

PRODUCT DATA SHEET

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Lipodisq™ Styrene:Maleic Acid Copolymer 1:1 [SMA-100]

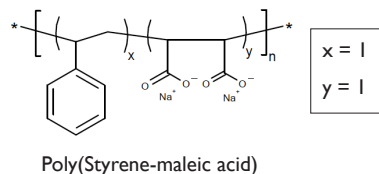
Cat. No.: IAX-700-201

Lot. No.:

Synonyms

Lipodisq™ Styrene:Maleic Anhydride Copolymer 1:1 (pre-hydrolyzed), [SMA-100], P(SMA) 1:1

Empirical Formula



Molecular Formula

$C_{12}H_{10}O_4Na_2$

Size

100mg

MW

Polymer MW average: M_w : 5,500 M_w : MW based on weight M_n : 2,100 M_n : MW based on number

CAS

26762-29-8

Purity

>98%

Solution pH

4.50 +/- 0.25 in 5% ddWater

Solubility

Soluble in water, and buffer solutions (pH 3.50-5.50) to allow the formulation of a proprietary, thermostable, aqueous lipid nanoparticle (Lipodisq™, Malvern Cosmeceutics Ltd., Malvern UK). Avoid the use of buffers with divalent ions such as Ca^{++} or Mg^{++} (>2mM) or pH <3.5 or >5.5, which can cause P(SMA) precipitation or interfere with SMA-Lipid Particle formulation or stability.

Formulation with Phospholipids

Lipodisq™ are nanosized lipid-based discoidal particles that can be manufactured with P(SMA)/SMA-LP and lipids such as DMPC (1,2-Dimyristoyl-sn-glycero-3-phosphocholine) (14:0 PC) (DMPC: IAX-700-400) to incorporate hydrophobic, poorly water-soluble active compounds, such as peptides, lipids, lipoproteins, transmembrane proteins and glycolipids. Applications of Lipodisq™ include functional and structural characterisation of the cargo and drug delivery with improved bioavailability, and biological half-life *in vivo* (PD/PK) or delivery of antigens preserved in their native conformation for immunization purposes.

Appearance

White powder

Handling

Keep dry. Avoid skin and eye contact.

Shipping

Ambient

Storage

2-8°C or 15-25°C

Stability

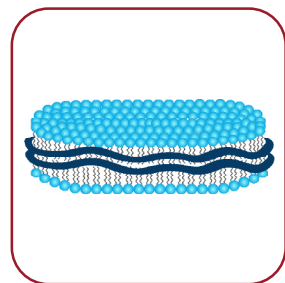
Upon receipt, store product at ambient temperature. Freezing is not recommended. In its unopened original vial, the product is stable for at least 36 months when stored at ambient temperature. Once the glass vial is opened, or solubilized in sterile, endotoxin-free ddWater (IAX-900-902) or for example in pH adjusted, diluted NaCl /acetate buffer (IAX-900-010) and aliquoted into sterile vials under sterile conditions, the SMA polymer solution (e.g. at 5%) remains stable for an additional one month when stored at 2-8°C and kept sterile.

MSDS

Available on request

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- A nanoparticle (11-40nm) drug delivery system comprising a discoidal phospholipid bilayer membrane stabilised by a chaperone molecular annulus.
- Lipodisq™ polymers are available as 4 defined structures (1:1, 2:1, 3:1 and 4:1 Styrene to Maleic Acid (SMA) ratios) each individually operational within a selected pH range for optimal working conditions.
- Components are batch-tested for Lipodisq™ formation using buffer systems available, which are tested for nano-formulated drug analysis by Dynamic Light Scattering (DLS). These buffer solutions are endotoxin-tested and sterile.
- Lipodisq™ formation is highly efficient and nanodiscs show a good safety profile and are suitable for *in vitro* and *in vivo* (in experimental animals) investigations.

Answering the call for Membrane Protein (MP) and MP reconstituting membrane simulants which do not necessitate the use of mediating detergent, is a copolymer of styrene and maleic anhydride subsequently hydrolysed into amphipathic polystyrene-co-maleic acid (SMA).

Lipodisq™ Styrene:Maleic Acid Copolymer / P(SMA) System: Introduction

Nanodiscs resulting from SMA are also referred to as styrene maleic acid lipid particles (SMALP). SMA-lipid particles are formed by directly extracting membrane proteins either from native cellular membranes (giving native nanodiscs) or from an intermediary MP-reconstituted synthetic membrane system to ultimately form self-assembled nanodisc structures of a general 10–12 nm diameter. The demonstration of SMA-lipid particles as a monodisperse MP reconstitution system was reported in 2009, although the interaction of SMA with phospholipids to generate disc shaped structures (now known as empty lipid nanodiscs), was previously established and investigated for use as a drug delivery system years preceding this discovery.

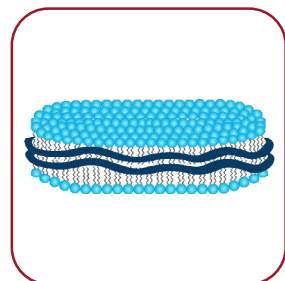
At a physiological pH of 7–8, SMA monomer ratios of 2:1 and 3:1 are most commonly used due to their optimal efficiency for nanodisc extraction from phospholipid bilayers. Abilities of SMA to create monodisperse nanodiscs with size flexibility facilitates the reconstitution of a range of oligomeric MPs and MP complexes for analysis by various studies including fluorescence microscopy, NMR and single-particle cryo-EM. Whereas in a statistical version of SMA, like the commercial Malvern polymer Lipodisq™, monomers are evenly distributed throughout the polymer chain sequence in proportion to their ratio as well as exhibiting greater dispersity in chain length.

Lipodisq™ Procedure Overview

- Published procedures using a ratio of 2:1 (w/w) for P(SMA)/SMA polymer and phospholipid, such as 100mg P(SMA)/SMA polymer and 50mg DMPC. Selected hydrophobic drug candidate or peptides or transmembrane proteins (MP) are mixed with the aqueous phospholipid emulsion making up 2.5%. This is stirred at temperatures above the phase transition temperature of the lipid (>24°C for DMPC), before aqueous P(SMA)/SMA polymer at 5% is added drop-wise and with pauses until a approximate volume ratio of 1:1 is reached and the lipid emulsion clears.
- Alternatively, P(SMA)/SMA polymer solutions are mixed with native (cell or bacterial) membranes to form native nanodiscs.
- Stirring time, pH, selected buffer type and strength (e.g. HEPES, NaCl, TRIS or PBS w/o Ca⁺⁺ and Mg⁺⁺) and temperature of the phospholipid emulsion containing the MP or active ingredient, need to be optimized.
- Further purification of formed Lipodisq™ can be achieved by ultracentrifugation at >100,000 x g to remove residual lipid, surplus P(SMA) polymer with the Lipodisq™ nanodiscs remaining in the supernatant. Alternatively, size exclusion column (SEC) methods can also be applied.

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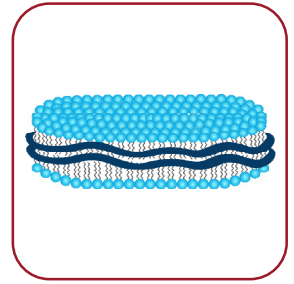
Catalogue	SMA Polymer Type	pH after dissolved in ddWater (5%)	Functional pH Range with Buffer	Molecular Weight (M_w/M_n)
IAX-700-201	Lipodisq™ Styrene:Maleic Acid Copolymer 1:1 [SMA-100]	4.25 +/- 0.25	3.50 - 5.50	5,500 / 2,100
IAX-700-202	Lipodisq™ Styrene:Maleic Acid Copolymer 2:1 [SMA-200]	7.00 +/- 0.25	5.50 - 8.00	7,500 / 2,700
IAX-700-203	Lipodisq™ Styrene:Maleic Acid Copolymer 3:1 [SMA-300]	7.50 +/- 0.25	6.00 - 9.00	9,500 / 3,050
IAX-700-204	Lipodisq™ Styrene:Maleic Acid Copolymer 4:1 [SMA-400]	9.25 +/- 0.25	8.00 - 10.50	10,500 / 4,500

Lipodisq™ SMA Polymer References

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- [2] *Applications of Synthetic Polymer Discoidal Lipid Nanoparticles to Biomedical Research.* Tanaka M. *Chem. Pharm. Bull.* (2022); 70:507
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Related Powered by Lipodisq™ Products for Nano-formulated Drug Delivery

IAX-700-100	Lipodisq™ Control Sterile Solution
IAX-700-101	Curcumin Lipodisq™ Sterile Solution
IAX-700-102	Melatonin Lipodisq™ Sterile Solution
IAX-700-103	Metformin Lipodisq™ Sterile Solution
IAX-700-104	Oxyresveratrol Lipodisq™ Sterile Solution
IAX-700-105	Resveratrol Lipodisq™ Sterile Solution
IAX-700-106	Umifenovir Lipodisq™ Sterile Solution
IAX-700-107	Dexamethasone Lipodisq™ Sterile Solution
IAX-700-108	Ambroxol Lipodisq™ Sterile Solution
IAX-700-109	Retinoic Acid Lipodisq™ Sterile Solution
IAX-700-201	Lipodisq™ Styrene:Maleic Acid Copolymer 1:1 [SMA-100]
IAX-700-202	Lipodisq™ Styrene:Maleic Acid Copolymer 2:1 [SMA-200]
IAX-700-203	Lipodisq™ Styrene:Maleic Acid Copolymer 3:1 [SMA-300]
IAX-700-204	Lipodisq™ Styrene:Maleic Acid Copolymer 4:1 [SMA-400]
IAX-700-400	DMPC (1,2-Dimyristoyl-sn-glycero-3-phosphocholine) (14:0 PC)

Endotoxin-free and Sterile Buffers and Related Products

IAX-900-001	PBS Endotoxin-free (sterile)
IAX-900-001DC	PBS Endotoxin-free (sterile) [For Nano-formulated Drug Analysis]
IAX-900-002	ddWater Endotoxin-free (sterile)
IAX-900-002DC	ddWater Endotoxin-free (sterile) [For Nano-formulated Drug Analysis]
IAX-900-003	Physiological Saline [Sodium Chloride 0.9% Endotoxin-free] (sterile)
IAX-900-003DC	Physiological Saline [Sodium Chloride 0.9% Endotoxin-free] (sterile) [For Nano-formulated Drug Analysis]
IAX-900-004	PBS with EDTA Endotoxin-free (sterile)
IAX-900-005	TRIS with EDTA [TE Buffer] (100x) Endotoxin-free (sterile)
IAX-900-006	EDTA (400mM) Endotoxin-free (sterile)
IAX-900-007	HEPES Buffer (500mM) Endotoxin-free (sterile)
IAX-900-008	DNA Loading Buffer with TRIS and EDTA (6x) (Blue)
IAX-900-009	HEPES Buffer (50mM) with NaCl [Sodium Chloride] (150mM) Endotoxin-free (sterile)
IAX-900-010	NaCl [Sodium Chloride] (1.5M) Endotoxin-free (sterile)
IAX-900-011	TRIS Buffer (1.5M) Endotoxin-free (sterile)
IAX-900-012	TRIS Buffer (30mM) with NaCl [Sodium Chloride] (150mM) Endotoxin-free (sterile)
IAX-900-013	PBS with Magnesium and Calcium Endotoxin-free (sterile)
IAX-900-014	ddWater with 0.9% Benzyl Alcohol [Bacteriostatic Water] Endotoxin-free (sterile)

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