



AseptiVent VF-y

Gamma Irradiatable PVDF Capsule Filters

for Sterile Filtration of Air/Gases in Biopharmaceuticals

Data Sheet

Biopharmaceutical manufacturing involves sterile filtration of air and gases for a multitude of critical processes such as air sparging, bioreactor venting, fermentor exhaust etc. The critical nature of biopharmaceutical processes and associated high costs require the highest degree of reliability for the filter device with regard to its retention efficiency, flow rates, service life and mechanical and thermal stability.

In order to do away with validation, energy and cleaning costs associated with reusable process assemblies and bioreactors, biopharma industry is moving towards single use disposable systems. Gamma sterilizable hydrophobic membrane filter devices offering high quality and reliability have become a necessity.

mdi gamma sterilizable *AseptiVent VF-*γ hydrophobic PVDF membrane capsule filters with a wide range of end connections and different sizes for linear scalability are specially designed for use with disposable single use assemblies for biopharmaceutical processes.

These filters are validated for microbial retention with liquid bacterial challenge test to ensure reliable performance under worst case conditions.

Applications

- > Sterile air sparging
- > Sterile venting
- > Fermentor exhaust

Key Features

- Absolute retention
- > 100% integrity tested
- High hydrophobicity
- > High air flow rates
- Low Bioburden, <1000 cfu/device</p>
- > Endotoxin level certified to be <0.25 EU/ml
- Widest range of end connections
- Products available for total scalability from seed reactors to process scale bioreactors/fermentors
- > Total traceability (unique serial number for each filter)
- > Individual certificate of quality for each device
- Sterilizable by Gamma irradiation or autoclaving

Quality Assurance

mdi's quality management system emphasizes on quality by design rather than by end product testing. Robust processes are developed for product manufacturing and are continuously monitored to ensure that the products meet their predetermined specifications and lot to lot reproducibility is ensured.

Certificate of Quality

Each AseptiVent VF- γ is accompanied by individual certificate of quality to ensure traceable documentation at user's end.

It certifies the product compliance to various regulatory as well as user requirements.

Validated for Microbial Retention

Even though AseptiVent $VF-\gamma$ is used for air/gas filtration, it is validated by liquid bacterial challenge test to subject the filter to most stringent conditions for higher degree of assurance.

Integrity test data have been correlated to actual microbial retention with Brevundimonas diminuta ATCC 19146 as per ASTM F838-05 to establish acceptable integrity test values.

Samples from each lot are subjected to microbial challenge test before final lot release.

100% Integrity Tested

Each AseptiVent VF- γ capsule filter is tested for integrity to comply with validated Acceptable Integrity Test Specifications.

Pressure, Temperature Endurance

AseptiVent VF- γ capsule filters are validated to endure high operating pressure and temperature conditions which may be encountered during use.

These filters are also validated to meet pre-determined burst pressure specifications to ensure user safety in case of inadvertent pressure build-up.

Bioburden Testing

Device bioburden is tested as per ISO 117 37-1 and assured to be <1000 cfu/device.

Endotoxin Testing

Aqeous extracts exhibit <0.25 EU/ml as established by Lumulus Amebocyte Lysate (LAL) test.

Gamma Sterilizability

AseptiVent VF- $\gamma\,$ are gamma sterilizable with up to 50 kGy of gamma irradiation.

Total Traceability

AseptiVent VF- γ capsule filters come with completely traceable lot numbers and unique identification number to facilitate easy and fast retrieval of manufacturing and quality control data associated with each filter.

These unique lot and identification numbers are laser etched on each filter device and also printed on the labels of the box in which individual filter is packed.

Packaging Integrity

AseptiVent VF- γ capsule filters are fitted with vent caps and are packed in double polyethylene bags to ensure package integrity during transit as well as to prevent particulate contamination while transferring to clean room assembly or process areas.

Other Regulatory Compliance

- Complies with USFDA 21 CFR 210.3(b)(6)
 for fiber release
- Complies with USFDA 21 CFR 177.1520 for indirect food additives
- Materials of construction tested for toxicity as per Biological Reactivity Tests, In vivo, USP <88> for class VI Plastics

Easy Connect

Datasheet

Widest Range of End Connections

Critical nature of biopharmaceutical processes involving steps such as sterile venting, air sparging, fermentor exhaust etc requires high quality, reliable, flexible and functionally convenient connectivity with filters.

mdi filters offer a wide range of reliable end connections for functional convenience and customized connectivity.

Validated for Performance

These end connections are manufactured with tight dimension tolerance and are validated for strength and connection integrity under extreme use conditions as well as for their ability to withstand prevalent sterilization methods including gamma irradiation and autoclaving.





1/2" HB



1/4" SHB



Male Luer Slip Outlet for 25 mm

1/2" Single Stepped HB

1¹/₂" Sanitary Flange



Quick Connector



Female Luer Lock Inlet for 25 mm

Some end connections available with AseptiVent VF-y

Customized Connectivity

mdi filters are available in a wide range of end connections and are also customized to offer different inlet-outlet combinations to meet the unique connectivity needs in biopharmaceutical process assemblies where, for example, stainless steel components with sanitary flange connections are sometimes required to be connected to single use disposable systems through quick-connectors or hose barb connections.



1¹/₂" Sanitary Flange to 1/2"Barb Hose

1¹/₂" Sanitary Flange to ¾" Sanitary Flange





HighSecurity ¹/₂" hose barb connection



DST DVLV01X1431D

Linear Upscaling from R&D to Production Process

Datasheet

Scientists in process development labs working with cell factories or small bioreactors require small area hydrophobic filters for air/gas filtration or sterile venting.

A scale up of these processes for larger productions requires larger area devices.

mdi offers a wide range of AseptiVent VF- γ Hydrophobic PVDF capsule filters to provide linear scale up from lab scale to pilot scale to full scale biopharmaceutical manufacturing processes. The appropriate size filter can be selected on the basis of the bioreactor size and required flow rates.



AseptiVent VF-γ 25 mm, 5 cm²



*AseptiVent VF-*γ 50 mm, 20cm²



*AseptiVent VF-*γ 1″, 250cm²



*AseptiVent VF-*γ 2″, 500cm²



*AseptiVent VF-*γ 5″, 1000cm²



AseptiVent VF-γ **8″, 2000cm²**

Bioreactor Size	Filter Devices	EFA* (Nominal)
200 ml Cell Factories	<i>AseptiVent VF-</i> γ 25 mm	5 cm ²
Up to 1 liter Cell Factories	<i>AseptiVent VF-</i> γ 37 mm	10 cm ²
Up to 5 liter	<i>AseptiVent VF-</i> γ 50 mm	20 cm ²
Up to 50 liter	AseptiVent VF-γ 1″	250 cm ²
Upto 100 liter	AseptiVent VF-γ 2"	500 cm ²
Upto 300 liter	AseptiVent VF-γ 5″	1000 cm ²
Upto 1000 liter	AseptiVent VF-γ 8"	2000 cm ²
Upto 5000 liter	AseptiVent VF-γ 10″	6000 cm ²

*Effective Filtration Area



*AseptiVent VF-*γ 10", 6000cm²

Specifications 0.2μm *AseptiVent* VF-γ

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		Constr	uction								
Size		25 mm	37 mm	50 mm							
Effective Filtratio	n Area (Nominal)	5 cm ²	10 cm ²	20 cm ²							
Membrane		0.2 μm Hydro	ophobic PVDF								
Support Layers		Poly	Polyester								
Plastic Parts		Gamma Stable	Gamma Stable Polypropylene								
Operational Rad	dius	15 mm	23 mm	28 mm							
		Opera	ational								
Max. Operating T	emperature	80° C @ <u><</u> 0.5 Kg/cm ² (7psi)									
Max. Differential	Pressure	1.5 Kg/cm² (22 psi) @ 30° C									
Minimum Acceptable Bubble Point with 50% IPA/Water		\geq 1.27 Kg/cm ² (18 psi)									
	By Irradiation	Gamma Irradiatable up to 50 kGy									
Sterilization	By Autoclave	Autoclavable at 125 °C for 30minutes Can not be in line steam sterilized	utoclavable at 125 °C for 30minutes, 1 Cycle after gamma irradiation. an not be in line steam sterilized								
		Assura	nce								
Toxicity		Passes biological reactivity test, In Vive	o, as per USP <88> for Class VI plastics								
Bioburden		Bioburden level is < 1000 cfu/filter device as per ANSI/AAMI/ISO 11737-1									
Bacterial Retention	on	LRV> 7 for <i>B. diminuta</i> per cm ² of filter area as per ASTM F 838-05 against liquid bacterial challenge									
Bacterial Endoto	xin	Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test as per USP <85>									
Non Fiber Releas	ing	Passes test as per USP and comply wit	Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release								
Particle Shedding	9	The filtrate complies with USP <788> test for particulate matter in injections									
Oxidizable Subst	ances	Passes test as per USP <1231>									
Indirect Food Ad	ditive	All Polypropylene components meet the FDA Indirect Food Additive requirements cited in 21 CFR 177.1520									
Good Manufactu	ring Practice	These products are manufactured in a facility which adheres to Good Manufacturing Practices									
Quality Manager	nent System	ISO-9001 Certified									
		DMF No. 015554									

Specifications 0.2μm *AseptiVent* VF-γ (1", 2", 5", 8")

Construction 1″ 2″ Size 5″ 8″ Effective Filtration Area (Nominal) 250cm² 500cm² 1000cm² 2000 cm² Membrane 0.2 µm Hydrophobic PVDF Support Layers Polyester Body and Core Gamma Stable Polypropylene **Operational Radius** 30 mm 65 mm 65 mm 65 mm (with Vent/ Drain) Vent and Drain 1/4" Hose Barb with Silicone "O" ring Operational 80° C @ 2 Kg/cm² (30psi) Max. Operating Temperature 4Kg/cm² (60psi) @ 30° C Max. Differential Pressure Minimum Acceptable \geq 1.27 Kg/cm² (18 psi) Bubble Point with 50% IPA Gamma Irradiatable up to 50 kGy Sterilization By Gamma Irradiation Assurance Toxicity Passes biological reactivity test, In Vivo, as per USP <88> for Class VI plastics Bioburden Bioburden level is < 1000 cfu/filter device as per ANSI/AAMI/ISO 11737-1 LRV> 7 for *B. diminuta* per cm² of filter area as per ASTM F 838-05 (liquid bacterial challenge) **Bacterial Retention** Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test **Bacterial Endotoxin** as per USP <85> Non Fiber Releasing Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release Particle Shedding The filtrate complies with USP <788> test for particulate matter in injections Oxidizable Substances Passes test as per USP <1231> Indirect Food Additive All Polypropylene components meet the FDA Indirect Food Additive requirements cited in 21 CFR 177.1520 These products are manufactured in a facility which adheres to Good Manufacturing Practices Good Manufacturing Practice **Quality Management System** ISO-9001 Certified USFDA DMF No. 015554

Datasheet

Specifications 0.2μm *AseptiVent* VF-γ 5", 10", 20", 30"

Construction 5″ 10″ Size 20″ 30″ Effective Filtration Area (Nominal) 3000cm² 6000cm² 12000cm² 18000 cm² Membrane 0.2 µm Hydrophobic PVDF Support Layers Polyester Body and Core Gamma Stable Polypropylene **Operational Radius** 78 mm 78 mm 78 mm 78 mm (with Vent/ Drain) Vent and Drain 1/4" Hose Barb with Silicone "O" ring Operational Max. Operating Temperature 80° C @ 2Kg/cm² (30psi) 4Kg/cm² (60psi) @ 30° C Max. Differential Pressure Minimum Acceptable \geq 1.27 Kg/cm² (18 psi) Bubble Point with 50% IPA Sterilization By Gamma Irradiation Gamma Irradiatable up to 50 kGy Assurance Toxicity Passes Biological reactivity test, In Vivo, as per USP <88> for Class VI plastics Bioburden Bioburden level is < 1000 cfu/filter device as per ANSI/AAMI/ISO 11737-1 LRV> 7 for *B. diminuta* per cm² of filter area as per ASTM F 838-05 (liquid bacterial challenge) **Bacterial Retention** Aqueous extracts exhibit < 0.25 EU/ml as established by Limulus Amebocyte Lysate (LAL) Test **Bacterial Endotoxin** as per USP <85> Passes test as per USP and comply with USFDA 21 CFR Part 210.3(b)(6) for fiber release Non Fiber Releasing Particle Shedding The filtrate complies with USP <788> test for particulate matter in injections **Oxidizable Substances** Passes test as per USP <1231> All Polypropylene components meet the FDA Indirect Food Additive requirements cited in 21 CFR 177.1520 Indirect Food Additive These products are manufactured in a facility which adheres to Good Manufacturing Practices Good Manufacturing Practice **Quality Management System** ISO-9001 Certified USFDA DMF No. 015554

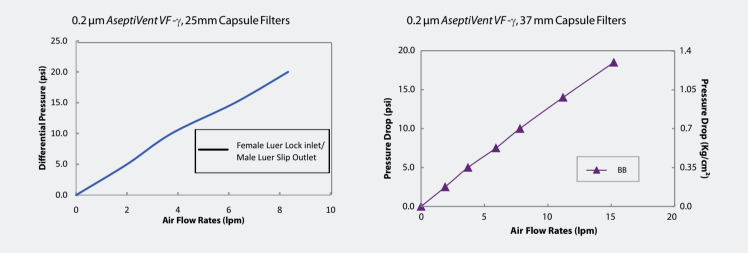
Datasheet

Typical Air Flow Rates

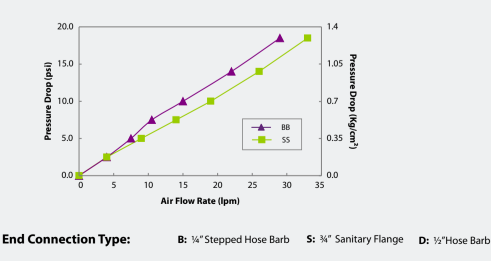
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AseptiVent VF-γ is produced using a high hydrophobicity PVDF membrane. This ensures good flow rates even with high moisture content in the inlet air.

AseptiVent VF-y capsule filters are designed to offer high air/gas flow rates at low differential pressures.



 $0.2\,\mu m$ AseptiVent VF- γ , 50 mm Capsule Filters

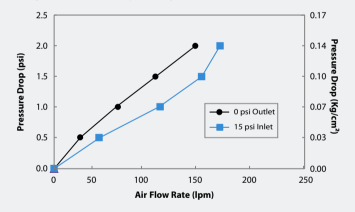


Typical Air Flow Rates

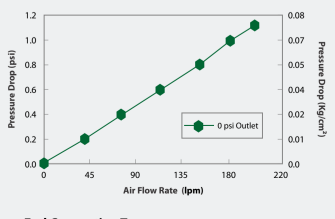
2.5 0.17 0.14 Pressure Drop (Kg/cm²) 0.07 2.0 Pressure Drop (psi) 1.5 1.0 0 psi Outlet 15 psi Inlet 0.03 0.5 0.0 0.00 21 35 0 7 14 28 Air Flow Rate (Ipm)

0.2 μm AseptiVent VF-γ, 1" Capsule Filters, EE Connection

0.2 μm AseptiVent VF-γ, 5" Capsule Filters, EE Connection

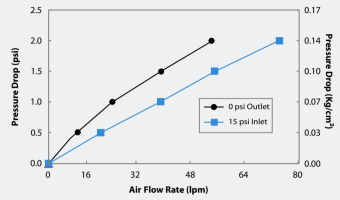


0.2 µm AseptiVent VF -7, 5" Large Capsule Filters, EE Connection

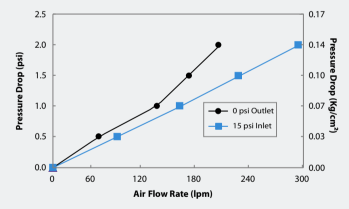


End Connection Type: E: 1½" Sanitary Flange

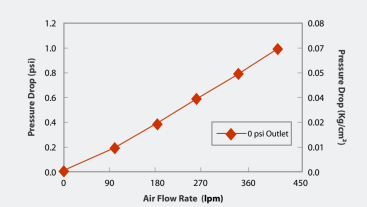
0.2 μm AseptiVent VF-γ, 2" Capsule Filters, EE Connection



0.2 µm AseptiVent VF - y, 8" Capsule Filters, EE Connection



0.2 μm AseptiVent VF-γ, 10" Capsule Filters, EE Connection



Datasheet

DST DVLV01X1431D

Ordering Information

Datasheet

0.2 μm AseptiVent VF-γ 25mm PVDF Membrane Capsule filter

Туре		Size		Pore S	ize	Inlet/Outle	et	Radiation Sterilizable		x	Sterility		Pack Size	
	Code		Code		Code		Code		Code			Code		Code
AseptiVent VF-γ	IVFX	25 mm	06	0.2µm	01	1/8" Hose Barb	Н	Yes	R		Non Sterile	1	100	04
						Female Luer Lock	М	No*	Х		Gamma Sterile	3		
						Male Luer Slip	N							
						Male Luer Lock L								
						1/4" Hose Barb B								
Example:						L								
IVFX		06		01		MN		R		Х	1		04	

* Gamma irradiated filters cannot be gamma sterilized again

Example for Non Sterile: IVFX0601MNRX104

Example for gamma Sterile: IVFX0601MNXX304

0.2 μ m AseptiVent VF- γ 37mm, 50mm PVDF Membrane Capsule filter

Туре		Size		Pore Size		Inlet/Outlet		Radiation Sterilizable				X	Sterility		Pack	Size
	Code		Code		Code		Code		Code			Code		Code		
AseptiVent VF-γ	IVFX	37 mm	08	0.2µm	01	1⁄4″ SHB	В	Yes	R		Non Sterile	1	10	02		
		50 mm	10			³ ⁄ ₄ " Sanitary Flange	S	No*	Х		Gamma Sterile	3				

Example:

IVFX 10 01	BB	R	Х	1	02
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* Gamma irradiated filters cannot be gamma sterilized again

Example for Non Sterile: IVFX0801BBRX102

Example for gamma Sterile: IVFX0801BBXX302

Note: Inlet/Outlet Connections and Pack Sizes available with different diameter filters as follows:

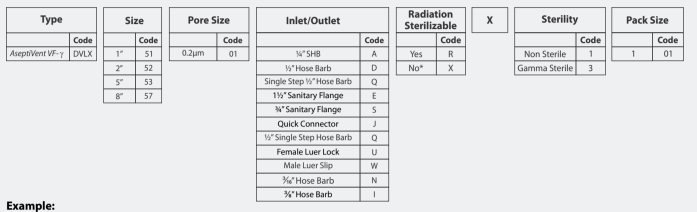
Connections Available									
Inlet/Outlet	25mm	37mm	50mm						
¹ / ₄ " - ³ / ₄ " Stepped Hose Barb	Х	\checkmark	\checkmark						
¾" Sanitary Flange	Х	х							
Female Luer Lock	Inlet Only	х	х						
Male Luer Slip Outlet Only		х	х						
1∕‰" Hose Barb		х	х						
Male Luer Lock	Outlet Only	х	Х						
1⁄4" Hose Barb		х	Х						

Dimension (in mm)	Inlin	e Capsule Fi	ilters
Inlet/ Outlet	25mm	37mm	50mm
1⁄4" - ¾" Stepped Hose Barb I/O	-	64	79
¼" Single Step Hose Barb I/O	38	-	-
³ ⁄4″ Sanitary Flange I/O	-	-	51
Female Luer Lock Inlet/ Male Luer Slip Outlet	23	-	-
¹ ∕ ₈ " Hose Barb I/O	36	-	-
Operational Radius	15	23	28

Ordering Information

Datasheet

0.2 μm AseptiVent VF-γ PVDF Membrane Capsule filter



DVLX 57	01	EE	R	х	1	01
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* Gamma irradiated filters cannot be gamma sterilized again

Example for Non Sterile: DVLX5301QQRX101 Example for gamma Sterile: DVLX5301QQXX301

Note: Inlet/Outlet Connections available with different Sizes/Length as follows:

Inlet/Outlet		Size/Le	ngth		Dimensions (in mm)		Small Capsule Filters				
	1″	2″	5″	8″	End Connections	1″	2″	5″	8″		
1/4" Stepped Hose Barb	\checkmark	\checkmark	\checkmark	\checkmark			400	170			
½"Hose Barb			V		1/4" SHB I/O	94	122	172	223		
	,	,	,		³ 4" Sanitary Flange Inlet I/O	91	103	155	205		
1½ "Sanitary Flange	\checkmark	\checkmark	\checkmark	\checkmark	Quick Connector	100	111	163	212		
¾" Sanitary Flange	\checkmark	\checkmark	\checkmark	\checkmark	1½" Sanitary Flange I/O	91	110	161	211		
Quick Connector					1/2" Hose Barb I/O	90	112	164	215		
½" Single Step Hose Barb	x				¹ / ₂ " Single Step Hose Barb I/O	-	115	165	217		
Female Luer Lock			√		11/2" Sanitary Flange Inlet 1/2" Single Step Hose Barb Outlet	-	111	162	212		
Male Luer Slip	Outlet Only	Х	x	х	3/8" Hose Barb I/O	-	115	165	217		
¾6″ Hose Barb			Outlet Only	х	Operational Radius	30	65	65	65		
‰" Hose Barb	х		\checkmark								

Ordering Information

Datasheet

0.2 μm AseptiVent VF-γ PVDF Membrane Capsule filter

Туре		Si	ze	Pore	Size	Inlet/Outlet		Radia Steriliz		Inline/T	-Line	
	Code		Code		Code		Code		Code		Code	
Aseptivent VF-γ	LVLX	5″	53	0.2µm	01	1/2" Single Step Hose Barb	Q	Yes	R	Inline	Х	
		10″	54			1½" Sanitary Flange	E	No*	Х	T-line**	Т	
		20″	55			³ 4" Sanitary Flange	S	•				
		30″	56			³∕‰" Hose Barb	I					
						1" Hose Barb	Z					

e	Sterility	'	Pack S	ize	
de		Code		Code	
(Non Sterile	1	1	01	
-	Gamma Sterile	3			

Example:

LVLX 54 01 EE R X 1	01	
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* Gamma irradiated filters cannot be gamma sterilized again

Example for Non Sterile: LVLX5401QQRX101 Example for gamma Sterile: LVLX5401QQXX301

** T-line is not available in 5" Capsule filter

** T-line Capsule Filter are available with 11/2" Sanitary Flange I/O Connections Only

Note: Inlet/Outlet Connections available with different Sizes/Length as follows:

	Inline				T-Line			Dimensions (in mm)		
Inlet/Outlet	5″	10″	20″	30″	10″	20″	30″	End Connections		
	1	,						1½" Sanitary Flange I/O		
¹ / ₂ " Single Step Hose Barb			V	\checkmark	X	X	Х	³ ⁄ ₄ " Sanitary Flange I/O		
1½" Sanitary Flange	\checkmark	½" Single Step Hose Barb I/O								
³ 4" Sanitary Flange			х	х	х	х	х	1½" Sanitary Flange Inlet ½" Hose Barb Outlet		
³‰" Hose Barb				V	x	x	х	³ ∕ ₈ " Hose Barb I/O		
	,	,		,				1" Hose Barb I/O		
1" Hose Barb	х	\checkmark	\checkmark	\checkmark	Х	х	х	Operational Radius		

Dimensions (in mm)	Inl	ine Cap	sule Filt	T-line Capsule Filters			
End Connections	5″	10″	20″	30″	10″	20″	30″
1½" Sanitary Flange I/O	207	326	605	865	338	588	848
¾" Sanitary Flange I/O	212	345	х	х	х	х	х
1/2" Single Step Hose Barb I/O	217	332	628	888	х	х	х
1½" Sanitary Flange Inlet ½" Hose Barb Outlet	203	332	618	878	x	x	х
¾"Hose Barb I/O	211	330	618	878	х	х	х
1" Hose Barb I/O	х	405	635	895	x	х	х
Operational Radius	78	78	78	78	78	78	78

Advanced Microdevices Pvt. Ltd.

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