



Conference call and webcast for investors and analysts November 12<sup>th</sup>, 2025





#### **Forward-Looking Statements**

This presentation contains forward-looking statements, including, without limitation, statements regarding Nanoform's strategy, business plans and focus. The words may," "will," "could," "would," "should," "expect," "plan," "anticipate," "intend," believe," "estimate," "predict," "project," "potential," "continue," "target" and similar expressions are intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. Any forward-looking statements in this presentation are based on management's current expectations and beliefs and are subject to a number of risks, uncertainties and important factors that may cause actual events or results to differ materially from those expressed or implied by any forward-looking statements contained in this presentation, including, without limitation, any related to Nanoform's business, operations, clinical trials, supply chain, strategy, goals and anticipated timelines, competition from other companies, and other risks described in the Report of the Board of Directors and Financial Statements for the year ended December 31, 2024 as well as our other past disclosures. Nanoform cautions you not to place undue reliance on any forward-looking statements, which speak only as of the date they are made. Nanoform disclaims any obligation to publicly update or revise any such statements to reflect any change in expectations or in events, conditions or circumstances on which any such statements may be based, or that may affect the likelihood that actual results will differ from those set forth in the forwardlooking statements. Any forward-looking statements contained in this presentation represent Nanoform's views only as of the date hereof and should not be relied upon as representing its views as of any subsequent date.

# COMMERCIALLY LICENCED MANUFACTURING FACTORY







### Nanoform key strategy

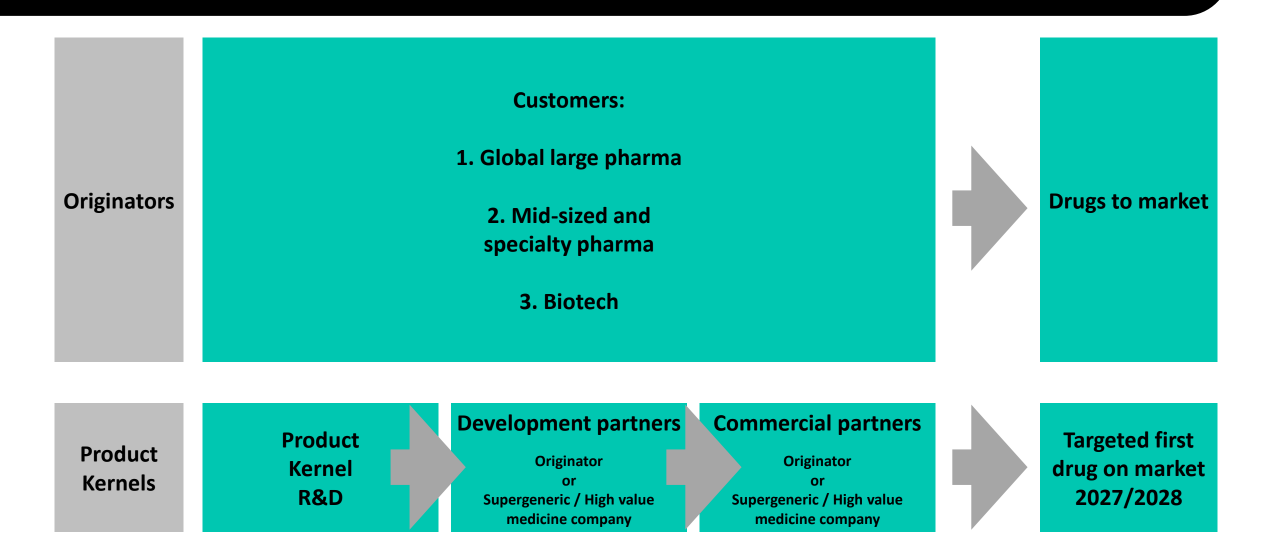
AII active pharmaceutical ingredients (API's) **should be Starmapped (AI)** 

Nanoform work with customers/partners to enable novel & existing molecules to become new and improved medicines

In parallel, to show a conservative industry the power of nanoforming, we create up to a dozen 'product kernels'



# Nanoform Technology – route to market





### **Proprietary technology platforms**

#### Small molecules

Proven CESS®\* nanotechnology enables new medicines through improved bioavailability, higher drug load & novel formulations

#### Large molecules

Unique **BIO** nanoparticles enable improved routes of administration with high drug load and longacting delivery

#### **Formulation**

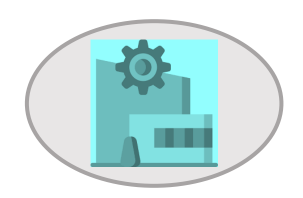
**Highly differentiated** novel formulations and unique drug delivery opportunities drive optimized therapeutic potential & patient convenience

STARMAP® 2.0 online picks best candidates and accelerates development by integrating deep expertise with sparse data Al



### Simplified value chain

#### High level overview of Nanoform's value chain and business model



#### **Clients**

- Global large pharma
- Mid-sized and specialty pharma
- **Biotech**



Launch of new drugs, improving existing drugs & reducing clinical attrition





#### Revenue

- Fixed fee per project
- Royalty as a % based on drug sales or supply price per kg

2012

2015

2020

2025

2028

**Discovery of** the CESS® process

**Nanoform** founded

**Clinical GMP** manufacturer license

European commercial cGMP manufacturing license

**Targeted first Nanoformed** medicine at market in Europe



### Nanoform key business highlights

- Nanoform granted European commercial cGMP manufacturing license
- Nancoencorafenib licensing and development agreement signed with A.forall (Riverside) and IMGA
- A new kernel, a nanoformulated combination of olaparib (Lynparza® by AstraZeneca Plc) and Ш temozolomide (Temodar® by Merck & Co Inc.) announced in partnership with Revio Therapeutics
- IV First target for 2026 announced: Cash burn below EUR 10m
- All 2025 near-term targets on track
- VI CMD date set for December, 16th, when new 2030 mid-term targets will be announced

## Nanoform granted European commercial cGMP manufacturing license

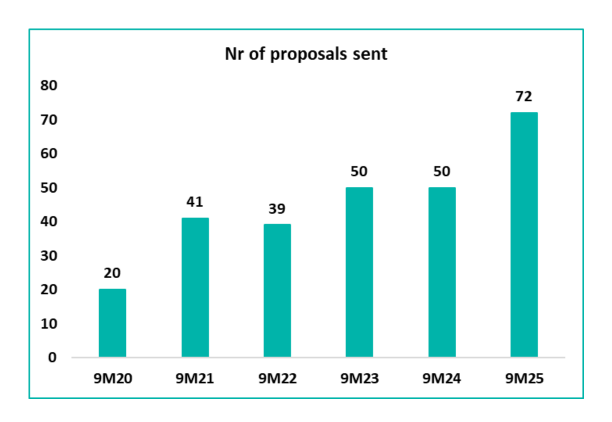
- **European commercial cGMP manufacturing license**
- For the production and quality control of nanoformed small molecule APIs (Active Pharmaceutical Ingredients)
- **Authorizes Nanoform to manufacture nanoformed APIs for** the European market and in countries in MENA, Asia, and Americas where mutual recognition applies to the European license

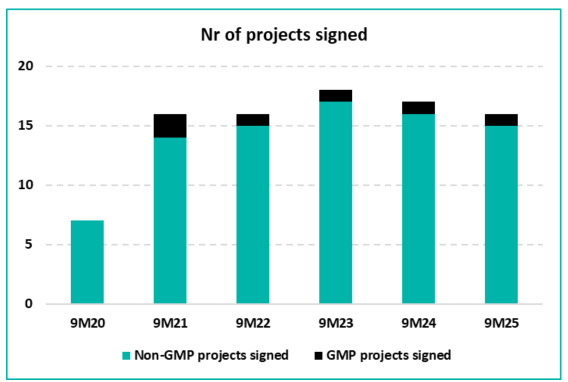






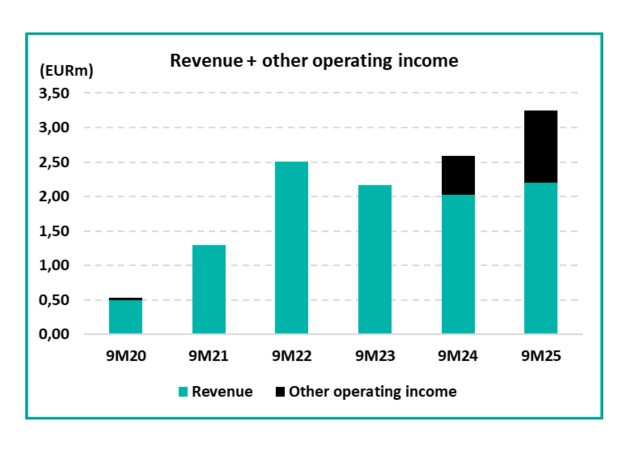
### Nr of proposals sent has jumped this year, projects signed stable despite difficult industry conditions for 4th year in a row

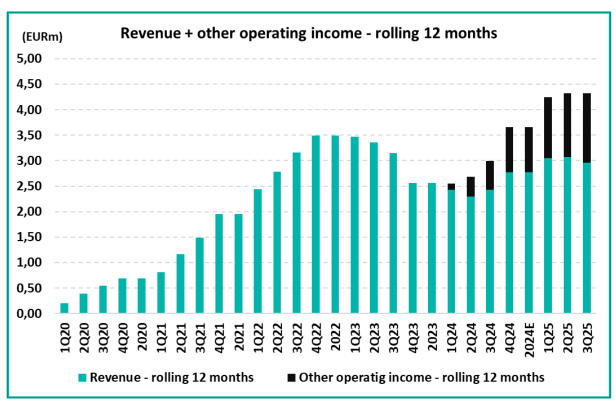






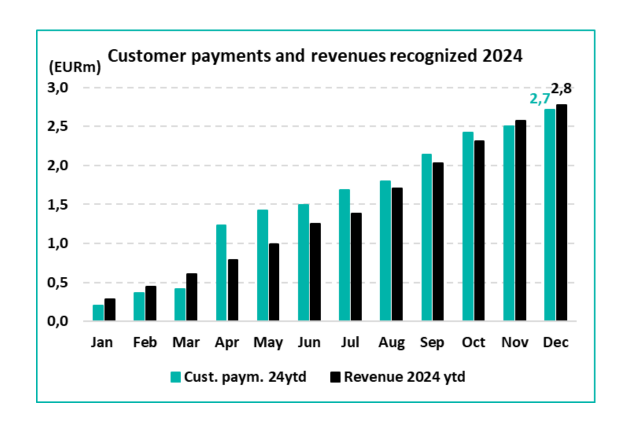
### Revenue and other income continues to grow

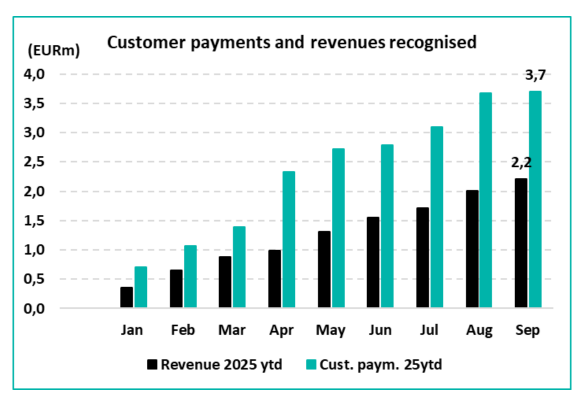






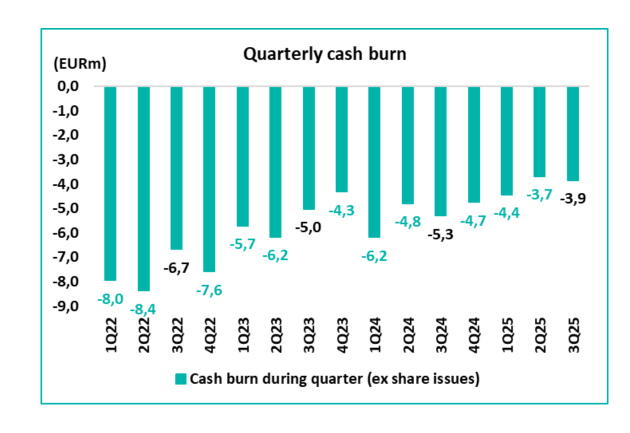
### Customer payments ytd exceed last year's payments and revenues recognized

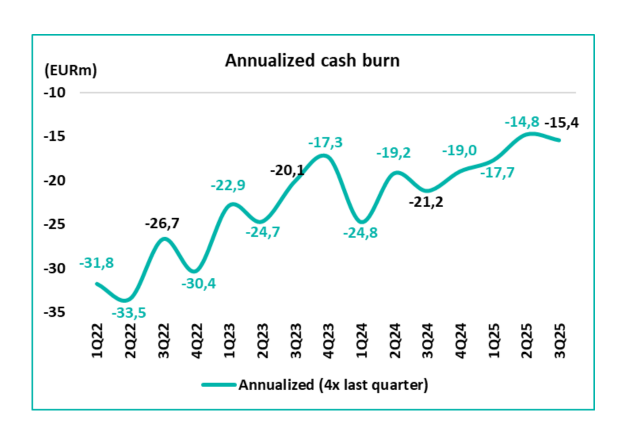






### Improvement in cash burn continues, target for 2026 < EUR 10m





- At the end of 3Q25, Nanoform had EUR 29.5m in cash
- Target for 2026: cash burn < EUR 10m

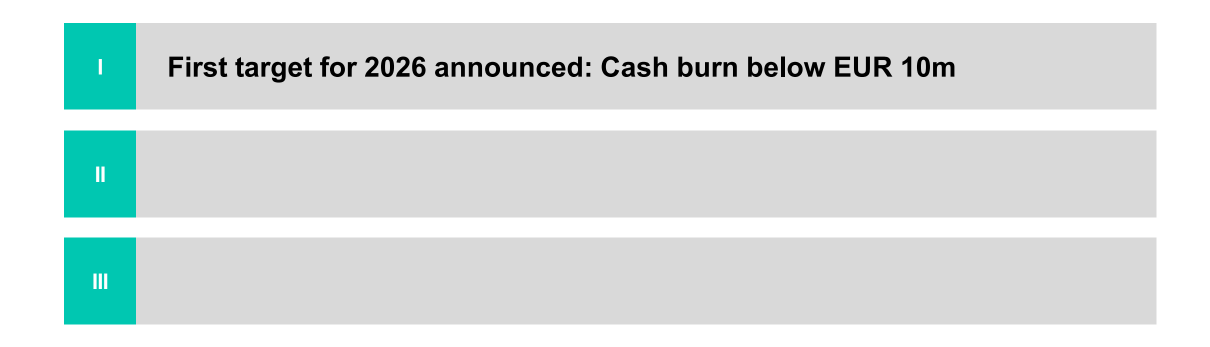


### Nanoform near-term business targets 2025 – all on track

- To sign several license/commercial supply agreements on several product kernels during 2025
  - First pivotal bioequivalence clinical study with a nanoformed medicine
- Increased number of non-GMP and GMP projects signed in 2025 vs 2024 Ш
- Improved free cash flow in 2025 vs 2024 IV



### Nanoform near-term business targets 2026









### Nanoform commercial highlights





### Nanoform partners with two specialist healthcare investors – October 14, 2025

- Nanoencorafenib is a patient-centric nanoformulation of encorafenib for melanoma and colorectal cancer
- Outlicensing, development and commercialisation agreement signed with A.forall (The Riverside) Company) and IMGA Futurum Tech Fund
- Encorafenib (Braftovi®, Pfizer) is an orally administered anti-cancer medication. Nanoform has developed a prototype nanoformulation with significantly higher drugload than that of the originator.
- The investment is expected to be sufficient to finance the clinical development of Nanoencorafenib up and until its commercialization and future outlicensing as an attractive patient-centric lifecycle management opportunity or a value-added generic medicine
- BRAFMed Ltd will pay Nanoform service fees, low single million development milestones, and up-to-midsingle digit tiered %-royalty. Nanoform's fully diluted ownership in BRAFMed is expected to be 40-50% (today 71%).



### Nanoform partners with Revio Therapeutics – October 27<sup>th</sup>, 2025

- Develop and commercialize GLIORA a locally-administered, long-acting, thermo-responsive hydrogel, for the treatment of high-grade glioma, a fast-growing and aggressive type of brain tumor
- GLIORA is a nano-formulated combination of olaparib (Lynparza® originally developed by AstraZeneca Plc) and temozolomide (Temodar® originally developed by Merck & Company Inc.)
- The program is expected to be in the clinic by 2H 2026. Subject to successful co-development and commercialization, GLIORA could be commercially available 2029-30
- Development costs and all licensing and commercial revenues will be shared equally between the partners, with Nanoform receiving an additional €1.5 million in accelerated revenue-share payments



### **Nanoform Product Kernel overview\***

			Nanoform Produ	Nanoform Product Kernels			Nanoform Pre-Clinical (non-GMP)			Nanoform Clinical (GMP)		Nanoform at Market		
Originator	Indication	Expected originator peak sales	Nanoformed API	Delivery route / dosage form	Nanoform ownership today	Development partnering status	Targeted commercial partnering	PoC*	Pre- formulation + in-vitro	Dosage form development + in vivo	PoP* / Dosage form development	Phase 1 / Pilot clinical trial	Pivotal - final - clinical trial	Earliest possible market launch
Astellas/ Pfizer	XTANDI®/Prostate cancer	~\$5bln	Nanoenzalutamide	Oral / Tablet	25 %	OnConcept Consortium	Ongoing							2027 US & 2028 EU
Johnson & Johnson	ERLEADA®/Prostate cancer	~\$5bln	Nanoapalutamide	Oral / Tablet	100 %	Ongoing	Ongoing					2026	2026-2027	2032 US & EU
Pfizer	BRAFTOVI®/Melanom a and colorectal cancer	~\$800mln	Nanoencorafenib	Oral / Tablet	71 %	BRAFMed Lda	Ongoing					2026	2027	2030 US & 2033 EU
Merck/AstraZ eneca	Glioma		NanoO+T (GLIORA)	Long Acting	50 %	Revio Therapeutics	2026-2027					2026	2028	2030 US & EU
Genentech/R oche	Oncology		Nanotrastuzumab	High Conc. Sub.Cut. Bio	100 %	2026	2026-2027							
Novo Nordisk	Obesity		Nanosemaglutide	Inhaled	100 %	2026	2027-2028							
Undisclosed	Inflammation		Undisclosed	Oral / Tablet	100 %	Partnered	2026-2027							
Undisclosed	Oncology		Undisclosed	Oral / Tablet	100 %	2026	2027-2028							
Undisclosed	Prostate cancer		Undisclosed	Long Acting	100 %	2026	2026-2027							

<sup>\*</sup> Only Product Kernel pipeline, i.e. not including customer projects

<sup>\*</sup> PoC = Proof of Concept

<sup>\*</sup> PoP = Proof of Process



### Industry shift and momentum in biologics delivery

#### **Recent industry news:**

Halozyme acquires Elektrofi for up to \$900m, including \$750m upfront (subject to regulatory approval)

- The transaction sends a strong signal that the value of advanced biologic formulation platforms is recognized by major players
- Nanoform's biological platform was designed with scalability, robustness, and patient-centric administration in mind
- Nanoform's biologics technology is complementary to the direction the market is moving - allowing for high concentration formulations, low injection volumes, and broad usability outside the clinic - for example enabling to switch from intravenous delivery to subcutaneous delivery

Nanoform high drug load suspension in a prefilled syringe (400-500mg/ml)





### **News from the pharma market**







'Significant interest in high concentration biologics from major pharma and leading Biotechs – Nanoform seen as a positive alternative'

#### PODD 2025, Boston



'Major CDMO peers confirm low project numbers in small molecule early development outsourcing'

'Biosimilar regulatory changes mean greater interest in this space from developers – faster and cheaper to market'

CPhI 2025, Frankfurt



BioEurope 2025, Vienna



### Nanoform expands into Asia

#### Japan

- **April 2024**
- **Exclusive partnership with CBC to bring best-in**class nanomedicine technology to Japan
- Shinya Miyairi, Managing Executive Officer at CBC: "CBC is honored to partner with Nanoform and to represent them in Japan. We believe that Nanoform's technologies represent a great fit for the Japanese market where the ability to provide medicines with higher bioavailability and fewer and smaller doses are important for success."



#### **South Korea**

- September 2025
- Nanoform expands commercial presence in Asia with A&LS Pharma in South Korea
- Mr. Won-Mook Kim, President of **A&LS Pharma:** "Nanoform's unique nanoparticle engineering services perfectly complement our mission to bring advanced solutions to Korean pharmaceutical and biotech clients. We look forward to helping our customers unlock the full potential of their molecules and deliver better medicines to patients"



### **Nanoform Capital Markets Day (CMD)**

#### December 16<sup>th</sup>, 2025, 09.00-13.00 EEST

@ Nanoform headquarter and commercial cGMP manufacturing site in Helsinki

Nanoform's leadership presents Nanoform's key priorities for its next strategy period together with new mid-term targets for 2030

Invitation to CMD will be press released Nov 13th, 2025





Edward Hæggström CEO



Johanna Kause Chief Quality Officer



Albert Hæggström CFO



Christian Jones



Peter Hänninen
General Counsel &
Chief Development
Officer





### Selection of upcoming events

Nanoform Q3 2025 report November 12

**SEB Healthcare Seminar, Stockholm** November 13

Jefferies Global Healthcare Conference, London November 18-20

November 25 Aktiespararna - Stora Aktiedagarna, Stockholm

DDL, Edinburgh December 10-12

December 16 Nanoform Capital Markets Day, Helsinki

JP Morgan 44<sup>th</sup> Annual Healthcare Conference, San Francisco **January 12-15** 

Nanoform Q4 and FY 2025 report February 2026

March 23-26 DCAT, New York



#### **Interesting short videos:**

Drug Delivery Leader Chief Editor Tom von Gunden sits down with Christian Jones, FRSC, Chief Commercial Officer at Nanoform, to discuss how nanoparticle technology is driving innovation in drug and biologics delivery:

<a href="https://www.drugdeliveryleader.com/doc/leveraging-nanoparticles-for-high-drug-load-delivery-with-nanoform-s-christian-jones-0001">https://www.drugdeliveryleader.com/doc/leveraging-nanoparticles-for-high-drug-load-delivery-with-nanoform-s-christian-jones-0001</a>

Nanoforming Biologics & GLP-1's: From IV to SUBQ and inhaled delivery <a href="https://event.on24.com/wcc/r/5024500/A18B95A0EFCAB2A1AFE07E1EB66A7233">https://event.on24.com/wcc/r/5024500/A18B95A0EFCAB2A1AFE07E1EB66A7233</a>

Nanoform high dose subcutaneous delivery of biologics: <a href="https://nanoform.com/en/nanoform-high-dose-subcutaneous-delivery-of-biologics/">https://nanoform.com/en/nanoform-high-dose-subcutaneous-delivery-of-biologics/</a>

Discover how Nanoformed API outperform traditional solid dispersions: <a href="https://nanoform.com/en/nanoform-cphi-milan-2024-tamas-solymosi/">https://nanoform.com/en/nanoform-cphi-milan-2024-tamas-solymosi/</a>

Nanoform's best-in-class nanodevelopment capabilities: <a href="https://nanoform.com/en/nanoform-development-capabilities/">https://nanoform.com/en/nanoform-development-capabilities/</a>

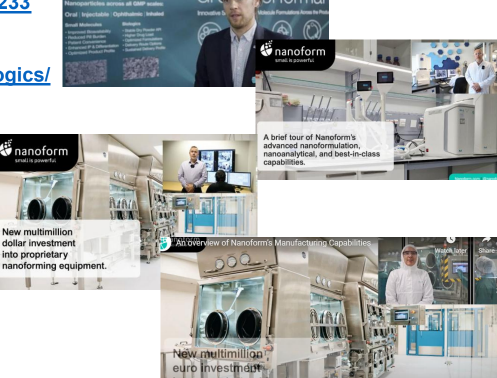
Nanoform's advanced nanoformulation, nanoanalytical, and best-in-class capabilities:

https://nanoform.com/en/nanoform-formulation-and-analytical-tour/

Nanoform's state-of-the-art manufacturing capabilities: https://nanoform.com/en/nanoform-dr-david-rowe-manufacturing-with-drone/



Leveraging Nanoparticles
For High Drug Load Delivery
With Nanoform's
Christian Jones



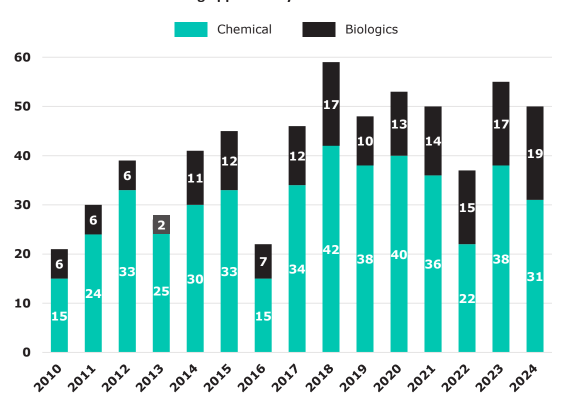


### The structural pharma R&D problem in the pharma industry

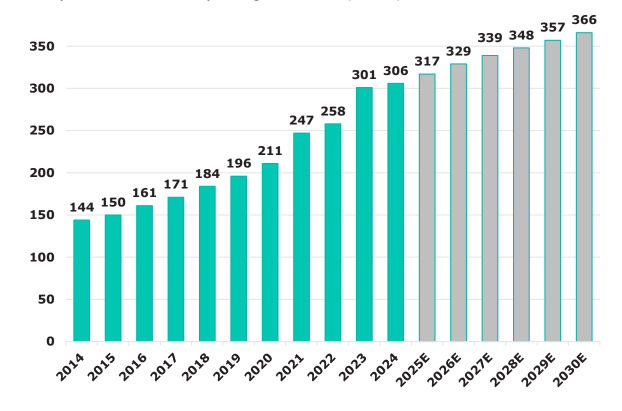
Fewer than 50 drugs approved in the US annually on average...

...while the global pharma industry R&D expenditure exceeds \$300B

Annual number of novel drug approvals by FDA 2010-2024



Global pharmaceutical R&D spending 2014-2030E (USDbn)



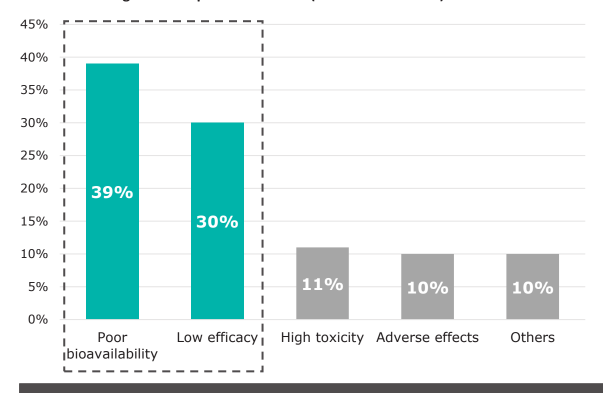
A game changer is needed to improve R&D yield



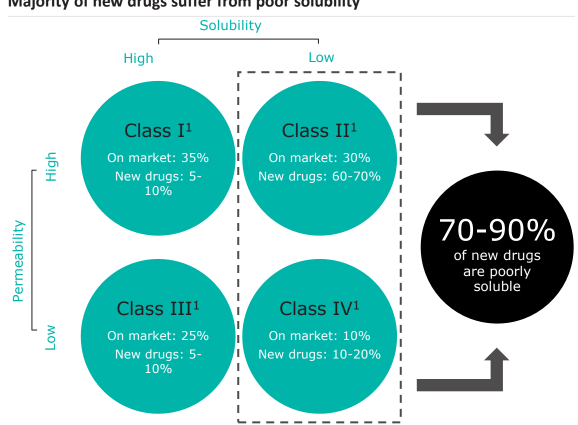
### Low bioavailability is the key issue

#### Poor bioavailability and low efficacy most common reasons for drug failure

#### Reasons for drug failure in pre-clinical trials (share of molecules)



#### Majority of new drugs suffer from poor solubility



#### > Nanoform can enhance the pharma industry output by targeting poorly soluble drugs

Source: GlobalData 2009, Cutting Edge Water-based Nanotechnology in Drug Development (Reasons for drug failure); Nikolakakis & Partheniadis (2017), Self-Emulsifying Granules and Pellets: Composition and Formation Mechanisms for Instant or Controlled Release (Share of poorly soluble drugs) 1) Classification of drug substance according to Biopharmaceutics Classification System (BCS)



### **Revenue drivers & industry attrition rates**

#### Nanoform pre-clinical and clinical revenue drivers

Non-GMP

#### Proof of Concept (PoC)

- # of active customers
- > # of APIs per customer
- Price per PoC per API

#### **Proof of Process** (PoP)

- Attrition between PoC and PoP
- Price per PoP per API
- > Time lag between PoC and PoP

#### **GMP**

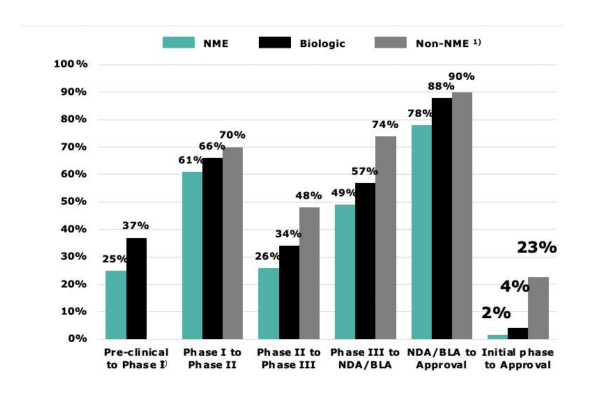
#### Phase I II & III and/or 505(b)(2)

- Attrition between previous and current phase
- Price per phase per API
- > Time lag between previous and current phase
- # of customers with 505(b)(2) strategy
- Proportion of new drug candidates and 505(b)(2) **APIs**

#### Drugs on the market

- # of drugs on the market using CESS®
- License fee & royalty level per drug
- Net revenues per drug
- Time lag Phase II and market (505b2)
- Time lag Phase III and market
- Speed of uptake on market

#### Global Pharmaceutical industry's pre-clinical and clinical success rates

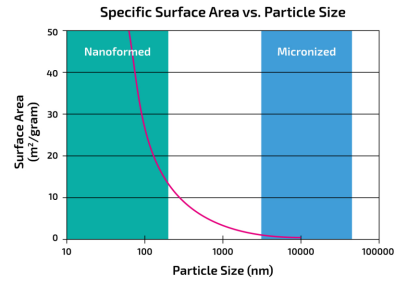


Timeline (years)	Pre-clinical	Phase I	Phase II	Phase III	Approval	Total
New drugs	~1-4	~2	~2	~3-4	~1	~9-13
Existing drugs	-	~1	~3-6			

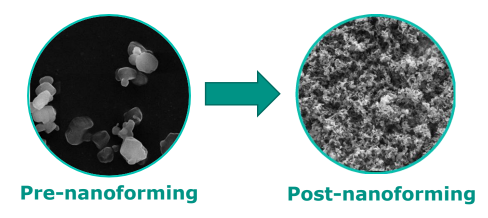


#### Particle size is key

#### Smaller particle size can improve a drug's bioavailability



- The surface area increases 30-fold from a 10 micron sized particle once the particle size is reduced to 100nm
- Reduction of particle size down to 50nm increases the surface area by 1,000-fold



- Smaller particles have a larger surface area
- Larger surface area of particles enables improved bioavailability of a drug
- Improved bioavailability implies increased absorption of a drug by the body's circular system
- CESS® can produce API with large surface areas which can significantly improve the bioavailability of drugs

> CESS® produced nanoparticles have a larger surface area and as such improved bioavailability.

Source: Company information 1 micron = 1,000nm

### Nanoform is here to fill the gap

**Enabling** new drugs

> 20,000 drugs in development\* **Improving** existing drugs

> 5,800 existing drugs\*

Giving unsuccessful drug candidates a second chance

> 58,000 failed drugs in the last 40 years\*



# Small molecules - Small is powerful®





## Nanoform has made substantial progress in Nanoforming solutions with in-vitro, in-vivo, and clinical study results

**Oncology:** Replaced amorphous solid dispersion (ASD) formulations with nanocrystalline high drug load formulations,

> matching bioequivalence for Enzalutamide and Apalutamide where life cycle management opportunities to reduce tablet burden to a single, smaller, easier-to-swallow tablet as well as working on Aprepitant in partnership with PlusVitech for

lung cancer to develop a regimen with substantially fewer tablets.

Inhalation: Engineering nanoformulations of both small and large molecules with excellent fine-particle dose (FPD) and fine-

particle fraction (FPF) performance in comparison to spray drying technologies. In biologics, Nanoform has shown FPF

>95% vs 50% with spray drying for delivering **high drug load** to the lungs.

**Biologics:** Demonstrated in partnership, with Takeda and other companies, ultra-high concentrations for subcutaneous drug

**delivery** with acceptable viscosity for injection (Takeda – Plasma Derived Therapies).

**Ophthalmic:** Multiple projects where nanoparticles have shown improved delivery potential. High drug load to the eye enabling

smaller implants with no requirement for mesh membranes, eye drop suspensions and ophthalmic inserts.

**Hydrogels: Shown high drug load** applications (5 x more than nanomilling) for post-surgical glioblastoma drug delivery and deep

penetration across the brain parenchyma enabling non-recurrence of glioblastoma where other formulations failed.

IP: Novel technologies, processes and formulations can enable market opportunities, lifecycle management and strong

launch strategies



# Nanoform customer projects – therapy area overview\*

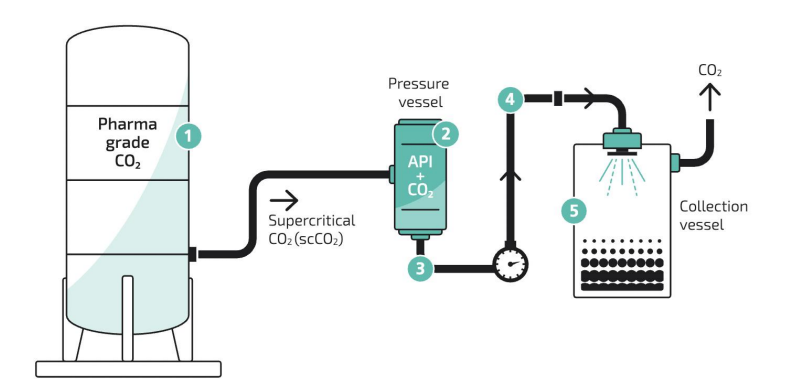
Pre-Clinical	Phase I	Phase II & III	Marketed/505b2
Cardiology	Immunology/Inflammation	Metabolism and Endocrinology	Infectious Disease
(e.g. Anemia)	(e.g. Cystic Fibrosis)	(e.g. Adrenal Hyperplasia)	(e.g. HIV)
Gastroenterology	Dermatology/Oncology	Neurology	Immunology/Inflammation
(e.g. Microbiome)	(e.g. Basal Cell Carcinoma)	(e.g. Schizophrenia)	(e.g. HEP B)
Immunology/Inflammation	Neurology	Oncology	Immunology/Inflammation )
(e.g. Psoriasis)	(e.g. Parkinsons)	(e.g. lung cancer)	(e.g. Cystic Fibrosis)
Infectious Disease	Oncology		Oncology
(e.g. HIV)	(e.g. Solid Tumors)		(e.g. Prostate Cancer)
Metabolism and Endocrinology	Ophthamology		Ophthamology
(e.g. Diabetes)	(e.g. Cataract)		(e.g. Glaucoma)
Neurology	Pain		
(e.g. Parkinsons)	(e.g. Post Operative Pain)		
Oncology	Infectious Disease		
(e.g. Multiple Myeloma)	(e.g. HIV)		
Ophthamology			
(e.g. Glaucoma)			
Respiratory			
(e.g. COPD)			

<sup>\*</sup> Shows the stage of customer molecule, not in which phase the project is at Nanoform (non-GMP, GMP, at market)

## **Small Molecules - Proprietary technology**

Green technology

### **Controlled Expansion of Supercritical Solutions - CESS®**



- Supercritical CO₂ is guided into a pressure vessel loaded with API
- Increasing the pressure and temperature in the vessel dissolves the API in supercritical CO<sub>2</sub>
- The CO<sub>2</sub> and the API are released from the pressure vessel and the flow, pressure and temperature profiles are accurately controlled
- The pressure and temperature is controlled to achieve a stable nucleation phase and formation of nanoparticles
- In a collection vessel the CO<sub>2</sub> is sublimated resulting in final nanoparticles ready for collection and formulation

> Relatively simple process developed through combining deep knowledge in physics, chemistry, and pharma



# **CESS® Superior to Existing Technologies**

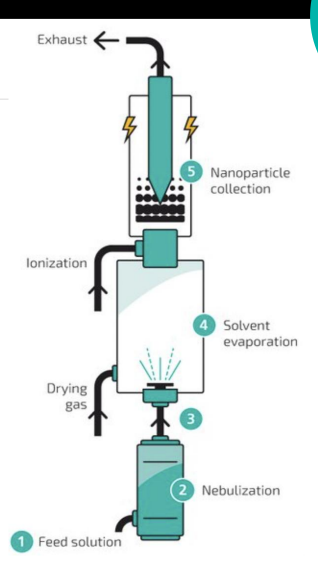
	Controlled Expansion of Supercritical Solutions (CESS <sup>®</sup> )	Solid dispersion (e.g. spray drying)	Jet milling	Nanomilling
Description	Extracts API from supercritical CO <sub>2</sub> by applying controlled reduction in pressure	API is dispersed into a solid material, which dissolves when exposed to an aqueous media	Application of energy to physically break down API particles to finer ones	API particle size is reduced in a liquid vehicle via grinding
Particle size	Down to 10nm	300nm-25μm	800nm-10μm	>150nm
Particle formation	Controlled crystalline or amorphous and stable	Amorphous (unstable without excipients)	Unstable (crystalline and amorphous structures)	Unstable (crystalline and amorphous – needs excipient to stabilise)
Ease of formulation	✓	×	×	×
Reproducibility	✓	✓	*	×
Free from excipients and solvents	✓	×	✓	×
Yield	High	Low	High	Low
Investment	Low	High	Low	Low

## **Large molecules - Proprietary technology**

**Green** technology

### **Nanoforming process for biologics**

- API containing feed solution is pumped into the nebulizer
- Peed solution is nebulized into a carrier gas
- Mist is transported into the drying chamber via a connection pipe
- Mist is dried using low-temperature drying gas
- Dried particles are charged by the ionizer and collected using electrostatic precipitation

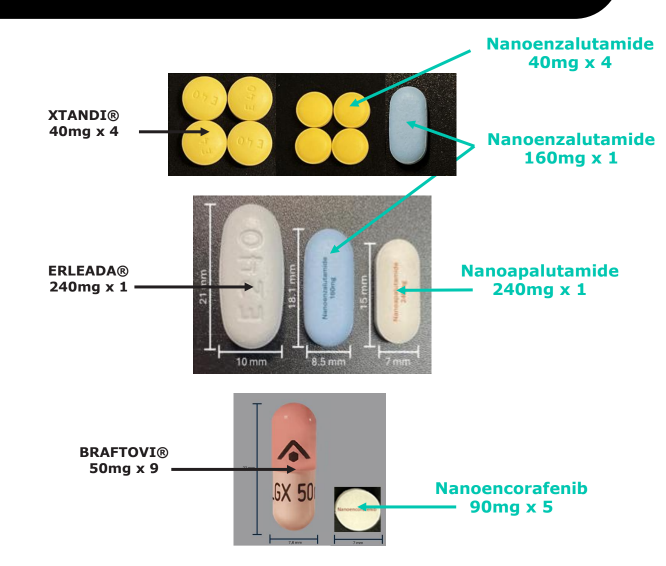


API = active pharmaceutical ingredient Nebulization = turns liquid into mist Ionization = particles electrically charged



### **Small molecules – Nanoform enables small/single pill strategy**

	Existing drug	Nanoformed version
	XTANDI®	Nanoenzalutamide
Formulation	ASD	Crystalline Nanoparticle
Drug load 160mg (x1)	-	40 %
Drug load 40mg (x4)	12 %	40 %
Size 160mg (x1)	-	18.1 x 8.6 mm
Size 40mg (x4)	10.1 mm	8.0 mm
	<b>ERLEADA®</b>	Nanoapaluatmide
Formulation	ASD	Crystalline Nanoparticle
Drug load 240mg (x1)	21 %	42 %
Drug load 60mg (x4)	7 %	42 %
Size 240mg (x1)	21 x 10 mm	15 x 7 mm
Size 60mg (x4)	17 x 9 mm	8 mm
	<b>BRAFTOVI</b> ®	Nanoencorafenib
Formulation	ASD	Crystalline Nanoparticle
Drug load 90mg (x5)	-	
Drug load 75mg (x6)	12 %	
Drug load 50mg (x9)	12 %	
Drug load 45mg (x10)	-	
Size 90mg (x5)	-	
Size 75mg (x6)	23 x 8.5 mm	
Size 50mg (x9)	22 x 7.6 mm	
Size 45mg (x10)	-	





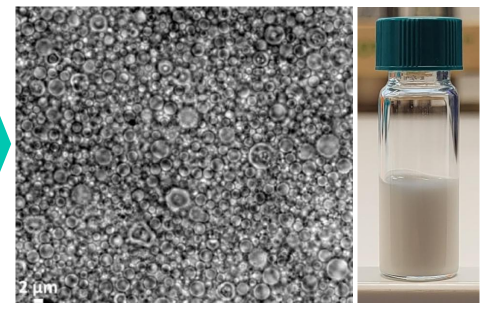
## Biologics - Game changing high drug load subcutaneous delivery (400-500 mg/ml)

### Nanoformed monoclonal antibody in dry powder

WD = 5.6 mm Signal A = InLens

Nanoformed high drug load monoclonal antibody in non-aqueous suspension

High drug load suspension in a prefilled syringe (400-500mg/ml)





- Non-aqueous suspension enables high protein load in a low volume (400-500 mg/ml)
- Intact and stable protein particles in suspension
- Good injectability of suspension with injection force below 20 N using a 27G needle

# **Nanoform Product Kernels**

Nanoform internal Product Kernel work	Development partners	Commercial partners
Value proposition around a medicine candidate, called 'Product Kernel'	Originator or Supergeneric / High value medicine company	Originator or Supergeneric / High value medicine company
2. New IP that Nanoform owns in an R&D phase	1. Upfront payments 2. Milestones 3. Revenue from Nanoforming the medicine for clinical trials	<ol> <li>Upfront payments</li> <li>Milestones</li> <li>Revenue from Nanoforming the medicine for clinical trials and commercial phase</li> <li>Royalties/profit share</li> </ol>



# Attractive revenue model with pharma and biotech customers

Phase	Proof of Concept / Proof of Process	Phase I – III clinical trials	Drugs on the market
Certification	Non-GMP	GMP	GMP
Description	<ul> <li>Proof of concept study - assessment of the possibility to nanoform a specific API</li> <li>Proof of process study - definition of parameters to establish the optimal process and controls for a specific API</li> </ul>	<ul> <li>API for clinical trials are manufactured in Nanoforms GMP facility</li> <li>Supply of material for customers' Phase I, II and III trials</li> </ul>	<ul> <li>Drugs that have passed the trials and reached commercialization</li> <li>Significant potential from patent extension (505b2 projects) of drugs already on market</li> </ul>
Revenue model	Fixed fee per project  Estimated project fee of EUR 50-500k  per API per project	Fixed fee per project  Estimated project fee of EUR 0.5-10m  per API per phase	Royalty as a % on drug sales or supply price per kg  Estimated royalty fee of 1-20%



## **Business case Amorphous Solid Dispersions (ASDs)**

Amorphous solid dispersion (ASD) medicines are currently the leading formulation strategy for poorly soluble APIs and there are ~50 marketed medicines globally that are ASDs and sell for ~\$50bln annually

Nanoformed and nanocrystalline medicines (e.g. nanoenzalutamide etc) offer an attractive alternative to ASD medicines (and other) with the following benefits to originators and supergeneric/high value medicines companies:

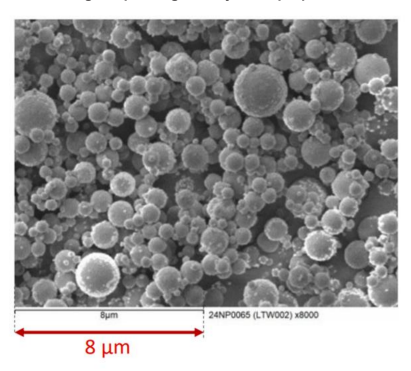
- green manufacturing process
- substantially <u>higher drug load</u> in the final drug product
- reduced pill burden for the patient
- opportunity to <u>extend IP protection</u> for the reformulated and improved product
- opportunity for <u>earlier market entry</u>
- possibility for <u>fixed dose combinations</u>



### Comparison of Nanoform's proprietary biologics technology vs existing technologies

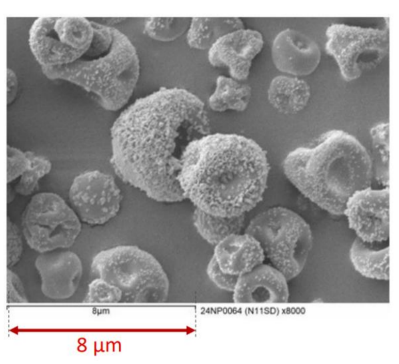
### **Nanoformed**

Perfect spheres, highly flowable and aerodynamic, great packing and injection properties



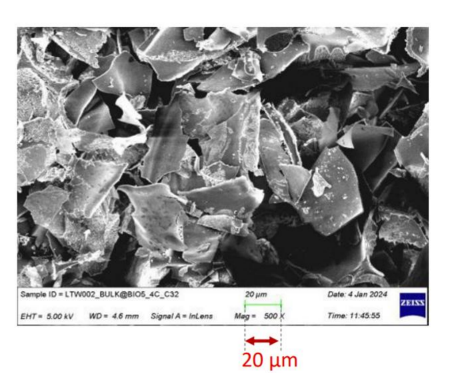
**Spray dried** 

Sticky, poor flowability, raisin shaped



### Lyophilized / freeze dried

Flaky morphology, dry cake, no flowability



Nanoforming biologics: Superior flowability, aerodynamic performance, high density packing, lower injection force properties, improved material quality and stability properties vs spray drying and lyophilization



### Nanotrastuzumab press release June 3, 2025

### Trastuzumab (Herceptin®)

- Genentech/Roche
- Monoclonal antibody (mAb)
- Breast and stomach cancer
- Intravenous administration
- In 2019, a hyaluronidaseenabled subcutaneous (SubQ) formulation (Herceptin HYLECTA™) of the product was approved, using Halozyme's ENHANZE® drug delivery technology¹
  - 1) Herceptin® is administered with 600mg every week intravenously (173min at oncology unit) Herceptin® HYLECTA™ is administered with 600mg subcutaneously every three weeks (53min at oncology unit)

### Nanotrastuzumab

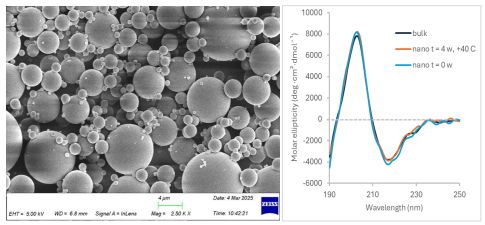
- Nanoform Finland Plc
- A high concentration (HC)
   nanoformulation of trastuzumab
- Proof-of-concept pre-clinical study
- Suitable for subcutaneous (subQ) injection/administration
- Above 400 mg/mL dosing
- Hyaluronidase-free
- Full dose in a single 2mL syringe

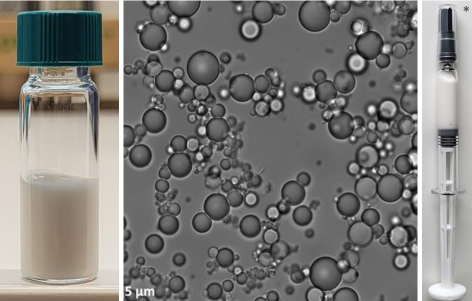
### Nanoform HC-SubQ benefits

- Potential to transform the industry of administration of biologic medicines
- Nanoform's formulation platform enables high concentration subcutaneous (SubQ) administration
- Replacing intravenous administration
- Significantly reduced healthcare costs, better patient experience and quality of life
- Potentially complementing Halozyme's technology



## Nanoformed Trastuzumab SubQ Suspension Formulation





From dry particles to 440 mg/ml stable Trastuzumab in suspension \*Svringe reference: Courtesy of Stevanato Group

- No significant changes can be detected in Trastuzumab primary or higher order structures and nanoformed mAb remains fully functional
- Up to 650 mg/ml of dry mass was reached with an injection force of below 10N
- Particles remain intact in the suspension after 4 weeks at +4°C and +25°C
- No sedimentation detected by Turbiscan technology at +4/+25C for 4 weeks
- **Injection force remain stable** after storing suspension in syringes at +4/+25C for 4 weeks

### June 2024

Nanoformed highconcentration biologics formulation for subcutaneous delivery results presented by Takeda at Drug **Delivery and Formulation Sumit** in Berlin

### August 2024

Nanoform and Takeda initiates collaboration on Takeda's plasmaderived therapy development

### June 2025

**Positive Nanoform** biologics respiratory data presented by Takeda at Drug **Delivery and Formulations Summit** in Berlin

## Sep & Oct 2025

Takeda to present on both high concentration subcutaneous data and respiratory data with Nanoformed Plasma Derived Therapies in **September at DDF Boston and in October** at PODD Boston



## Project Glioblastoma (hydrogel for central nervous system cancer)

Nanoform customer TargTex S.A. was granted Orphan Drug Designation by FDA for its nanoformed drug candidate TTX101 to be used in patients with malignant gliomas (October 2023). The orphan drug designation follows the generation of a preclinical rodent data package in which a survival advantage was shown for this nanoform-enabled medicine candidate.

The hydrogel nanoformulation developed by Nanoform enabled a 200-fold increase in drug load compared to bulk and a 5-fold increase in drug load compared to nanomilling.

In November 2023, the European Innovation Council and SMEs Executive Agency (EISMEA) awarded TargTex €14m in funding.

TargTex is currently raising additional funds to take this innovative treatment to clinic and is planning a phase 1/2a clinical trial in recurrent glioblastoma (GBM) patients across the US and EU, in which nanoformed TTX101 is applied as adjunct to surgery after tumour excision.



### Management team: Multi-disciplinary with international merits



#### CEO & Co-founder; Ph.D. (Applied physics), MBA Edward Hæggström



- Professor at the University of Helsinki, Head of Electronics Research Lab. within the Dept. of Physics
- Previously visiting professor at Harvard Medical School, visiting scholar at Stanford University and project leader at CERN
- Has led large number of scientific projects
- Current ownership: 5,409,405 shares and 408,000 options



#### CCO; M.Sc. (Chemistry) **Christian Jones**



- · Previously Commercial Director and member of the Senior Leadership
- Team for the Global Health Sector at Johnson Matthey
- Senior roles at Dr. Reddy's Global Custom Pharma Solutions and Prosonix
- **Key area of responsibility:** Commercial strategy and business development
- Current ownership: 284,000 options



#### General Counsel & Chief Development Officer; LL.M Peter Hänninen



- Previously Attorney, Borenius Attorneys
- Successful track-record of advising technology companies from founding to exit in key transactions and collaborations
- Key area of Responsibility: Legal, Compliance, IPR, HR, IT
- Current ownership: 173,125 shares and 580,000 options



Chief Quality Officer, M.Sc. (Pharmacology) Johanna Kause



- · Previously Head of Quality, Regulatory and Safety for Finland and the Baltics at Takeda Pharmaceuticals
- · 25 years of experience in Quality Management in the Pharma
- **Key area of responsibility:** Quality Management, GMP, GDP
- Current ownership: 130,000 options



#### CFO and member of the Board; B.Sc. (Economics) Albert Hæggström



- 20 years of finance and investing experience
- Prior roles include positions at Alfred Berg, BNP Paribas, Nordea and
- Current ownership: 805,779 shares and 690,000 options



Head of Manufacturing; Ph.D. (Chemistry) **David Rowe** 



- Previously Particle Size Reduction Lead for GlaxoSmithKline
- Chaired the PSR Centre of Excellence
- **Key area of responsibility:** Technical leadership within new chemical entities and commercial assets
- Current ownership: 313,720 options



#### **Chief of Business Operations (Chemistry and Quality)** Antonio da Silva



- Degree in Chemistry from Lisbon University and Master degree in Quality from the University Aberta of Lisbon
- Extensive background in the CDMO and particle engineering space (19 vears at Hovione)
- **Key area of responsinility:** Pharmaceutical product launches
- Current ownership: 25,051 shares and 228,032 options



## **Board of directors: Top executives from leading industry positions**



### **Miguel Calado** Chairman of the Board





- Previously CFO at international particle engineering CDMO company Hovione Group
- Other previous roles include CFO at PepsiCo International and President International Operations at Dean Foods
- Experienced Board member in both the EU and the US
- Current ownership: 167,544 shares and 230,000 options
- Key experience:



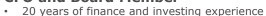






### **Albert Hæggström**





- Prior roles include positions at Alfred Berg, BNP Paribas, Nordea and SEB
- Current ownership: 805,779 shares and 690,000 options
- Key experience:





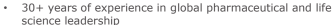






### Jeanne Thoma

#### **Board Member**



- Prior roles include executive positions at BASF Inc, Lonza AG and SPI Pharmaceuticals
- Current ownership: 91,263 shares and 38,630 options
- Key experience:

Lonza We create chemistry







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