



**Annual Meeting of
Shareholders
September 14, 2010**

CONFIDENTIAL

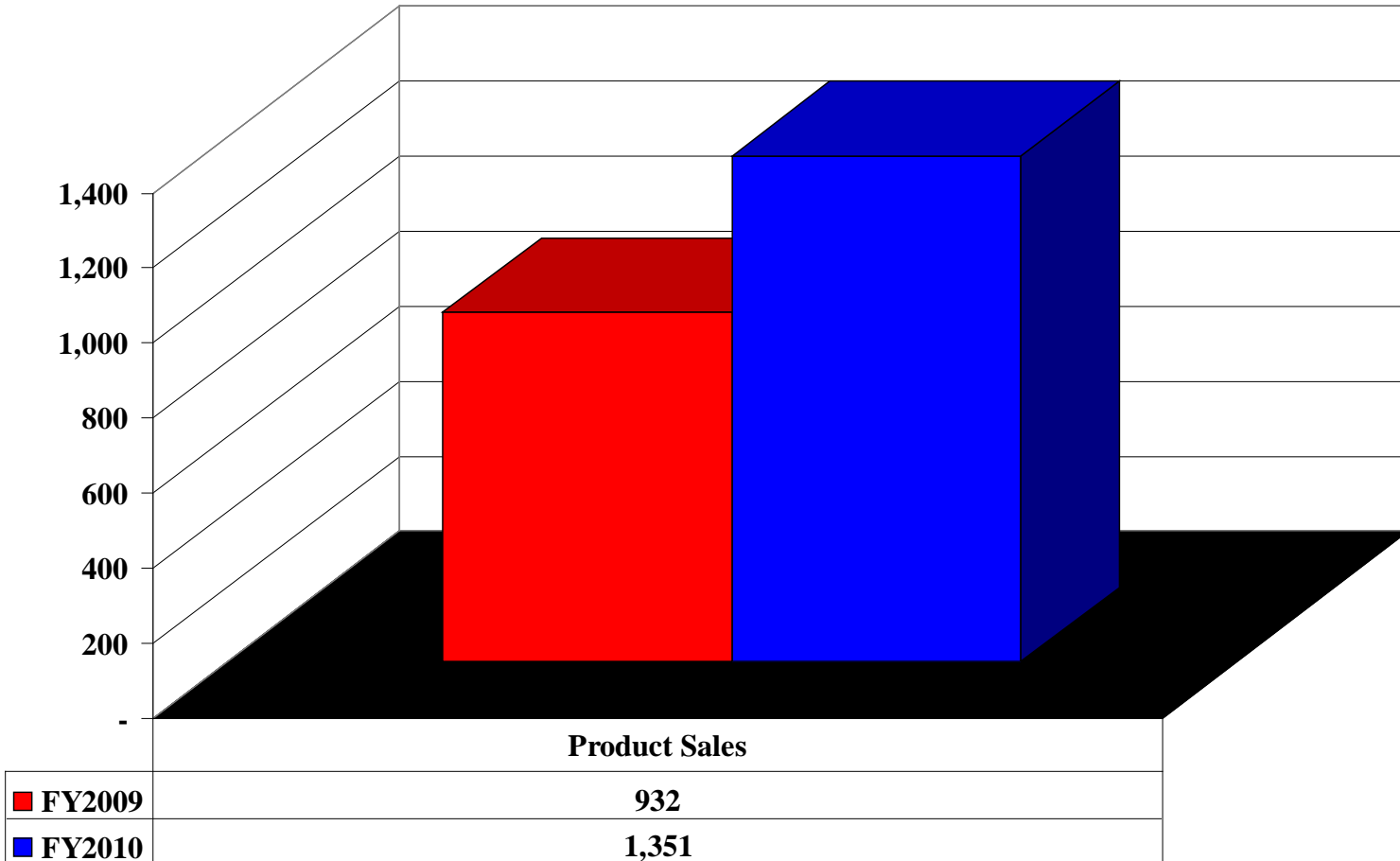
Forward-Looking Statement

This slide presentation may contain projections or other forward-looking statements regarding future events or future financial performance. These statements are only predictions and reflect the current beliefs and expectations of AdvanSource Biomaterials Corporation. Actual events or results may differ materially from those contained in the projections or forward-looking statements. Factors that could cause actual results to differ materially from forward-looking statements contained herein are detailed in documents AdvanSource Biomaterials files from time to time with the Securities and Exchange Commission, including recent filings on Form 10-K and 10-Q. Forward-looking statements in this presentation are made pursuant to the safe harbor provisions contained in the Private Securities Litigation Reform Act of 1995.

Financial

- No Debt
 - Positive Gross Profit on Product Sales
 - Restructured Existing License, Royalty and Development
 - 15% Reduction in Operating Expenses
 - \$2.1MM of Cash as of September 13, 2010
 - Growth in Product Sales
-

Financial



Significant Events

- Medos Settlement
 - Close CDT Sale Escrow
 - CardioPass
 - Patents:
 - Awarded Anti-Microbial Polymer Patent
 - Filing of ChronoSil Patent Application
 - Submission of NYSE Amex Plan to Regain Compliance with Listing Standards
-

Commercial Activities

- Record # of Negotiated Agreements
 - Significant Increase in Customer Base
 - Expanded Sales and CS Presence
 - Establishment of Alliance Partnerships
 - Value-Added Service/s Market Presence
 - Aggressive International Expansion
 - Thought Leadership Recognition
-

Data Sheet Progression

CT BIOMATERIALS



Technical Fact Sheet

***A Premium Grade of Long Term Implantable Thermoplastic Polyurethane.**

Introduction

ChronoFlex®AL is a family of premium polycarbonate aliphatic biodegradable thermoplastic polyurethane elastomers developed by CT Biomaterials, Div. of CardioTech International Inc. ChronoFlex®AL medical-grade elastomers are manufactured under strict GMP conditions. ChronoFlex®AL has been developed to overcome the in vivo formation of stress-induced microfissures. This makes ChronoFlex®AL an excellent material choice for medical device applications requiring long term durability.

Product Advantages

ChronoFlex®AL elastomers are custom synthesized from reactive monomers. CT Biomaterials, Div. of CardioTech International tailors polymers to specific customer applications. ChronoFlex®AL polymers can be compounded with various radiopacifiers at the prepolymerization stage to ensure maximum dispersion of additives. This enhanced dispersion results in surfaces that are considered to be smoother and glossier than those provided in conventional compounding techniques. ChronoFlex®AL

polymers are offered in hardnesses ranging from a soft 80 Shore A to a very hard 75 Shore D.

Physical Properties

Shore Hardness	80A	55D	65D
Appearance	Clear	Clear	Clear
Modulus at 100%	650	2900	3200
Modulus at 300%	660	----	----
Tensile Strength (psi)	5500	8400	9000
Elongation (%)	585	325	300

Drying Recommendations

ChronoFlex®AL pellets should be dried to a moisture content equal to or less than 0.01% by weight prior to melt processing. Most extrusion problems can be traced to improperly dried pellets. We recommend a desiccant-type hopper drier capable of reaching and maintaining a dew point of minus 40°F. Typically, the pellets should be dried for at least 3 - 4 hours with a drier inlet temperature of 160° - 185°F. If the pellets can be pre-dried for extended periods, such as overnight, an inlet temperature of 110 - 130°F is generally

sufficient.

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AdvanSource
biomaterials

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Fact Sheet: ChronoFlex® AL

ChronoFlex AL is a family of biodegradable thermoplastic polyurethane elastomers developed by us. These medical-grade elastomers have been specifically synthesized to overcome the in vivo formation of stress-induced microfissures. Surface microfissuring is a major weakness of present polyurethane elastomers which may lead to catastrophic failures in long-term implanted devices.

Product Advantages

In the early 1980's it became clear that ether-based polyurethane elastomers were susceptible to biologically induced environmental stress cracking. Microcracks formed on the surface of polyurethanes as a result of exposure to the biological environment, particularly where the polymer was subjected to mechanical stress. ChronoFlex AL is at the forefront of a new generation of aliphatic, ether-free, thermoplastic polymers which do not undergo biologically induced environmental stress cracking. ChronoFlex AL derives inherent biodegradability from its polycarbonate-based structure.

Physical Properties

	80A	55D	65D
Modulus at 50% (psi)	500	----	----
Modulus at 100%	650	2900	3200
Modulus at 300%	600	----	----
Tensile Strength (psi)	5300	8250	9000
Elongation (%)	585	325	425

Handling Extruded Tubing

Following extrusion, ChronoFlex AL tubing will remain sticky for about 10-15 minutes. The tubing must be protected during this time frame. Afterwards, the tubing surface will become tack-free. Surface tack will diminish as surface crystallization of the hard segments takes place.

Drying Recommendations

ChronoFlex AL pellets should be dried to a moisture content equal to or less than 0.01% by weight prior to melt processing. Most extrusion problems can be traced to improperly dried pellets.

ChronoFlex® ChronoThane™ HydroMed™ HydroThane™ PolyBlend™

ALIPHATIC POLYCARBONATE-BASED URETHANES

ChronoFlex AL™

Exceptionally strong, safe and dependable.

Description

ChronoFlex AL is a family of biodegradable aliphatic polycarbonate-based thermoplastic polyurethanes designed to overcome surface degradation such as stress-induced microfissures.

With a long history of reliable performance in both long and short term devices, this medical grade polymer has the versatility to be used across a broad range of application areas ranging from oncology and neurology to cardiovascular disease management.

These ether-free polyurethane elastomers are biostable and display a low modulus of elasticity, excellent solvent resistance and limited softening in-vivo.

These products are adaptable to most standard manufacturing processes and are available in hardnesses ranging from 75 Shore A to 75 Shore D.



CHRONOFLEX AL IN PELLET FORM

The ASB Advantage

AdvanSource Biomaterials synthesizes and manufactures medical grade materials offering the ability to tailor physical and mechanical characteristics to support and enhance your end product design.

This characteristic's, critical to the design and development of medical devices, can incorporate a wide range of physical and chemical properties. While maintaining core characteristics such as biodegradability and biocompatibility, specialized characteristics such as the addition of radiopacifier or colorant agents and antimicrobial properties can be added in the formulation such to provide a homogenous material and limit secondary processing steps.

With an expanding range of secondary operations including custom solution development and prototype coating services, ASB's expert team of chemists, scientists, engineers and industry professionals assists in every stage of customers' projects, from concept initiation through full-scale manufacture.

An ASB product

- EUROPEAN COUNCIL DECISION - 1999/54/EC - COMPLIANT
- EXCELLENT SOLVENT RESISTANCE
- BIOCURABLE
- LIMITED IN-VIVO SOFTENING
- INHERENT MATERIAL STRENGTH
- BIOCOMPATIBLE
- RELIABLE PERFORMANCE IN LONG AND SHORT TERM IMPLANTABLE DEVICES
- LOW MODULUS OF ELASTICITY
- ESC RESISTANT

AdvanSource
Biomaterials

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Ask us about our animal-free origin products.

- COMPLIANT WITH EUROPEAN COUNCIL DECISION 1999/534/EC
- VERIFIED RAW MATERIAL SUPPLY
- PROACTIVE RESPONSE TO INDUSTRY NEEDS



AdvanSource Biomaterials upholds the highest risk assessment through rigorous processes and performance criteria while advocating efficient and effective control measures at every step of our manufacturing process.

ASB...proactively partnering to protect our clients... and yours.



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- COMPLIANT WITH EUROPEAN COUNCIL DECISION 1999/534/EC
- VERIFIED RAW MATERIAL SUPPLY
- PROACTIVE RESPONSE TO INDUSTRY NEEDS



AdvanSource Biomaterials mitigates the risk of infectious agent contamination through controlled sourcing and qualified evaluation of all our starting materials.

ASB...taking one more step to ensure the safety of medical devices.



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Enhanced Global Expansion

SHOW PREVIEW: BIOMEDevice

to ISO 9001 and the four US plants are certified to ISO 13485. With more than 30 years of experience developing flexible packaging for the medical device industry, the company is proficient in the industry's regulatory, quality and CGMP requirements.

Ancor Flexibles
Stand 022

Packaging

Bischof + Klein GmbH & Co. KG supplies flexible plastic packaging and technical films to medical device companies. Operating a number of

cleanroom production facilities, the firm offers low-germ and low-particle packaging materials. The company's CleanFlex product line conforms with the requirements of the pharmaceutical, medical device and related industries for flexible cleanroom packaging. The CleanFlex products are available as single-wound and tubular films, open-mouth and side gusseted mitred sealed bags, two- and three-ply bags and Tyvek bags with three side seals that are autoclavable. The company also offers extrusion, printing and converting services under cleanroom conditions.

Internal laboratory and testing facilities are available onsite. The company is certified to ISO 9001:2000 and complies with US FDA CGMP guidelines.
Bischof + Klein GmbH & Co. KG
Stand C19

Valve-testing equipment

CB Automation has developed a machine designed to ensure the correct assembly of drug-delivery valves. The machine also can determine the valves' dose by analysing air pressure. The unit's dose control system is composed of a pair of mechanical heads, pressure transducers and electronic cards for data acquisition. The machine creates a light pressure in the transducer chamber by compressing the head gasket. This preliminary operation is carried out to check for leakage in the valve's upper chamber. Afterwards, an upper cylinder opens the



valve and the resulting air input creates pressure proportional to dose volume. The pressure can be monitored on the PC that is included with the system. The control parameters (time on the x axis and pressure on the y axis) allow the identification of dose volume and the detection of valves that are not compliant.
CB Automation – Division of Bettinelli
Stand C13

Custom plastic components

Formed in 1989, APE Medical is a plastery company that specialises in the manufacture of plastic technical components for the pharmaceutical, medical and diagnostics industries. The manufacturer has more than 15 years of experience serving the pharmaceutical and medical industries. The firm offers a variety of services including moulding under Class 10,000 cleanroom conditions,

klassische Spitzeneinsatz zur Seitenanspritzung im 90°-Winkel zur Entformungsrichtung ist nun auch mit 60-, 45- und 30°-Winkelung verfügbar. In der Medizintechnik lässt sich so beispielsweise die oftmals bei der



seitlichen Anspritzung dünnwandiger, schlanker rohrförmiger Bauteile wie Pipetten auftretende einseitige Belastung des Formkerns durch den Schmelzedruck mit daraus resultierendem Kernversatz reduzieren – durch gewinkelte Spitzen kann der Anspritzpunkt näher am Kernlager positioniert werden.

Die Schmelzuführung erfolgt seitlich, der Anschnitt ist aber in Entformungsrichtung (0°) positioniert. Diese Technik wurde besonders für Bauteile entwickelt, bei denen eine Anspritzung auf waagerechten Flächen in direkter Nähe aufsteigender Konturen gefordert ist, beispielsweise auf Flanschen von Probengefäßen oder auf Halteplatten von Spritzen. Dies kann sowohl mit rechteckig gekröpften Wärmeleitspitzen, als

auch mit Nadelverschlussmechanik erfolgen. Bei beiden Verfahren kann der Anspritzpunkt bis zu 3 mm nah an der aufsteigenden Kontur platziert werden. Beim Nadelverschlussmechanik erfolgt die Nadelbetätigung synchron über eine Hubplattenmechanik. Eine Neuheit sind die kombinierten Nadeldichtungen und -führungen, die anschnittnah im gekühlten, einteiligen Formeinsatz eingeschraubt sind. Neben hoher Dichtigkeit des Systems können so eine kurze freie Nadellänge und minimale Nadelbelastung erreicht werden. Anstelle der beim offenen System eingesetzten Wärmeleitspitzen werden gebogelte

Wärmeleitelemente verwendet, die durch ihre spezielle Formgebung die Nadel direkt vor dem Anschnitt großflächig und berührungsfrei umschließen und eine gleichmäßige Wärmeverteilung im Anschnittbereich sicherstellen.
EWIKON Heißkanalsysteme GmbH & Co. KG
Frankenberg, Deutschland
Stand 459

Hochleistungskeramik für die Dentalmedizin

Zirkonoxidexperten aus der Schweiz haben Altbewährtes neu definiert und das Ziraldent-Implantatsystem in Zusammenarbeit mit der Abteilung

Creating Technology.



Enabling Success.

AdvanSource Biomaterials continues advancing material technology through product innovation to provide applicationally focused materials, targeted polymer synthesis, second source alternatives, and an expanding range of secondary operations.

In addition to our polycarbonate-based urethanes and ether-based elastomers, key advancements in our product portfolio include thermoplastic silicone polycarbonate urethanes, liquid polymers, extrudable & solution-based hydrophilic materials, homogeneous colorant technologies, and a full line of antimicrobial products.

In support of the engineering community we offer our commitment of a consultative partnership, on-time delivery, and ease of manufacturability through a broadened range of products spanning the durometer range from 5 Shore A to 75 Shore D.

AdvanSource Biomaterials – where partnership makes a material difference.

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Schaffung von Technologie.



Ermöglichen des Erfolgs.

AdvanSource Biomaterials führt fort, materielle Technologie durch Produktionnovation voranzubringen, um applicationally fokussierte Materialien, gerichtete Plastiksynthese, Zulieferalternativen und eine erweiterte Palette der Sekundärbetriebe zur Verfügung zu stellen.

Zusätzlich zusätzlich unseren Polycarbonat-gegründeten Uräthanen und zu Äther-gegründeten Elastomeren schließen Schlüsseltechnologien in unserer Produktmappe thermoplastische Silikonpolycarbonat-Äthane, flüssige Polymer-Plastiken, extrudierbares & mit ein Lösung-gegründete hydrophile Materialien, homogene Farbstofftechnologien und eine volle Linie der antibiotischen Produkte.

Zur Unterstützung der Technikgemeinschaft bieten wir unsere Verpflichtung einer breiten Teilhaberschaft, Einschaltzeit-Anlieferung an, und Mühelosigkeit des manufacturability durch eine erweiterte Produktpalette die Härtemesserskala von 5 Ufer A überspannend 75 Ufer D.

AdvanSource Biomaterials - wo Teilhaberschaft materielle unterschreibt.

AdvanSource
biomaterials

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Enhanced Global Expansion



开创技术。
推动成功。

AdvanSource Biomaterials是一家专业从事聚氨酯技术开发的公司, 可提供量身定制的目标聚合物合成和材料科学专业知识, 以优化医疗器械的设计和开发过程。

AdvanSource Biomaterials专注于开发材料科学领域的创新型先进技术, 全心全意应对医疗行业最为重要的挑战。

产品和服务: • 结构工程聚合物 • 抗菌聚合物 • 用于挤压和涂层的亲水性材料 • 液态聚合物 • 目标机械配方 • 均质着色剂技术 • 原型涂层能力 • 定制合解决方案

AdvanSource Biomaterials致力于为工程和医疗器械行业提供咨询合作关系、按时交货服务、批次间一致性、专业技术团队以及硬度范围为5 Shore A至75 Shore D的材料。

AdvanSource Biomaterials——合作让材料与众不同。



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Advertising

创造科技 | 迈向成功!

公司概况

AdvanSource Biomaterials ("ASB") 是一家通过 ISO 认证的材料科技公司, 致力于医用器材的研发、工艺放大和制造, 可依据需要定制物理和机械特性, 支持更美的产品设计需求。

通过对产业需求与趋势的综合分析, 以及对高分子化学制剂的不断钻研, ASB 开发的新型生物材料可用作结构工程高分子材料, 还可用作众多材料减薄的涂层。

ASB 的专有材料长期以来应用于微创型及大型型的介入医疗器械, 例如支架、人工心脏、导管和血管通路接口。

创造科技 | 迈向成功!

AdvanSource Biomaterials 将最初的设计概念完美地转化为最优化的器械。

创造科技, 引领医疗和制药行业多个领域的客户共同迈向创新。

迈向成功, 让精益制造产能开发并支持拥有先进技术的产品, 应对各种手术器械需求和器械改良。

企业实力:

产品创新

ASB 通过产品创新不断研发新材料, 提供了应用针对多种材料、目标高分子材料合成以及第二单体替代。

ASB 一直在不断地扩大产品组合, 包括聚酯醚醚基聚氨酯; 芳香族和脂肪族聚氨酯; 脂肪族和芳香族聚氨酯; 可塑性的以及基于常用的亲水材料; 涂层与粘接剂; 弹性高分子材料以及完整的医疗器械系列。ASB 还在继续开发所有新的材料科学专业知识, 提高加工能力, 致力于优化产品质量。

生产

ASB 在生产过程的每个阶段均严格执行《良好操作规范》, 公司通过了 ISO 9001:2000 以及 ISO 13485:2003 认证, 符合《医疗器械生产》(CFR) 的要求, 并可根据您一定的技术和资源需求, 确保您的产品始终如一。作为生物材料行业领军企业, 我们拥有先进的生产设施, 为产品开发的每个阶段都进行了有效谨慎的监控, 致力于为您的客户提供技术, 增加进一步加工处理以及上市的质量, 效率和交货速度。

合作材料开发

ASB 致力于与客户建立战略合作关系, 保证产品的按时交付。同时, ASB 的售前产品专家具备材料科学的化学及机械特性, 提供极具的协助性。公司还将根据您的特定需求制定产品, 具备量身定制的耐用性。ASB 产品所具有的特性: 离子材料中可加入特殊限制 (例如加入适当的着色剂和抗菌特性), 以提供符合材料并减少二次处理步骤。

此外, 还可将 X 射线造影剂融入配方, 进一步提高产品性能。X 射线造影剂的含量最高为 40%, 提供可对比的射线成像造影。

ASB 配备了二次加工立身的设备 (包括定制开发者和标准涂层服务), 积极促进团队合作。

ASB 优势

ASB 专家团队由科学家、化学家、开发工程师、质量控制以及技术人员组成, 累积了 150 多年的综合经验, 在整个开发阶段为您提供全程的指导, 以客户为中心的专业技术支持。

ASB 提供高质量的可重复性, 各批次产品的一致性, 专业的团队以及极具竞争力的材料成本 (从 50 磅 A 到 75 磅 D)。

AdvanSource Biomaterials 的产品只由指定、易于生产、符合严格的法规性要求, 能够克服难以加工 (ESPC), 且具有较好的耐用性, 易于植入。因此, ASB 始终为您提供生物持久性及用原材料领域的领先者。

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Company Overview

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聚氨酯醚基聚氨酯

ChronoFlex C (芳族) 和 ChronoFlex AL (脂肪族) 产品系列是设计用于生物相容性 (如血液相容性) 并可提供高弹性力 (ESPC) 的生物持久性聚氨酯醚基聚氨酯。两个产品系列均不含第二单体。此外, 两种产品均符合欧盟理事会进行 1999/53/EC 决议的规定。

在临床中, 这些聚氨酯醚基子材料在血液和天然液体环境中表现出优异的性能, 如生物持久性, 良好的生物相容性, 弹性以及出色的耐用性。它们具有良好的稳定性, 表现出优异的弹性、出色的耐用性, 并且植入后不会发生降解。硬度范围从 75 磅 A 到 75 磅 D。

ChronoFlex AR1 产品系列为聚氨酯醚基聚氨酯, 可完全在液体形式存在, 设计用于涂层、油墨以及涂层应用。此外, 可进行静置以形成水乳状液。这些材料对 X 射线造影剂具有持久性 (例如人工心脏瓣膜、人造血管) 的应用或高清晰度成像的用途。ChronoFlex AR1 是高度透明的涂层, 能够稳定长期的寿命。这种独特性使其适用于 W/O 和 O/W 乳液等应用。

ChronoFlex AR 和 ChronoFlex ARLT 的不含邻苯二甲酸, 而且都是非动物源性认证产品。根据产品需求, 可提供具有各种物理/化学形式的产品。

ChronoSR 是一种聚氨酯醚基聚氨酯。它保留了聚氨酯醚基聚氨酯的所有优点, 包括生物持久性、耐降解性 (ESPC)、抗降解性以及优异的化学稳定性与耐压缩性; 并增加了新的优点, 例如: 最佳的弹性、优异的耐用性以及高清晰度。这些优点得到 FDA 的认证。ChronoSR 不含邻苯二甲酸, 符合欧盟理事会进行 1999/53/EC 决议的规定。

ChronoSR 还保持了其其它产品的优点, 包括定向机械特性、生物相容性以及优异的机械强度 (从 75 磅 A 到 75 磅 D)。

聚氨酯基聚氨酯

ChronoThane P (芳族) 和 ChronoThane T (脂肪族) 产品系列是聚氨酯醚基聚氨酯。这些生物相容性材料具有多种特性, 例如: 最佳的弹性、优异的化学稳定性、生物持久性、优异的耐用性、ESPC、生产批次间不含第二单体。两个产品系列均不含邻苯二甲酸。ChronoThane T 符合欧盟理事会进行 1999/53/EC 决议的规定。

这些高分子材料易于生产, 并可长期稳定地确定其机械性能。两种产品, 应用范围包括导管、接口及连接器等。材料的硬度范围为 75 磅 A 到 75 磅 D。

亲水性材料

HydroThane 产品系列包括可挤压的亲水性聚氨酯醚基聚氨酯。根本的硬度为 75 磅 A。这些材料具有优异的弹性、优异的耐用性、良好的机械强度和优异的化学稳定性。这些材料具有出色的生物相容性。同时, 两种产品均符合欧盟理事会进行 1999/53/EC 决议的规定。

可挤压的亲水性产品系列在提供独特的特性, 适用于制造上述提到的应用。产品具有一般的亲水表面亲水性以及低摩擦系数, 从而避免了一次性重复使用。

HydroThane 即使在湿润状态下也具有弹性, 无弹性存储条件。目前的适用范围为 80 磅 A 和 80 磅 A。

产品中不含邻苯二甲酸, 符合欧盟理事会进行 1999/53/EC 决议的规定。

HydroMed D 系列聚氨酯醚基聚氨酯具有出色的弹性/粘弹性。可根据最终产品的需要定制成各种材料, 使之具有不同的亲水性和耐水性。适合用于公众接触 (包括导管、软管、介入器械以及泵) 的涂层。与常规水凝胶相比, 因此为需要长时间重复使用的器械提供了更好的选择。

产品中不含邻苯二甲酸, 符合欧盟理事会进行 1999/53/EC 决议的规定。这确保了具有卓越性能、优异稳定性、优异耐用性; 此外, 优异的机械强度和优异的化学稳定性使其具有优异的弹性。PolyBend 的硬度范围为 45 磅 A 至 80 磅 A。

ChronoPrene 生物相容性聚氨酯醚基聚氨酯具有优异的弹性、优异的耐用性、优异的机械强度和优异的化学稳定性。此外, 优异的机械强度和优异的化学稳定性使其具有优异的弹性。PolyBend 的硬度范围为 45 磅 A 至 80 磅 A。

ChronoPrene 是一种容易成型的高弹性材料, 具有优异的挤出、注射和挤压方法; 对于弹性特性的应用 (例如: 导管/连接器) 以及内窥镜 (例如: 导管) 等。这些材料具有优异的弹性、优异的耐用性以及优异的机械强度。ChronoPrene 展现出优异的亲水性和优异的弹性。这些材料在多种应用中可替代传统材料。

产品中不含邻苯二甲酸, 符合欧盟理事会进行 1999/53/EC 决议的规定。ChronoPrene 表现出优异的弹性、耐久性和优异的稳定性, 硬度范围为 5 磅 A 至 40 磅 A。

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Thank you for your time.
