% H ₃ PO ₄	228	8	same as above	,	203	2	•	fabric did not burn fabric unevenly stained	•	•	2
2-2	same as above	225	ફ	same as above		200	2	•	(abric badly stained, did not burn	,	ı	8

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TABLE VIII-2

STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC (1)

		1	į			į	ļ				S Strieter	
g ś	Rogent	Tomp [©] F (Min	.:	Medium	Time (Mine.)	Kedtum	Medium (Mins.)	Additional Treatment	Remarks	Trees Selfy	Over-all Shrinkage	By Thermal Exposure
ž	⁷ од ⁸ и 3.98	210	3	wader Na, CO,	\ '_	8	e —	4	tensile strength 50 lbs/in.	·	Ŀ	-
74-2	SE K_CR_O, is 400 mt 85%	ħ	•	same as above		8	2	•	tensile strength 60 lbs/in.	,	'	
2-3	K3FO4								•			
) !		ä	\$	same as above		28	<u>e</u>	•	tenalle strength 20 lbs/in.	,	,	•
7+7	855 H3PO	22	3	same as above	•	88	2	•	tensile strength 75 lbs/in.	,	,	
74-5	Ses H3PO	210	8					•	tennile strength 80 Be/in.	,	'	, 1
į		\$22	ន	same as above		§	9	•	teasile strength 60 lbs/in.	•	•	,
3	50-50 by volume 40° Be HNO ₃ and 36% H ₂ 50 ₄	۰	2	2-1/2% NB ₃ in water	51	8	2	•	ı	\$	•	5
ž		•	2	same as above	55	8	2	•	•	2	•	a
<u>2</u>	25:75 by volume 40° Be HNO ₃ and 96% H ₂ 50 ₄	#	ž	KOH sol'n	35	§ 	2	HgO rimee 51	risse solution seld	\$	•	a
2-2	same as above	32	22	buffer sol's	51	§	2	same as above	rinee solution acid	\$		#
82-3	same as above	23	82	KOH sol's until pH of fabric is water = 6	9 m	438	15	H ₂ O rime	•	•	,	•
ž	1	same as above					1	after H ₂ O rinse boiled in Altanox for 10 mins.	,	•	,	=
144	25:75 40° Be HNO ₃ and 96% H ₂ 8O ₄	21	15	WH4OH sol's	₹ 8	\$ 	22	rines in H ₂ O and test pH of sample	fabric yellow after rines	•		2
		same as above			# 1/2 1/2	\$	15	•	see above	,	ı	:
		same as above			to pH 7-1/2	***	12	•	fabric turns brown when	1		2
		same as above			→ 6 pH	\$	22	•	ass above	,	•	*
		same as above			¥6 pH	2	=	•	see abore			*

TABLE VIII-3

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STUDY OF TREATMENTS FOR REDUCTION OF THERMAL SHRINKAGE OF POLYBENZIMIDAZOLE FABRIC

NOTES:

% thermal shrinkage is calculated on the area of the treated sample. The thermal shrinkage test procedure is given in Appendix A. The See Table I for a description of 6-2 the fabric used in these tests. £

TABLE IX-1 STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT $^{(1)}$

																\$ }	THE STATE OF	. 1				
:	ist Treatment		Drying	Brinkage		2nd Treatment Temp Time		Drying Temp Time		읡	3rd Treatment Temp Time	atment np Time	Brinkage	Drying Temp Time	Ğ	Mrinkage NH OH	i ege	A REC	Temp Time	Tonaile Erength Ibe/in.	Thermal Brinkago(3)	Ploutbilly of Ch.
Solution	0	- 1	O S	-	Solution SG CIO.SH	. E		<u>د</u> ع	1	13.6	Solution C	110 10	3.5	8.	1	#.4		5 FE 8	er er		37.4	Bride
ĕğ	10 200 100 100 100 100 100 100 100 100 1		- 		in Poct ₃					- 7:	<u> </u>		9,0			11 3 11	11.5 \$ 4		\$	•	31.6	
	<u>:</u>									21.8			27.2			37.9	11.5 9.5	*	\$ 1	•	37.5	
j	: :			:						17.6			27.2			2 ° 2	•	3 220	•• ~	=	40.0	
1	io			5.1						15.9			25,9			31.0	11 45 %.	*		•	.	
	1			:						2,5			18.2			22 3 11	11.4	:		•	41.2	
	: E			÷	-	-	- -		_ a				21.8			22 0 22	11.40	2		5	7:2	
1		<u>.</u>											23, 1				11.45			2	41.3	
11 -	Tree and			:								R	22.1			3 ·	• • •		_	=	*	
• •				2								Š.	16, x	-		25.3	ê 2+ II	c c		2	41.8	
				•								<u>.</u>	71.x		-	1 5 27	st 12		-	ī	÷.0	

TABLE IX-2 STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT $^{(1)}$

	Observations	Fabric darker than any of above except no apole 1 to 3	Spots patters of alum. abelf				foure black deposit on fabric after treatment is POCI3 ClOyde, Fabric somewhat heritie after treatment, also quite date.	Considerable black deposit on fibric after POCI ₂ CIO ₂ Mt treatment. Fabric brittle after treatment and online dark.				
	Flexibility of Char	Britte									وسز →	golf.
	Thermal Shrinkage(3)	47.5	60.0	42.6	42.3	40.0	32.6	0.07	37.5	38.	20.0	16,3
	Tensile Rrength 15s/in	2	3 2	92	82	3	89	•	•	•		•
	Temp Time	3066	š —						62		Panilita	201
50 H	PH of Rinsed Fabric	**	n •	:	:	:	ć.	6.	•	e.		
Rinsing in 5. of NH ₄ OH	Nii oi Affer Rines	11.49	:	=	=	11.5	•	11 45	:	=		1
	Brink ye.	72.4	27.	22	22	ş	17. •	20 01	23.8	2	27.9	7 92
	Drying Temp Time	2	8						23			
	S rinkage	_	2.1	2	•	•	19.1	18.5	21.8	21.0	23 2	2
	Time	2	8 –									
	3rd Treatment Temp Time	*0										
	Time Temp Time Seriakage	•	19.3	13.9	18.2	4 .2	5.3	18.2	50. 50	\$.0\$	3	;
	Time.	a	8-						 g -			
	g F.	S.	<u>=</u> -						 8 -			
	F 3	2	2-						 2-			
	1st Treatment Temp	E	=	211	711	***	112	=======================================	=======================================	21	112	:
	Polerice	Same but 112 from sol's		 					- 1 .			
	1 §	(2	2	2	=	5	=	- 1.	8	ដ	:

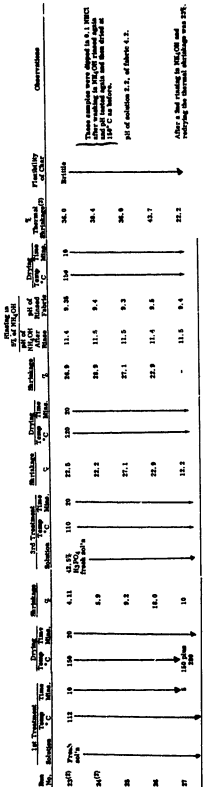
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TABLE IX-3

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STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT $^{(1)}$



HOTES:

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Clock sample 73 (see Table D) mad for all russ. Except for samples 1, 2, and 3, before drying the samples after treatment is H3PQ, they were assistances and cross and covered by a "requer of 4.32" history plate and placed on the blumians was able in a 270 plate in the drying the red or as at all times. Before that they were removed with the sample and were allowed to cool. For \$13 a \$" equare alumin was plate, and for \$14 and after, the acressa were associated between two 3" equare aluminam plates .035" thick. Except for the 7 best shrinkage all 7 shrinkage.

The cloth for these runs was dried for 18 mins. at 156°C before treating. € €

See Appendix A for test precedure. The S thermal shrivkage is bused on the area of the trested sample

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Table X-1 study of chlorosulfonic acid foci $_3$ treatment $^{(1)}$

Fabric Let Trestment Let Drying E			p Time Temp 7 (Mine.) C (1)	Top 7		M 53	e c	Heading i- Temp I	desting inp Time Strink C (Mins.) ages	量品	2nd Treatment Temp Ti	omb C C	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Sarink - Tagno Time Sarink	2nd Drying Temp Time C (Miss	Zing Time (Miss.)	전 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Stantog to SS NH OH PH of the ringed fabric	ard Drying Temp Tiu	89	Hatt Berieb	Flord- Mility	Observations
yes 5% Ct H3O ₃ II2 10 I50 10 5.1 280 in POCt ₃	150 10 5.1	150 10 5.1	150 10 5.1	5.1	5.1		280		81	7.56	42.5% H ₃ PO ₄	911	2	12.5	80	2	25.5	:	150	2	17.0	Ę	Temperature of oven used the heat fabric of minutes @ 200 may have been genetiderably higher than 200
yes 5% Ct H3O ₂ 112 10 150 10 7.90 280 in POCt 3	150 10 7.90	150 10 7.90	150 10 7.90	10 7.90	10 7.90				84	9.07	42.5% H ₃ PO ₄	91	2	20.4	200	2	36 . 10	6.2	921	2	27.5	27.5 brittle	
no 5% Ct HNO ₃ 112 10 150 10 20.00 280 in POCt 3	112 10 150 10 20.00	112 10 150 10 20.00	10 150 10 20.00	150 10 20.00	10 20.00				81	34.30	42.5% H ₃ PO	2	2	34.3	200	2	3.3		23	2	31.3	31.3 brittle	Looked dark ad hed some black spots
20 2% C. H. H. M. 150 10 15.25 280 28 P. C. M. 1 150 10 15.25 280	112 10 150 10 15.25 }	112 10 150 10 15.25 }	10 150 10 15.25)	150 10 15.25	10 15.25				N	18.40 A	42.5 H ₃ PO ₄	91	2	20.0	88	2	23. 80	6	150	2	3.4	34.4 brittle	Looked dark had some black spots
112 10 150 10 7.70 280	112 10 150 10 7.70 280	112 10 150 10 7.70 280	10 150 10 7.70 280	150 10 7.70 280	10 7.70 280	280	280	**	84	10.10 A	42.5% H ₃ PO ₄	ន្ព	2	12.4	200	2	19.25	9.2	33	2	6.0	60.0 brittle	
yee 10% C4.EBO ₃ 112 10 150 10 6.00 280 2 in POCC ₃	112 10 150 10 6.00 280	112 10 150 10 6.00 280	10 150 10 6.00 280	150 10 6.00 280	10 6.00 280	780	780	84		7.10	42.5% H ₃ PO	91	2	15.5	200	9	18.65	6.3	120	2	40.0	10.0 brittle	
34 300° 300° - 400° 2	3000	,006 300	,006 300	, 300	- 300	- 3000-400	300 -4002	% م		,		,		,			•		,	,	3		
yes 6%Ct BFO ₂ IF. 10 150 10 4.50 280 2 in POCt 3	IF. 10 150 10 4.50 280	IF. 10 150 10 4.50 280	10 150 10 4.50 280	10 4.50 280	10 4.50 280	280	280	81		4.50	42.5% H ₂ PO ₄	음	2	19.3	802	. 01	26.50	9.2	32	2	8.	36.0 brittle	Some dark brittle arear ther appear after H ₃ PO ₄ treatmen and over day
yes 6% Ct HBO ₃ 112 10 150 10 4,90 250 2 in POCC 3	10 150 10 4.90 250	10 150 10 4.90 250	10 150 10 4.90 250	10 4.90 280	10 4.90 280	361	361	89		5.80	42.5% H ₂ PO ₄	음	2	21.9	200	2	28.40	9.2	3	2	37.6	37.6 brittle	eee spore
yes 5% Cl HBO 112 10 150 10 3.50 300 2 ta POCl 3	000 03.50 00 051 01 21	000 03.50 00 051 01 21	10 150 10 3.50 300	10 3.50 300	3.50 300	300	300	**	81	6.40	42.5% H ₂ PO ₄	유	9	24.1	200	2	26.20	9.0	32	2	31.5	37.5 brittle	see above
yes 5% Cf HBO ₃ 112 10 150 10 3.50 300 2 In POCf 3	10 150 10 3.50 300	10 150 10 3.50 300	10 150 10 3.50 300	10 3.50 300	3.50 300	300	300	8		6.20	42.5% H PO	91	2	17.6	200	2	22.00 9.2	9.3	33	9	36.1	36.1 brittle	Somewhat darker than 35,36,37,40,41,42 after rguffle furnace freatment at 287 - 340

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TABLE X-2

STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT $^{(1)}$

	Observations	see commen; on 35	see above	see abore	This was beated less than 10 minutes in ${ m H_3^{PO}}_4$	Some dark brittle spots which appeared after H. PO, treat- ment and oven dify. Both therice	ware mightly grows and the war- backing at 400 C but of berrelee locked of vy. Both samples were limad a second time is SS NE, OR and restricted and seals the themal	harinking was 19%. After a second NE Off rises and redrying the thermal shrinkage was 33%.		The weight change of the two sal-pies after let treatment +31% +27%	after 2nd treatment +121% +124%
	Flexi- Hilty	37.6 brittle	35.9 brittle	29.9 brittle	35.9 brittle	ti ot	ų į		#o#	40.5 brittle	40.5 brittle
į	Arrink(2)	37.6	35.9	29.9	65.0	10.0	10.0	21.3	19.0 soft	40.5	40.5
N.	np Time Shrinkz)	01	2	2	2	2	2	91	9	2	0
밁	E	150	120	120	150	3 2	150	10.6(5) 150	150	82	150
Rineing in SS, NH, OH	ringed fabric	9.3	4.0	6	9.6	9.4	9.4	10.6(5	e.	ø.	e. 8
28	Sprink-	25.2	24.2	26.0	26.5	23.2	25.8	•	•	30.5	26.3
Ä	Time :	2	2	2	9	2	9	. 2	6	9	2
2nd Drying	Temp C. C.	58	200	200	200	200	200 10	200 10	dling fo	200 10	200 10
**,	Shrink- Tgmp Time Shrink- rinsed age C (Mins.) age fabric &	17.2	19.8	22.2	20.1	21.0	6. 6.	12 N H 20 N 12 N	ater at bo	23.2	24.3
뉨	Temp Time C (Mins.)	2	2	9	2	2	2	vs	» 20 00	2	2
2nd Treatment	Temp C C	£	91	81	91	8	011	110	ric in 2	110	8
T Luz	Solution	42.5% H.PO.4	42.5% H P O	42.5% H_PO	42.5% H ₂ PO	42.5% H 204	10.30 42.5% H ₂ PO	neutralized with NH, OH 2 dried and 42.5% retreated H ₂ PO ₄	5% dye on weight of fabric in 200 cc water at boiling for 5	11.90 42.5% H ₂ PO ₄	42.53 HPO4
	o Time Shrink- (Mins.) age	5.60	3.50	6.20	3.70	7.30	10.30	ith NH cried and streated	e on we	11. 90	16.10
J	Time (Mins.	8	69	81	84	84	84	A F G L		*	64
Heatin	Temp Dines Temp Time Shrink- Temp C (Mine.) C (Mine.) age C	320	320	£	350	400	0	340	fast blue	380	280
	Shrink #6	9.1	3.5	e.	3.1	5	9.1	•	lyamid	2.4	1.8
,	Time (Mine.)	2	2	2	2	2	2	ĸ	odpes	6	w
lst Trestment lst Drylag	Temp	8	150	150	130	150	22	85	lon blue	120	120
	Time (Mins.)	2	\$	2	2	2	Ce.	w	ig in	14	'n
etine	ရှိသ	£.	A	ਬ •••	ם **-	. 2°	21	21 E		a 2° ≈	5 T
H	Fabric Rus dried 10 No. @ 150 C Solution	SE CUESO, 11:) In POCU	SS CC HSO, 112	SS, Ct HSO, 112	SECURSO, UZ	SECUENCE IIS III POCE 3	5% CV ESO, 112 in Poct 3	5% Ct H5O ₃ 112 in POCt ₃	\$29 after immersion is maxilion blue sadpolyamid fast blue minutes, rinsed and dried.	5% CV HBO ₃ 112 In POCY ₃	5% Ct H803 112 in POCt 3
	Pabric entrado 150 C	2	Ĕ	Ě	ž	£	ž	ž	428 at minut	Ĕ	ķ
	Rus No.	8	\$	7	3	3 46	\$	\$	\$!	\$

TABLE X-3

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STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT $^{(1)}$

15

NOTES

(i) The sampées of cirth '2-1 (ass Table I) serve heated in the oven after H. FO, treatment anndwithed between sheets of 40 mesh stainless steel serven and 6 " x 6" x 1/32" alumisum plates:
a 2070 gram had weight. The absets and wrights were in the oven at all times. Sabrinkages, except for the best shrink test, are based on the original area of the fabric.

(2) Samples in brackets were all treated to the same fatch of C ℓ HSO $_3$, POC $_3$ solution.

(3) See Appendix A for hest procedure. The Sahrithage is based on the area of the treated sample.

(4) Same relution as that used for samples 15 to 42.

(5) pal of time selution. This is forest than normal. See Table VII.

Table XI-1 study of chlorosulfonic acid Pocl_3 treatment⁽¹⁾

Plendalite	Shrinkage (3) al over	brittie	brittle	flexibia	flexible						very brittle	very brittle	brittle	brittle	brittle	semi-brittie	brittle	brittle
į	hrinkage (3)	38.7	37.5	7.5	3.5	51.3	2 .8	47.5	8.8	25.8	87.0	2.0	92.0	32.0	7.12	18.8	30.0	23.3
	Observations	Fabric slightly darkened	Fabric darkened more than I	Fabric black, flexible	Fabric black, flexible	Fabric light	Fabric light	Fabric darkened about like t	Fabric light similar to 5	Fabric light similar to S	Fabric light brown same as 6	Fabric light brown about same as 10	Fabric dark brown about same as 7	Fabric light brown but darker than 10	Fabric light brown same as 13	Fabric dark brown about same	Fabric medium brown about same	Fabric a little bit darker
	L.O.1.(2)	•	•	ı	٠			,	•	•	0.36			0.35	•	,	- HD	
	Stiffness				•		•		•		ş	Š	Slightly s	ŏ	ĕ	Stiffenex	Slightly stiff	Stiffened
	Color		•	•			•	•	•		ŧ	₹	Slightly darker Slightly stiff	ğ	S X	Slightly darker Stiffened	ŧ	Slightly darker Stiffened
	's weight add-on	•	•	•	1	•	•	•	•	•	2.04	3.10	2.40 \$	6,95	6.40	10.00	11.50	10.50
	Time (Nins.)	8	30	90	30	20	\$	8	40	'n	2	20	2	8	20	&	20	20
Drying	Temp ^o C (Mins.)	35	340	90	9	320	320	320	360	360	320	340	340	320	340	340	320	340
	Time (Mins.)	s	20	ın	20	40	n	ø	w	ø	s	w	w	ю	10	o	so.	17
Treatment	Time Temp C (Mins.)	F.	RT	RT	RT	R	RT	RT	TX	9110	RT	RT	R T	S	8	ទ	911	2
1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	ia POĆt solution 3	so.	10	so	•	*	*	**	**	ĸ	•	**	₩,	₩	w	w	w	'n
Dried	at :20% (Miss.)	v	w	•	•	•	•	'n	•	•	N)	so.	••	•	w	v	w	s
	₹ .é	_	~	•	*	•	•	•	•	•	2	=	n	2	3	22	*	11

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TABLE XI-2

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MANAGEMENT STATEMENT CONTROL

STUDY OF CHLOROSULFONIC ACID POCI $_3$ TREATMENT⁽¹⁾

	1							
	of char	brittle	brittle	brittle	semi-brittle	semi-brittle	seed-brittle	sond-brittle
	Shrinkage (3) Flormaliny Shrinkage of char	17.5	90.0	54.7	26.0	s 21	14.A	2e 14.5
	Observations Sh	Dark because of speet POC! 3	Light brown good texture	Light brown good cleth-like texture 34.7	Dark brown good cleth-like texture 26.0	Medium-brown tenture somethat	Medium-brows tenture somewhat	Dark brown tenture somewhat coarse 14.5
	L. O. I. (2)	•	0.35	0.33	0.34	0.39	0.36	
	Color Stiffness L.O.I.(2)	Stiffened	OK	ŧ	Darkened Slightly stiff 0.34	Stiffened	Stiffened	Darkened Stiffened
		Darkened Stiffened	¥	¥	Darkened	Slightly dark Stiffened	Slightly dark Stiffened	Darkened
	% weight add-on	•	6.50	€.00	9.20	16.50	17.00	20.50
*	Time C (Mins.)	2	2	2	2	2	2	2
Deyta	Temp ^o C	3	320	340	8	Š	3	9
	Time Temp [©] C (Miss.)	w	•	6	ø	•	10	10
-	Temp C	2	3	3	3	97	90	91
1. T. O. 28.	is PCCt 3	w	2	2	2	2	2	2
Dried	# F	*0	so.	s	10	•	•	•
	1 ź	2	2	2	=	#	#	ጃ 49

NOTES.

(i) Cloth 73-1 (see Table i) used after drying 5 minutes at 220°C before treatment. Temperature in drying oven questionable for runs 1 to 9 and $\overset{+}{\sim}$ °C thereafter.

(3) LOI (finiting caypes index) is minimum percent caypen by volume in a nitrogen caypen mixture in which the sample will burn.

(3) See Appendix A for procedure. Percent shrinkage is based on the area of the treated material.

it was noted that when preforming the flame shrinkage test on the 2 X 2 equares that a lesser amount of visible fumes were emitted from the fabric when the shrinkage was low, 1. e. the more the

Mark

TABLE XII

Properties of 7 Yards of Fabric Submitted to Air Force for Further Evaluation Treated for Evaluation by Technical Monitor Using Optimum Process(1)

Sample Description(2)	Tensile Strength(3) lbs/in.	Elongation at Break(3)	% Shrinkage Due to Processing	Weight Add-on(4)	% Moisture Regain 50% RH 65% RH	sture ain 65% RH	Thermal Strinkage (6)	L. O. I.(7)
Untreated	88	13	1	ı	7	7 12	7 9	36
Sample #1	66	27	ı	ı	ı	ı	93	53
Sample #2	95	26	i	i	i	ı	31	i
Sample #3	68	27	14	2	œ	14	27	53
Sample #4	92	56	ı	ŧ	1	1	29	i
Sample #5	89	27	ı	ŧ	1	1	27	28

NOTES:

- room temperature, residence time 40 seconds per 2" length, solution feed rate 33 cc/min. The leader section broke at the beginning of the run, therefore, the first 3 feet of fabric was pulled through the 3 banks of 250 watt infrared lamps, o lamps per bank, setting of 2500 watt radiant heater 5 (with this together and treated in the full scale apparatus shown in Section 3.2. The fabric drying section had setting one end of the heater was hotter than the other), solution 5% chlorosulfonic acid in POCl3 at After treatment, the cloth was heated for 1 1/4 hours at 300°C. The oven temperature dropped to Two pieces of cloth 118-1 described in Table I, one 2 yards long and one 5 yards long, were sewn reactor by hand. The temperature of the heating solution increased during the run to about 80°C. 285°C when the fabric was introduced. $\widehat{\Xi}$
- (2) The sample locations were as follows:
- 1 initial 3-foot section pulled through by hand
 - 2 end of first 2-yard section

TABLE XII (Continued)

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NOTES: (Cont.)

- 4 same as 3 except taken from the opposite side of the section where the radiant heater was the hottest - beginning of second 5-yard section taken from side where radiant heater was at a lower temperature 5 - end of 5-yard section at the same side as 4
 - Fed. Test Method Std. 191 Method 5104, three specimens warp direction only. ල
- (4) Based on original area.
- room at $73 \pm 2^{\circ}$ F and $50 \pm 5\%$ relative humidity. Figures 3 and 4 show the plots of 5 moisture regain and the fluctuations in relative humidity during the period the samples were exposed. The relative Dry weight determined as in Fed. Test Method Std. 191, Method 2600 and moisture regain determined by perfodic weighings until sample reaches a constant weight when exposed in a constant temperature humidity was measured each morning. 3
 - % shrinkage is based on the area of the treated sample. See Appendix A for procedure. 9
- LOI (limiting oxygen index) is the minimum percent oxygen in a mixture of oxygen and nitrogen in which the sample will burn. 3
- Same as note (5) except after drying cloth was conditioned for 20 hours in a dessicator at 78.5°F over a solution of sulfuric acid having a density of 1.27. 8

Section 4 DISCUSSION OF RESULTS

4.1 Phosphorylation

Initial work was conducted with phosphorus oxychloride (Table XIII). This reagent gave non-uniform treatments resulting in thermal shrinkages generally greater than 40%. Moisture present on the fabric appeared to enhance weight gain and aid in the stabilization of the material. Since water decomposes POCl₃ to phosphoric and hydrochloric acids, these acids were examined individually and in combination with POCl₃. Treatments of the fabric with hydrochloric acid were ineffective in reducing thermal shrinkage; but treatments with phosphoric acid were successful. A summary of the effects of various combinations of POCl₃ and phosphoric acid has been selected from Table II and given in Table XI.

Of all these treatments, direct reaction with 85% phosphoric acid at 210°-300°F appeared to provide a treated fabric with the best resistance to thermal shrinkage. In addition, an increase in the oxidation resistance of the fabric was noted - an untreated sample had a Limiting Oxygen Index of 0.36 while the treated sample had one of 0.47. However, significant fabric shrinkage occurred during the treatment process itself. For example, a sample treated 30 minutes in 85% H₃PO₄ at 218°F shrank another 23%. While the combined shrinkages of this sample was still lower than the shrinkage of an untreated sample on thermal exposure along (60-65%), the shrinkage on immersion in phosphoric acid was considered too large to be acceptable.

4.2 Candidate Crosslinking Agents

Of all the candidate crosslinking agents evaluated when the study was broadened, chlorosulfonic acid produced the greatest improvement in the dimensional stability of the fabric at the least expense to the desired fabric physical properties. This treatment appears to be completely resistant to laundering or washing in alkaline solutions. Table XIV summarizes the thermal shrinkage results of fabrics treated by the candidate systems.

	Treatment	The	ermal Shrinkage
1a.	POCl ₃ (vapor) + PBI	nitrogen	42
20.	rooig (vapor) · rzr	225 - 245°F	
b.	POCl ₃ (vapor) + PBI	0xygen 235 - 300° F	54
	v		
c.	POCl ₃ (vapor) + PBI	oxygen 225 - 235°F, H ₃ PO ₄	48
		223 - 233 F, n ₃ FO ₄	
2a.	POCl ₃ (reflux) + PBI	oxygen 218 - 228° F	50
	3		
b.	POCl ₃ (reflux) + PBI	oxygen 218 - 228°F, H ₃ PO ₄	23
c.	POCl ₃ (reflux) + PBI	oxygen	41
		218 - 228°F, POCl ₃ (liq)	
		heat 700°F 3 hrs. in air	→ 28
		3 hrs. in air	
3.	POCl ₃ (liquid) + PBI	70 - 230° F	45
		pyridine	
4.	POCl ₃ (liquid) + PBI	240°F	49
5.	H ₃ PO ₄ (85%) + PBI		23
	-3-4 (00 N) 1 L DI	219 - 300°F	20

Table XIV-1
Results of Treatments Utilizing Candidate Reagents (1)

Reagent	Conditions	Thermal Shrinkage**	Comments
10% Potassium Dichromate Sol'n	pH = 1, 212°F 5 min*	42%	Fabric burns in air due to residual dichromate
1% KMnO ₄ Sol'n	pH ≅ 12, 212° F 1 min*	42%	Fabric turns very dark - burns in air
0.7% Nitric Acid	212°F, 5 min*	56%	Fabric neutralized with NH ₄ OH prior to oven heating - no fabric color change
0.7% Nitric Acid	212°F, 5 min*	22%	Fabric turned deep purple color in oven
30%, Hydrogen Peroxide	400°F, 15 min	64%	
5% Chlorosulfonic Acid in Phosphorus Oxychloride	230°F, 5 min followed by oven treatment at 360° for 10 min	15% ° C	Sl. darkening of fabric
$10\%~\mathrm{H_2SO_4}$	400° F, 5 min	45%	Fabric shrinks
250 cc Phosphorus Oxychloride 10 g Terephthalic Acid 25 cc POCl ₃ 10 g Aluminum Chloride	230° F, 10 min	55%	
Same as above excep Maleic Anhydride substituted for Terephthalic Acid	pt 230° F, 10 min 220° C, 10 min	48%	
Het Anhydride substituted for Maleic Anhydride	230° F, 10 min 220° C, 10 min	31%	
100% Lupersol 130	200° C, 10 min	65%	

Table XIV-2

Reagent	Conditions	Thermal Shrinkage	Comments
Radiation, 12 Mrads, 3 MeV electrons	500° F	60%	
Phosphorus Oxychloride	230° F, O ₂ gas	48%	
Phosphoric Acid	260° F, 30 min	23%	32% shrinkage occurred on treatment
Hydroquinone	200° C, 10 min	65%	
Silane Y-4310	200° C, 15 min	60%	
Silar.e A-1100	200° C, 15 min	60%	
37% Formaldehyde Solution	boiled, 5 min 210°C, 10 min	61%	
2.1% Sodium Hypo- chlorite solution - 0.4% Sodium Hydroxide	boiled, 5 min 210° C, 5 min	37%	

⁽¹⁾ Data selected from Tables III to VII.

^{*}Following immersions in reagent solution, fabric was heated in oven to 390-400°F for about 15 min.

^{**}Determined by exposing 2" x 2" fabric swatch between metal gauze restraints to

^{~ 1300°}F for 1 min. Shrinkage quoted as initial area - final area x 100%.

Except for nitric acid, the oxidizing agents were largely ineffectual. Of particular note in the case of dichromate and permanganate solutions is the tenacious pick-up of these agents by the fiber. Even after extensive washing with soaps and reducing agents (oxalic acid, stannous chloride - hydrochloric acid, sodium thiosulfate), a substantial percentage of the dichromate and permanganate remained in the fiber. Since benzimidazoles are known to form a wide variety of complexes with metals (Ref. 4), it is conceivable that the chromium and manganese are chelated by PBI. If these metals are in fact chelated, they may be used as mordants to permit the uniform dyeing of the fiber - which is a prime requisite for the development of commercial interest in the fiber.

Treatment of PBI with nitric acid and heat produced a purplish tinge to the fabric. Concomitant with the purpling was a reduction in the thermal shrinkage to 22%. The coloration and thermal stabilization could be prevented by neutralizing the acid treatment with ammonium hydroxide solution before heating the fabric in the oven. It was found that fabric that had been turned purple could not be dyed with the dyes normally used with PBI. On the other hand, fabric, first dyed, could not be turned purple or stabilized through treatment with nitric acid and heat. A hypothesis consistent with these observations is that both the acid and the dye react with the same site on PBI - probably the amido nitrogen.

Shapeter hand towns a transaction to the

4.3 Chlorosulfonic Acid Treatment

In the batch-chlorosulfonic acid treatment, PBI fabric*, dried at 300° F for ten minutes, is immersed in a solution of 5% by volume chlorosulfonic acid (Eastman) in phosphorus oxychloride (Matheson Coleman and Bell) for 5 minutes. The fabric is removed and padded with absorbent towels to approximately a 50% wet weight addition. The fabric is then placed in a 340° C muffle furnace in 30 minutes. Following the heat treatment, the fabric is neutralized with a 10% armunium hydroxide solution, rinsed, and dried.

An optimization of the chlorosulfonic acid process was undertaken to determine the reaction conditions that would give maximum dimensional stability to a minimum change in PBI fabric flammability, color, and tensile strength.

^{*}Fabric dry cleaned and then washed with 1% neutral chip soap, 1% sodium silicate solution in commercial laundering equipment.

Table XV which gives data selected from Tables X and XI shows the effect of the temperature of the chlorosulfonic solution on the thermal shrinkage of 2/1 twill fabric.

As the solution temperature is increased, so is the permanent weight addition to the fabric. In addition, the oven temperature has a decided effect on the weight gain. Figure 7 shows the correlation between the per cent shrinkage (in the thermal shrinkage test) and the per cent weight gain. Of particular note is the effect of oven temperature. In fact, a trade-off between oven temperature and time was found. For example, identical thermal shrinkage results are obtained for fabric heated at

400°C for 2 min 360°C for 10 min 340°C for 25 min

This implies that the reaction rate increases 1.5 times for every 10°C increase in temperature.

The graphic presentation of Figure 8 shows how different conditions of time and temperature of treatment may be selected to obtain equivalent levels of stabilization of PBI through treatment with chlorosulfonic acid. Of course, stabilization is not the only result of exposure to the treatment. Other changes resulting in darkening and reduction in physical strength occur at the same time. Indeed, changes of this kind may occur as a result of treatments which do not yield optimum stabilization. This combination of behavior may be represented as shown in Figure 9, where the influence of time and temperature on the coexisting properties of stabilization and physical attributes are shown. Over-all optimization of treatment requires selection of a desired field within such a chart.

A theory of reaction consistent with the above information is that two reactions occur. The first takes place on immersion of the fabric in the chlorosulfonic acid solution.

If the fabric at this stage is noutralized with 10% $\rm NH_4OH$ solution, the fabric shows no weight gain.

Table XV

Effect of Temperature of 5% Chlorosulfonic Acid in Phosphorus Oxychloride on Fabric Thermal Shrinkage

	I HODPHOI	is oxyomorius o				
Run No.	Solution Temperature C	Time of Immersion (Min.)	Temp	en /Time Mins.)	Percent Weight Gain	Thermal Shrinkage (1)
10 11 12 13	room temp. room temp. room temp. 65	5 5 5 5	320 340 340 320	20 20 80 80	2.04 3.10 2.40 6.95	57 54 52 32
14 15 16	65 65 110	5 5 5	340 340 320	20 80 20	6.40 10.0 11.50	27.7 19.8 30.0 23.3
17 25 26 27 28	110 110 110 110 110	5 5 5 5	340 340 340 340 340	20 20 20 20 20	10.50 26.2 11.0 8.6 8.4	10.5 14.7 19.0 19.0
40	110	<i>.</i>	540		···	

NOTES:

(1) See Appendix A for Test Procedure.

عوقت معماله وموافق مايكومة وميدي يعطونك فالمسارة مدارة طومتناه فالأفاع فالأمام فالمبادة ماليوهو والموافقة الإولامة

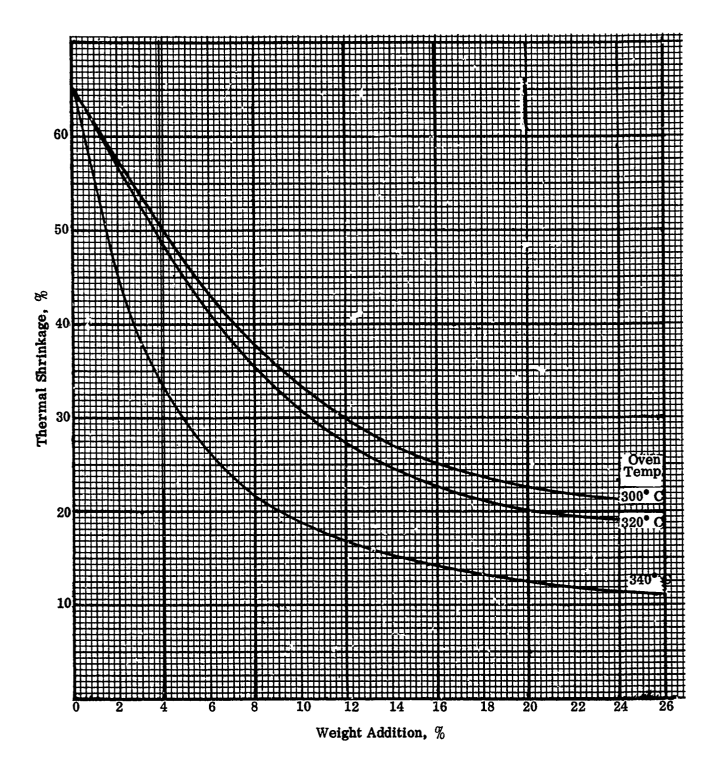
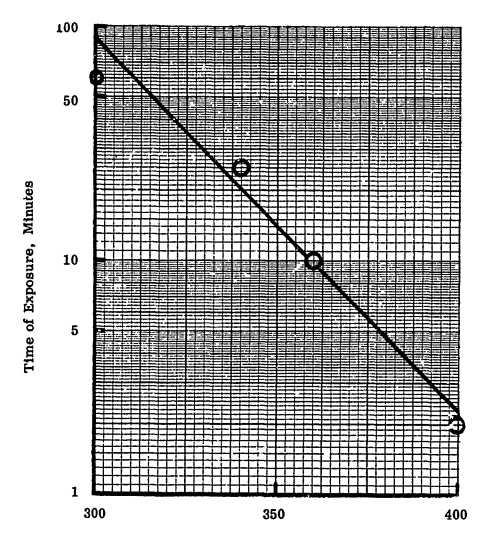


FIGURE 7

Fabric Thermal Shrinkage Versus Weight Addition



Oven Temperature, ° C

Figure 8

Conditions of Time and Temperature of Treatment

Producing Equivalent Stabilization of PBI with

Chlorosulfonic Acid

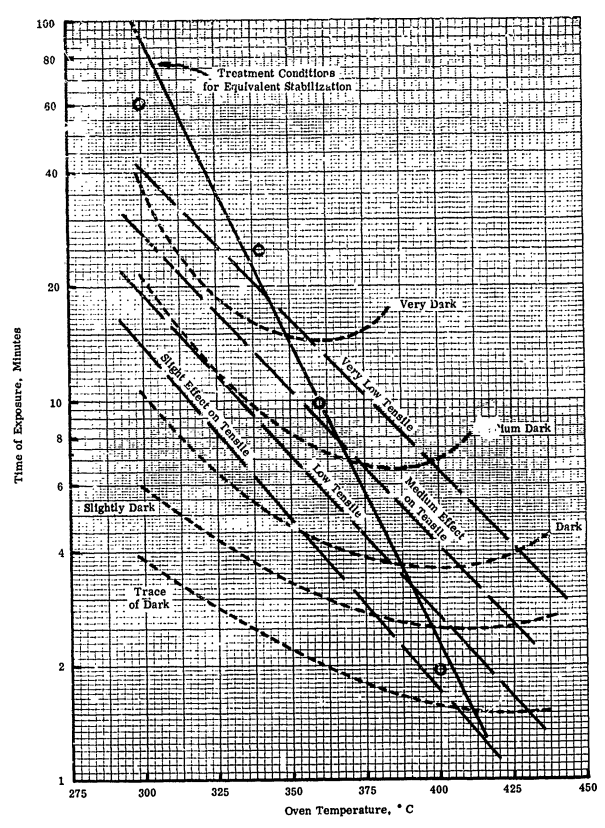


Figure 9

Representation of Effects on Physical Properties in Stabilization Treatment of PBI

If, however, the material is heated, the following rearrangement may occur:

A sulfonate group on the ring is not susceptible to hydrolysis with ammonium hydroxide — thus, a permanent weight gain, durable to laundering, is obtained after the heat treatment.

Lending credence to the proposed theory is the fact that nitrogen-substituted benzimidazoles are known to rearrange under the influence of heat (Ref. 5). For example,

Also well documented is the sulfonation of benzimidazoles with sulfuric acid and chlorosulfonic acid (Ref. 6).

In the patent literature these sulfonations generally involve dissolving the benzimidazoles in the sulfonating reagents. Since both sulfuric and chlorosulfonic acids are excellent solvents for the polymer, direct treatment with these reagents causes the fabric to be reduced to a jellied mass. To minimize the dissolution, solutions of chlorosulfonic acid in various non-solvents for the fabric were evaluated. Phosphorus oxychloride proved to be the best because it was non-protonating, completely miscible with chlorosulfonic acid, and effective in rapidly penetrating the PBI.

The cross-linking of the fabric appears to occur simultaneously with the rearrangement of the sulfonate group. Cross-linking through sulfonate groups probably occurs as shown below.

4.4 Seven Yards of Fabric Treated for Evaluation by Technical Monitor

As shown by the test results in Table XII, the seven yards of fabric treated by the chlorosulfonic acid in POCl₃ meets all the goals for Phase I. It has tensile and elongation properties equal or better than the control, essentially the same flammarbility, and the treatment shrank the fabric only 14%. The color of the fabric was affected only very slightly, the fabric is still dyeable and the hand and moisture regain properties are excellent. The fabric has 27% thermal shrinkage by our very severe thermal shrinkage test, however, our test produced about three times higher shrinkages than the open pit flame test. Consequently, shrinkages of the order of 8-10% are anticipated in the pit test. In fact, another lot of PBI fabric treated in this way under contract F33615-71-C-0342 and fabricated into flight suits performed outstandingly in pit test exposures at Maynard, Mass., August, 1971.

As shown in Tables X and XI it is possible, using this chlorosulfonic acid treatment, to produce fabric having much better resistance to thermal shrinkage than 27%, but problems are encountered with fabric darkening and stiffening. It is probable, however, that with commercial equipment and careful choice of reaction conditions it will be possible to produce light colored fabric with a good hand having less than 20% thermal shrinkage even by our test.

4.5 Tow Treatment Facility

The tow processing line was constructed using the fabric treatment facility as a model; however, provisions were made on the tow line so than precise control could be maintained of processing times, temperatures, and the gas environment to which the tow was exposed.

Initial trial runs on the equipment demonstrated the effectiveness and uniformity of the treatment. Twisted PBI tow, 30 plies of 550 fil fiber with about 1 1/2 twists per inch, was run through the continuous process at three speeds, 1.18 in/sec, 1.27 in/sec, and 1.88 in/sec. An oven temperature of 475° C and a nitrogen gas purge was employed. Thermal shrinkage results were obtained on tow sections by placing individual fibers in glass tubes, and then heating the tubes until the fiber became a blackened char (about 400° C for 1 hour).

,这么一样是是是我们的是是我们的是是我们的,我们是是我们的,我们是是我们的,我们是是我们的的,我们是我们的的,我们是我们的,我们们的是我们的,我们们的是是我们的,我

Table XVI summarizes the thermal shrinkage results, obtained at two processing speeds, of fibers selected from the cross-section of the tow.

Table XVI. Thermal Shrinkage of 30 Ply Tow

Processing Speed	Residence Time in Oven	Oven Temp.	Fiber No.	Thermal Shrinkage
1.18 in/sec	35.7 sec	475° C	1	22.1%
Ī	1	1	2	23 %
j	!	}	3	23%
1	•		4	22.6%
l	l l		5	22.9%
			6	23.4%
[7	24.8%
			8	30.6%
1			9	25.3%
1			10	19.8%
			11	19.7%
•			12	19.7%
lacksquare	$lack \psi$	$lack {f V}$	13	17%
	-	64		

Processing Speed	Residence Time in Oven	Oven Temp.	Fiber No.	Thermal Shrinkage
1.88 in/sec	22. 3 sec	↓ 475° C	Average Control A Control B	21. 8% 62. 2% 58. 3% 28. 6%
			2 3 4 Average Control A Control B	27. 7% 23. 5% 23. 6% 25. 9% 52% 54. 5%

The treated fiber produced in these first trials had a chocolate brown appearance: to minimize fiber discoloration, the oven temperature was lowered. Various temperature settings and oven residence times were evaluated to produce low thermal shrinkage fiber with minimum discoloration. During these trials it was found that a nitrogen purge did not appreciably prevent discoloration, and the purge was eliminated as a processing step. The process condition that were finally established are given in Table. XVII.

Table XVII
Final Processing Conditions Established for Tow Line

Fiber:	16,500 filament tow; twisted \sim 1.5 $\frac{\text{twists}}{\text{inch}}$
Line Speed:	10 ft/min
Chlorosulfonic Acid Make-up:	5% ClO ₃ SH (Eastman): 95% POCl ₃ (Ventron)
Residence Time in Acid Saturator:	~2 sec.
Tube Furnace Temperature:	430° C
Nitrogen Purge in Furnace:	No
Residence Time in Furnace:	21 sec.
Ammonium Hydroxide Concentration:	~58% NH ₄ OH

Ammonium Hydroxide Concentration: ~58% NH₄OH

Residence Time in Ammonium Hydroxide: ~9 sec.

Drying Oven Temperature:* room temp.

*The final dry step was accomplished by placing the fiber (on a bobbin) in a 300° F oven for 12 hours.

Fibers selected from the cross-section of the treated tow showed the thermal shrinkages and tensile strengths listed in Table XVIII.

Table XVIII

Thermal Shrinkage of Individual Fibers

Control	60.2%
Fiber A	18.6
Fiber B	23.3
Fiber C	21.4
Fiber D	22.0
Average for treated fibers	21.3%

Tensile Strength*

Control 1	52*
Control 2	51
Control 3	59
Fiber D	54
Fiber E	51
Fiber F	51

^{*}arbitrary units

The average shrinkage of 21.3%, the light color, and the maintenance of the tensile strength of the fiber : cated that the process had been developed to the point where a large scale production run could be made with confidence.

One and one-half pounds of tow were processed under these conditions and submitted to the Air Force Materials Laboratory for approval. Approval of the tow and to proceed with the processing of 30 lbs of PBI was received on July 16, 1971.

4.6 Production of Treated Tow

Five-thousand filament tow (1.5 denier/filament) was plied to 15000 filament tow having approximately 1-1/2 turns per inch. This tow was unreeled from 3 inch diameter bobbins into the continuous processing line. The saturator was filled with 5% by weight chlorosulfonic acid in phosphorus oxychloride. At the selected line speed the tow remained immersed in the 6 inch saturator for 3 seconds. The furnace following the saturator was set at 430° C. The period of immersion in the ten-inch neutralization tank was about 5 seconds. The tow was arranged to make seven double passes through the water rinse tank, giving a residence time of about 2 minutes. The drying oven was set at 420° C. The initial bobbin was processed with the above arrangement, and second bobbin was processed similarly, with the exception that the drying oven temperature was raised to 460° C.

The color of the fiber produced on the first and second bobbins was quite light, and it was concluded that a somewhat higher oven temperature could be tolerated. The oven temperature accordingly was raised to 448° C, and 4 additional bobbins were produced with this single change in processing conditions. The first two bobbins were designated Lot 1 and the last four bobbins were designated Lot 2.

4.7 Conversion of Tow into Fabric

The PBI tow, processed as described in 4.6, was supplied as two lots wound on six bobbins to Celanese Research Company to be crimped and cut to 2-inch length. The material could be handled satisfactorily in the crimping and cutting operations, but it was observed that the fiber affected the metal parts of the processing equipment. Following the processing it was found that aluminum parts were pitted and steel parts were heavily rusted. This action suggests that some of the fiber contained a residue of alkali while another part remained acid through incomplete neutralization. Support of this explanation is furnished by the observations of those handling the fiber at Celanese that the smaller lot, which had been processed at 430° C had a slight odor of ammonia. The larger lot, which had been processed at 448° C was characterized as smelling of "nitrates", i.e., gave an impression of acidity. It is evident that the

neutralization and water-washing steps in the continuous processing facility varied in effectiveness. During the processing, pH measurements of the damp tow leaving the rinse tank were made with indicator paper, but indications of neutrality obtained in this way obviously were not totally adequate.

The crimped and cut tow was sent to Textile Research Services, Inc., Raleigh, North Carolina, for weaving into cloth. In preparation for carding the fiber was sprayed with Nopco 2152P antistat. Upon stempting to card and draw the fiber it was found that the card loaded up with fiber after was found to build up on metal parts continuation. A relatively hard, powdery substance was found to build up on metal parts as it was attempted to process the fiber. This behavior is believed to be a further consequence of the presence of residual alkali or acid on the fiber as noted at Celanese Research Company. It is probable that the antistat material reacted with the acid and alkali to produce a coating on the fibers having adhesive qualities. This adhesiveness probably caused the card loading, and the material observed to build up on metal parts may well have been the reaction product of the antistat.

In order to overcome the behavior just noted the fiber was scoured. The scouring procedure was carried out with Igepon P in water at 190° F. The scouring cycle consisted of 15 minutes washing, two rinses of twenty minute duration, and 5 minutes extraction. After drying and conditioning Nopco 2152P antistat was again sprayed on the fabric. Following this treatment the fiber processed normally. The procedures followed in converting the fiber to cloth at Textile Research Services, Inc., are summarized in Table XIX.

Table XIX
Spinning and Weaving Procedure Applied to Crimped and Cut Tow

Step	Lot 1	Lot 2
1. Card, draw, and rove to spin and ply	Yarn no. 37/2 Twist 3.85 T.M. 'Z' x 14.5 T.P.I. 'S'	Yarn no. 37/2 Twist 3.85 T.M. 'Z' x 14.5 T.P.I. 'S'
2. Number spool	140 spools 380 yards/spool	140 spools 700 yards/spool
3. Make 2 warps via the silk system	140 ends/band 20 bands 3 turns/band	140 ends/band 20 bands 6 turns/band
4. Draw and reed	8 body harness (skip draw) 4 selvage harness 80 total selvage ends (40/side) 26.5/2 Reed 52" Reed width	Same
5. Weave	2 x 2 L. H. Twill 54 x 52 Off Loom Const. 4.4 oz/yd ² Fitting same as warps	Same

4.8 Performance of Fabric

The fabric produced from the tow processed as described in 4.6 had excellent color and hand, appearing entirely like other samples of fabric examined previously. Measurements were made of the tensile strength, limiting oxygen index and thermal shrinkage of samples taken from the two lots of cl. th. These data are presented in Table XX. It will be seen that the two lots differed importantly in their response to thermal exposure.

Table XX

Evaluation of Fabric Woven from Treated Tow

Property	Lot 1	Lot 2	Control
Tensile Strength, lb/inch width	74	73	94-96
LOI	0.40	0.50	0.41
Shrinkage, %	57	32	61

Lot 1 differed very little from an untreated control, whereas Lot 2 showed much improved shrinkage and a significant increase in oxygen index. The shrinkage obtained with Lot 2 was like that obtained earlier in treating already formed fabric. The oxygen index of the Lot 2 fabric prepared from tow was found to be higher than that of fabric treated as such.

Since the essential difference in preparing lots 1 and 2 was the oven temperature, no other reason for the differences in behavior can be advanced. The discussion of PBI behavior has made it clear that the thermal exposure following chemical treatment is important in determining response, but the levels selected in tow processing were within the range found useful. The temperature level of 436 selected initially and utilized in processing Lot 1 was selected on the basis of preliminary work (using single fibers for evaluation) as the lowest effective temperature and therefore a level at which fiber discoloration would be least. Upon observing that the fiber produced as Lot 1 was very little affected in color the decision was made to raise oven temperature 18° C, as reported. The color of the fiber so processed was not affected importantly and it is evident that this change in oven temperature was of profound importance.

It is apparent that the conditions of treatment chosen in processing the tow were too conservative. Referring to Figure 9, it may be appreciated that other conditions, specifically those involving higher oven temperatures, at the fixed residence time chosen, would have given consistently adequate stabilization at the cost of some additional darkening of the fiber. The limited experience in working with tow contributed to the conservative choice of conditions; in subsequent treatment of tow it will be possible to raise the level of stabilization substantially. Further work on tow processing and preparation of fabric from treated tow is recommended in order to achieve the desired level of stabilization in conjunction with acceptable fabric properties.

Section 5

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CONCLUSIONS

The shrinkage which occurs when polybenzimidazole (PBI) fabrics are exposed to the high temperatures encountered in flames can be reduced substantially through suitable chemical treatment of the fibers or of finished cloth. A treatment involving immersion in a solution of 5% by weight chlorosulfonic acid in phosphorus oxychloride followed by thermal treatment at 300°-450° C was found to reduce the area shrinkage of treated fabric by 50% without affecting the hand and drape, color, or physical properties of the fabric to an important degree. The flammability of the treated fabric is essentially unaffected. The treatment is durable, permitting the fabric to be laundered normally.

The process for thermally stabilizing PBI is suited to continuous processing of fiber in the form of tow or fabric. A lot of PBI fabric, treded as described, and fabricated into flying suits performed exceptionally well on exposure in flame pit tests, indicating that the treatment provides additional properties of high value to PBI fabric. PBI tow was processed on a continuous basis with the treatment, the tow being subsequently woven into cloth. The thermal stabilization process developed has been shown to provide the necessary fabric improvement while enabling fabric color and physical properties to be acceptably maintained.

The success achieved in developing a practical stabilization process for PBI supports the recommendation that the parameters involved in satisfactory treatment of PBI tow be subjected to additional investigation. Such further work on tow treatment will enable full-range utilization of this important improvement in manufacturing PBI fabric for all Air Force applications.

Appendix A

TEST FOR FABRIC SHRINKAGE DURING FLAME IMPINGEMENT

Equipment

Meker Burner-Cenco Cat #11043

Two 2 1/4" square pieces of 40 mesh stainless steel screen 300 gram weight-made by melting a 60/30 mixture of lead and 50/50 solder in a 600 ml stainless steel Beaker, Fisher Cat #2-583

4 x 4 Corning Ware Ceramic Dish (P-41 Petite Pan)

2 x 2 Steel die which cuts a sample 4.0 in² in area

Procedure

- 1. Mount the ceramic dish on a ring stand so that it is 3/8" above the Meker burner and adjust gas to give a temperature of 800° F \pm 10° F as measured by a copper constantan thermocouple touching the center of the ceramic dish.
- 2. Warm the weight to $450^{\circ} F \pm 10^{\circ} F$ (about the melting point of the lead solder mixture).
- 3. Place a $2'' \times 2''$ cloth sample $(A_0 = 4.0 \text{ in}^2)$ die cut from the sample to be tested between the pieces of wire screen and lay it in the hot ceramic dish and cover with the weight. Remove after 1 min. + 5 secs.
- 4. After the cloth is cool, measure area A_1 by tracing an outline on a piece of 10 x 10 per 1/2" rectangular coordinate graph paper and counting the number of squares. Calculate % shrinkage.

% shrinkage =
$$\frac{A_0 - A_1 \times 100}{A_0}$$

- 5. Repeat above and report the average shrinkage of the two samples.
- 6. The data below obtained on untreated fabric shows the precision to be expected from the test.

Fabric	Original Area	Area After Heating	% Change (based on original area)
Continuous Filament	$4 in^2$	$1.50 in^2$	-62.5%
Continuous Filament	4 in ²	1.44	-64.0%
2.85 oz/yd^2	4 in^2	1,56	-61.0%
6.17 oz/yd^2	4 in ²	1.62	-59.5%

Appendix B

FIBER TENSILE TEST METHOD

Equipment

Tensile Testing Machine suited for measurements of 0-15 lb * 12 inches/minute.

Sample jaws suited to clamp one end of a loop formed from 6-7 inch fiber and suspend other end of loop.

Procedure

- 1. Form a loop of a length of fiber. holding the free ends between thumb and forefinger.
- 2. Clamp end of fiber loop formed at free ends in tester jaw.
- 3. Hook blind end of loop over suspending rod on other jaw of tester.
- 4. Check that loop is secure with jaw separation 2-3/4'' 3-1/4''.
- 5. Set tester speed at 12 inches/minute.
- 6. Record tensile at break.
- 7. Repeat, determining results for 5 fibers.
- 8. Calculate and report the average tensile at break for the 5 samples tested.

Appendix C

PROCEDURE FOR DETERMINING SHRINKAGE OF POLYBENZIMIDAZOLE FIBER

Standard Testing Procedure, Number 25 - Dec. 1970

Equipment

3 glass thes, 1 ft. long, 1.2 mm ID. Circulating Air Oven at 500° C.

Procedure

- 1. Thread fiber through the glass tube using slight suction, cut the fiber at both ends of the tube; measure its length to the nearest 1/32".
- 2. Repeat step 1 with two more fibers in two more tubes.
- 3. Place the tubes in an oven set at 500° C \pm 5° C as measured with a calibrated or thermocouple for 10 mins.
- 4. Remove the tube from the oven and remeasure fiber lengths to the nearest 1/32".
- 5. Determine linear shrinkage based on 1' initial length and length as measured in tube after shrinkage.

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