

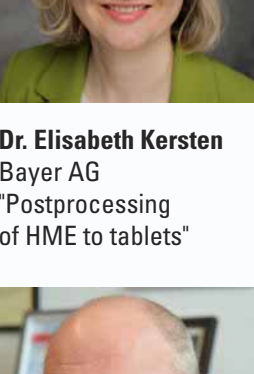
Dear Reader,

we are pleased to share an exciting update on Shin-Etsu Pharma's most recent activities including: the upcoming 4th Technical Seminar on Solubility Enhancement: from Screening to Downstreaming in May 2023 at Shin-Etsu's site in Wiesbaden, Germany; the most recent research articles pursued or contributed to by the Shin-Etsu Pharma Team; as well as the newly released Technical Information.

Announcement | Product Portfolio | Research Articles | Technical Informations | Exhibitions | Links | Contact

Announcement | 4th Technical Seminar on Solubility Enhancement

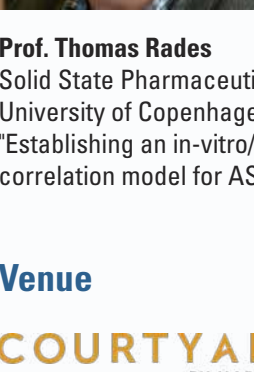
We are glad to announce our 4th Seminar on Amorphous Solid Dispersion (ASD). As technology provider in the field of ASD, Shin-Etsu, together with partners Alexanderwerk, Frewitt, PROCEPT and Thermo Fisher Scientific will share the latest insights on ASD development steps ranging from initial screening phase up to downstream processing including spray drying, hot melt extrusion (HME), roller compaction, milling and 3D printing. The program is comprised of practical lab demonstrations featuring the leading polymer for ASDs - Shin-Etsu AQOAT® (HPMCAS) and lectures from invited speakers with expertise in the field.



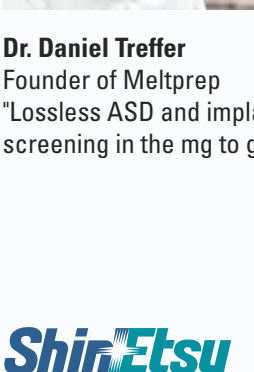
4th Technical Seminar on Solubility Enhancement From Screening to Downstreaming

SAVE THE DATE 11th-12th May 2023

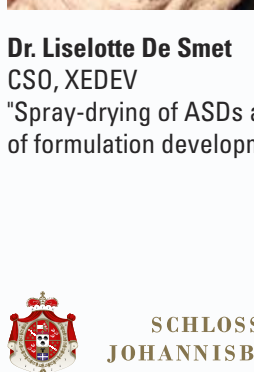
Confirmed speakers



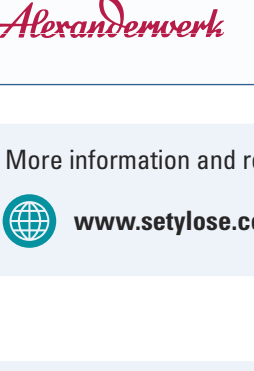
Dr. Elisabeth Kersten
Bayer AG
"Postprocessing of HME to tablets"



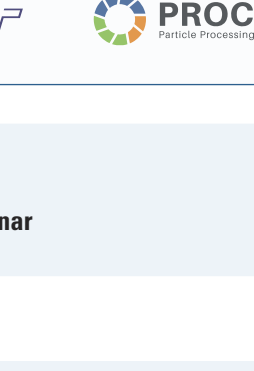
Dr. Christian Lübbert
Amofor GmbH
"Predicting the physical stability of amorphous solid dispersions"



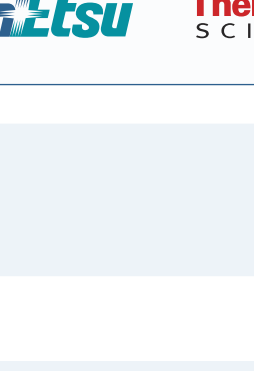
Dr. Alvaro Goyanes
UCL School of Pharmacy
"3D printing technologies to improve drug solubility"



Prof. Thomas Rades
Solid State Pharmaceuticals, University of Copenhagen
"Establishing an in-vitro/in-vivo correlation model for ASDs"




Dr. Daniel Treffer
Amofor Meltrep
"Lossless ASD and implant screening in the mg to g scale"




Dr. Liselotte De Smet
CSO, XEDEV
"Spray-drying of ASDs as a part of formulation development"


Venue



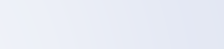




Lecture Venue
Courtyard by Marriott
Ostring 9, Wiesbaden-Nordenstadt
65203 Wiesbaden, Germany



Demonstration Venue
SE Tylose GmbH & Co. KG
Kasteler Straße 45
65203 Wiesbaden, Germany



Networking Event
Johannisberg Castle
65366 Geisenheim
Germany



More information and registration:
www.setylose.com/en/pharma-seminar

Product Portfolio

Cellulose Ethers

for Pharmaceutical Applications

METOLOSE®
Methylcellulose, Hypromellose
USP 1 EP 1 JP

TYLOPUR®
Hypromellose
USP 1 EP 1 JP

PHARMACOAT®
Hypromellose
USP 1 EP 1 JP

HPMCP
Hypromellose Phthalate
NF 1 EP 1 JP

METOLOSE® SR
Hypromellose
USP 1 EP 1 JP

TYLOPUR® SR
Hypromellose
USP 1 EP 1 JP

L-HPC
Low-Substituted Hydroxypropylcellulose
NF 1 EP 1 JP

Shin-Etsu AQOAT®
Hypromellose Acetate Succinate
NF 1 JP

* product available in EMEA, North America and LATAM

L-HPC

Low substituted Hydroxypropyl Cellulose NF, JP, EP



Dual Functions, Multiple Benefits...

- Good compressibility
- Disintegrate into smaller particles
- Higher stability
- No peroxide
- Non ionic in nature
- Low water activity
- Simplified formulations

Shin-Etsu was established in 1926 and began producing cellulose derivatives in 1962. Pharmaceutical grades of cellulose ethers have been manufactured since 1971. Regulated grades of cellulose ethers used in pharmaceutical applications are manufactured in Naotsu, Japan (METOLOSE®, METOLOSE® SR, PHARMACOAT®, L-HPC, HPMCP and Shin-Etsu AQOAT®) and Wiesbaden, Germany (TYLOPUR® and TYLOPUR® SR).

L-HPC, low-substituted hydroxypropyl cellulose NF, JP, EP, was developed by Shin-Etsu and first approved in 1977 as disintegrant for tablets in Japan. L-HPC is a non-ionic, multi-functional excipient based on a cellulose backbone with a low amount of hydroxypropyl groups. Thus, L-HPC is not soluble in water but manifests swelling properties in this media.

Typical applications of L-HPC are: binding and disintegrating, stability enhancement, pellet extrusion and continuous manufacturing. On top of that, L-HPC reduces the residual die-wall force and ejection force resolving capping issues, a typical problem in the tableting process.

L-HPC manifests excellent properties as binder for direct compression, wet granulation and dry granulation (roller compaction).

L-HPC can simplify immediate release tablet formulation as only one excipient is needed to assure good tablet hardness and quick

disintegration. Thus L-HPC is a key excipient for reducing the tablet size (mini-tablet formulation).

One of the additional benefits of L-HPC is linked to its non-ionic-nature and absence of peroxide leading to a better stability of the drugs by avoiding interactions with the API.

L-HPC is also applicable for pellet extrusion (micronized grades). Here, L-HPC provides wet mass with a "buffer effect" – wet mass accepts a wider range of water content, and plasticizes it to enhance extrusion speed and yield.

For capsule filling, the swelling property of L-HPC actively contributes to improving the disintegration and dissolution of the cake formed during the capsule filling process.

L-HPC also plays a key role in continuous manufacturing. Its optimized particle shape and particle size distribution is designed to maximize the flow property and multifunctionality required for direct compression, wet granulation and dry granulation performed in a continuous manner.

There are 9 commercially available grades of L-HPC, which vary in physico-chemical properties fitting various pharmaceutical applications.

More information and registration:
www.setylose.com/en/products/healthcare/l-hpc

Research Articles

1. Downstream Processing of Itraconazole:HPMCAS Amorphous Solid Dispersion: From Hot-Melt Extrudate to Tablet Using a Quality by Design Approach

Despite the importance of hot-melt extrusion in the preparation of amorphous solid dispersions (ASDs) for solubility improvement of poorly soluble drugs, there are few reports on milling of hot-melt extrudates. In a recently published peer-reviewed article, the Shin-Etsu team in Totowa (NJ, USA) in cooperation with Thermo Fisher Scientific in Karlsruhe (Germany) reports on the milling of ASD extrudates using a design of experiments (Scheme 1). The study comprises the preparation of three different milling feedstocks of itraconazole and hypromellose acetate succinate (Shin-Etsu AQOAT(R)) ASD by variation of the downstream process after hot melt extrusion (hand-cut, pelletization, chill roll flakes).

These different feedstocks were subjected to milling in a pilot scale hammer mill and the milled extrudates were compressed into tablets. Interestingly, it was found that by careful selection of milling speed, a similar particle size distribution (PSD) was observed after the milling of hand-cut filaments in comparison with the milling of pellets or chill roll flakes, demonstrating an easy transfer from early-formulation hand-cut processing at the lab scale to industrial-applicable processes. Furthermore, the tablet formulation with milled chill roll flakes showed improved compactibility over milled hand-cut filaments, despite a similar PSD. In summary, the study demonstrates the facile milling of an hot-melt extruded ASD based on HPMCAS as carrier polymer at ambient temperature.

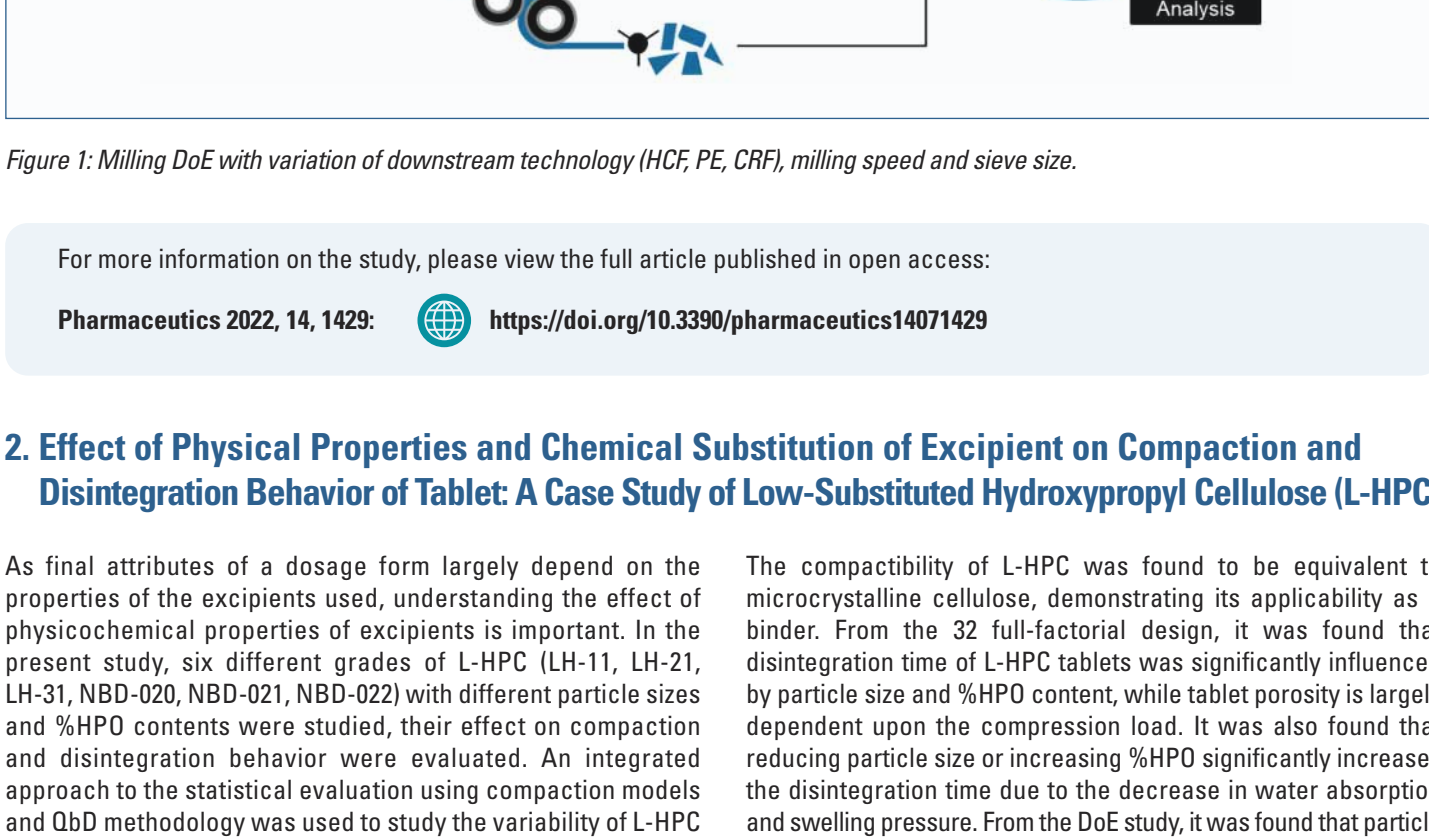


Figure 1: Milling DoE with variation of downstream technology (HCF, PE, CRF), milling speed and sieve size.

For more information on the study, please view the full article published in open access:
Pharmaceutics 2022, 14, 1429: <https://doi.org/10.3390/pharmaceutics14071429>

2. Effect of Physical Properties and Chemical Substitution of Excipient on Compaction and Disintegration Behavior of Tablet: A Case Study of Low-Substituted Hydroxypropyl Cellulose (L-HPC)

As final attributes of a dosage form largely depend on the properties of the excipients used, understanding the effect of physicochemical properties of excipients is important. In the present study, six different grades of L-HPC (LH-11, LH-21, LH-31, NBD-020, NBD-021, NBD-022) with different particle sizes and %HPO contents were studied, their effect on compaction and disintegration behavior were evaluated. An integrated approach to the statistical evaluation using compaction models and QbD methodology was used to study the variability of L-HPC grades on the quality attributes of tablets. It was found that all tested grades of L-HPC are highly compressible, undergoing predominantly plastic deformation. Additionally, it was found that particle size has a large effect on deformation behavior/ compressibility of L-HPC, and LH-31 with smaller particles was found to be less compressible compared to other grades of L-HPC such as LH-21 and LH-11.

The compactibility of L-HPC was found to be equivalent to microcrystalline cellulose, demonstrating its applicability as a binder. From the 32 full-factorial design, it was found that disintegration time of L-HPC tablets was significantly influenced by particle size and %HPO content, while tablet porosity is largely dependent upon the compression load. It was also found that reducing particle size or increasing %HPO significantly increases the disintegration time due to the decrease in water absorption and swelling pressure. From the DoC study, it was found that particle size of L-HPC grades demonstrates a larger degree of effect compared to %HPO on disintegration time, showing a higher coefficient value. Thus, based on the present study, it can be concluded that variability in L-HPC can have a significant influence on critical quality attributes of tablets such as compaction and disintegration behavior, and thus the selection of the correct grade of L-HPC is critical for successful formulation development.

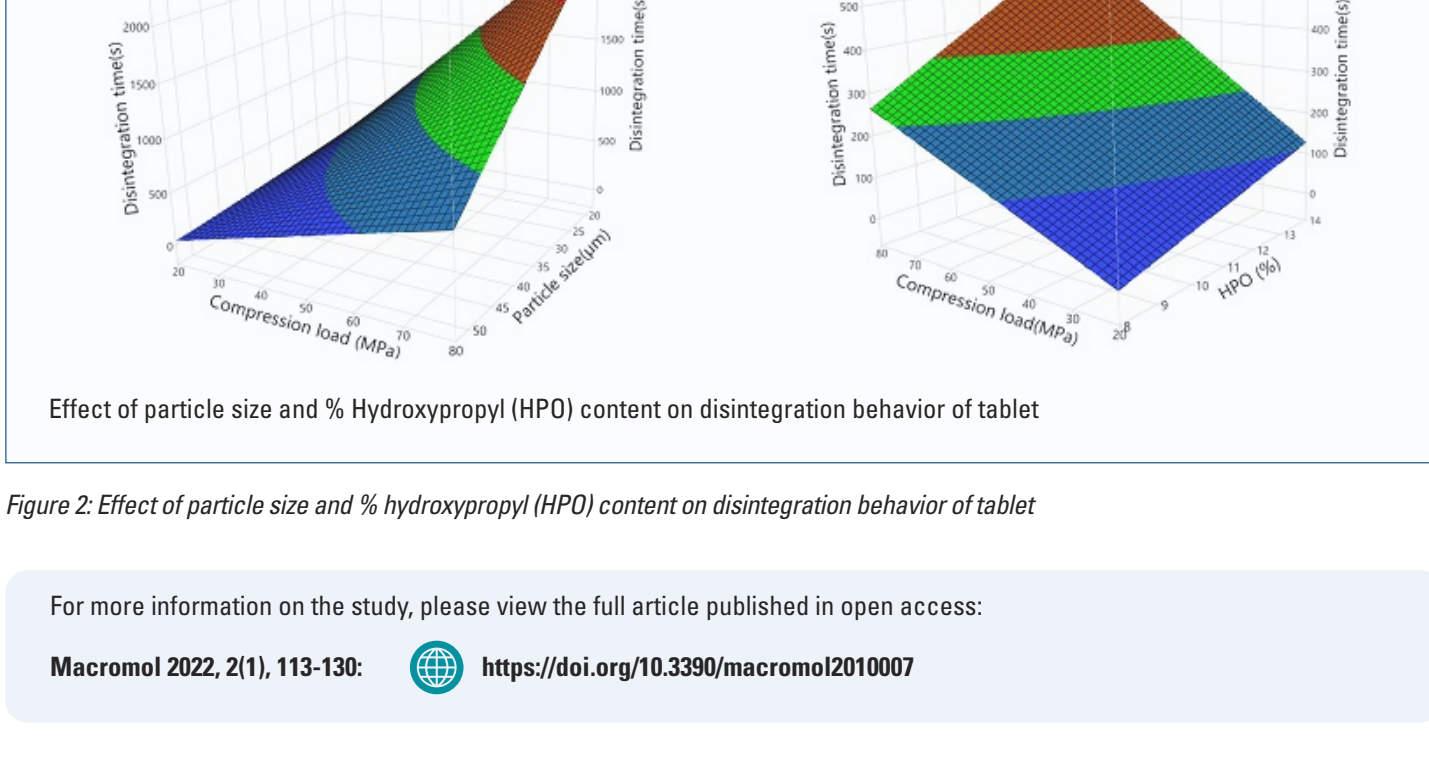


Figure 2: Effect of particle size and % hydroxypropyl (HPO) content on disintegration behavior of tablet

For more information on the study, please view the full article published in open access:
Macromol 2022, 2(1), 113-130: <https://doi.org/10.3390/macromol2010007>

3. Stability and intrinsic dissolution of vacuum compression molded amorphous solid dispersions of efavirenz

In this research paper from the University of Copenhagen, co-authored with Shin-Etsu, Losan Pharma and Habke Pharma, the effect of polymer type and polymer loading on stability and intrinsic dissolution of amorphous solid dispersions is discussed. The amorphous solid dispersions of efavirenz, an HIV drug, and the different polymer carriers were prepared by vacuum compression molding using the Meltrep device. Melting point depression studies yielded the solubility of efavirenz in the respective polymer and ASDs were prepared around the solubility limit. ASDs with efavirenz loading below or close to the solubility limit did not crystallize at different stability conditions for seven months. All ASDs above the solubility limit crystallized at high humidity. ASDs above the solubility limit with cellulose derivatives (Shin-Etsu AQOAT® HPMCAS, PHARMACOAT® HPMC) were kinetically stable at 22°C/23% RH over the course of the study, while others showed signs of crystallization. The intrinsic dissolution study gave insights into the solubility behavior of the ASD discs which depends on polymer solubility and API load in the ASD.



Figure 3: Study design

For more information on the study, please view the full article published in open access:
Int. J. Pharm. 2023, 632, 122564: <https://doi.org/10.1016/j.ijpharm.2022.122564>

4. Shin-Etsu's solutions for the typical tablet defect

During the manufacturing of tablets, defects like capping, lamination, sticking, etc. can occasionally be encountered. These tablet defects can originate from any upstream operation units and from the tablet press. In the upstream part of the process, the quality and concentration of raw materials as well as process optimization are the key variables for manufacturing of good quality tablets. Tablet defects, if not avoided at the right time, may lead to the recall of product, loss of time and decreased productivity. The attached troubleshooting guide shows the most common tablet defects that can be easily avoided by choosing appropriate Shin-Etsu Pharma excipients.



Figure 4: The six major defects during tableting.

More information:
<https://www.setylose.com/en/knowledge-base/healthcare/brochures#l-hpc-10253>

New Technical Information

Shin-Etsu is continuously striving to deliver new technical data for your formulation development. Please take a look at our new technical information sheets that are available from your local Shin-Etsu representative.

Shin-Etsu AQOAT®

A-073
Mechanical Properties of Hot Melt Extruded Amorphous Solid Dispersion with Shin-Etsu AQOAT®

www.metoelose.jp/en

<https://www.setylose.com/en/knowledge-base/healthcare/technical-information#shin-etsu-aqoat-10399>

HPMCP

H-023
Application of Hypromellose Phthalate in Extended Release (ER) Matrix Tablet formulation

www.metoelose.jp/en

<https://www.setylose.com/en/knowledge-base/healthcare/technical-information#hpmcp-10400>

METOLOSE® SR

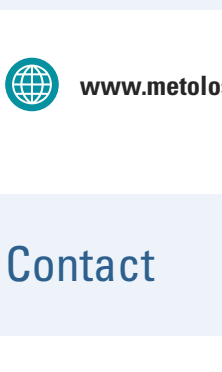
SR-021
Impact of combination of METOLOSE® SR grades in Carbamazepine Extended Release (ER) Matrix Tablet formulation

www.metoelose.jp/en

<https://www.setylose.com/en/knowledge-base/healthcare/technical-information#metolose-sr-10398>

Exhibitions 2023

Please come visit us at the following trade shows. We will have technical teams available who will be happy to answer any questions, and we can also book private or confidential meetings.




4th European Conference on Pharmaceutics
Advanced technologies enabling new therapies
MARSEILLE FRANCE
20 - 21 March 2023
europeanmeeting.org

20th – 21st March 2023 | 4th European Conference on Pharmaceutics – Marseille, France

Talk
Dr. Vanessa Havenith
"Continuous Manufacturing: Rheological Powder Characterisation of Excipients to understand their Behavior during the Feeding Step"
Monday, 20.03.2023 at 16:00 at the Pharmaceutical manufacturing and engineering; Continuous manufacturing Session


Poster
Dr. Andreas Sauer
"Downstream Processing of Itraconazole: HPMCAS Amorphous Solid Dispersion: From Hot-Melt Extrudate to Tablet Using a Quality by Design Approach"
Tuesday, 21.03.2023



DCAT Week
The Premiere Event for the Global Bio/Pharmaceutical Manufacturing Value Chain
20th – 23rd March 2023 | DCAT – New York, USA

公益社団法人 日本薬剤学会
The Academy of Pharmaceutical Science and Technology, Japan
16th – 18th May 2023 | (APSTJ) – Japan
Pharmaceutical Application Laboratory
Science and Technology

14th Global DDMF Summit
Drug Delivery & Formulation
31st May – 2nd June 2023 | DDF – Berlin, Germany



in-PHARMA JAPAN
Int'l Pharmaceutical and Cosmetics Ingredients Expo
5th – 7th July 2023 | in-Pharma Japan – Tokyo, Japan

More Information

www.metoelose.jp/en www.setylose.com www.linkedin.com www.youtube.com

Contact

Japan
Shin-Etsu Chemical Co., Ltd.
Commercial:
Cellulose & Pharmaceutical Excipient Dept.
Address: 4-1, Marunouchi 1-chome,
Chiyoda-ku, Tokyo, 100-0005, Japan
Phone: +81-3-6812-2441
Technical:
Cellulose Technical Support Center
Address: YBP Technical center, 134, Godo-cho,
Hodogaya-ku, Yokohama, 240-0005, Japan
Phone: +81-45-459-5415
Web: www.metoelose.jp/en/

EMEA
SE Tylose GmbH & Co. KG
Address: Kasteler Strasse 45,
65203 Wiesbaden, Germany
Phone: +49-611-962-6345
E-mail: contact@setylose.com
Web: www.setylose.com

North America, Canada and Mexico
SE Tylose USA, Inc.
Address: 140 Commerce Way, Suite H, Totowa, NJ 07512, USA
E-mail: contact-pharma@setyloseusa.com
Customer Service
Phone: +1-225-309-0110 ext. 5714
Pharmaceutical Application Laboratory
Phone: +1-973-837-8001
Web: www.setylose.com

LATAM
Shin-Etsu do Brasil Representação de Produtos Químicos Ltda.
Address: Rua Coronel Oscar Porto, 736,
8^o Andar - Sala 84, Bairro Paraíso
CEP: 04003-003 - São Paulo, Brasil
E-mail: contact@setylose.com
Phone: +55-11-3939-0695
Fax: +55-11-3052-3904
www.setylose.com

India
Shin-Etsu Chemical Tylose India Pvt. Ltd.
Address: Office no. B, 7th Floor, D Building, MBC Park,
Ghodunder Road, Kasarwadavali
Thane West- 400615, India
E-mail: pharmaindia@setylosein.com
Phone: +91-22-6283-3008
Web: www.setylose.com