# Scanning analysis of semi-solvent impact using sequential solvent vapor (SASSI-SSV): Assay of poly(lactide-co-glycolide)-naltrexone (PLGA-NTX) microparticles

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### Introduction

The microstructural arrangement of PLGA microparticles is a critical facet of their drug release performance. Combining the selective solubility of semi-solvents with advanced imaging techniques allows analysis of PLGA microparticle structural arrangement based on morphological changes to the particles in response to semisolvent exposure [1]. It was found that a series of semi-solvent vapors can be used to dissolve PLGA layers from the surface for rapid, automated tracking of particle morphological (related to the microparticle structural) changes.

**Emulsion-**

Figure 2. Exa	ample images	s in laser-inte	nsity mode o	f selected mi	croparticles.	
(Lactide% by	HNMR and	M <sub>n</sub> by Gel Pe	ermeation Ch	romatograph	l <b>y</b>	
quaternary detection, Red scale bar = $50 \ \mu m$ ).						
Sample	Dry	EI	TOL	2PE	PA	
$(L\%, M_{n})$						
Emulsion						
Blank -no			<b>.</b>			
drug (75%,						
33,941 Da)						
	50.000µm	50.000µm	50.000µm	50.000μm	50.000µm	

Results

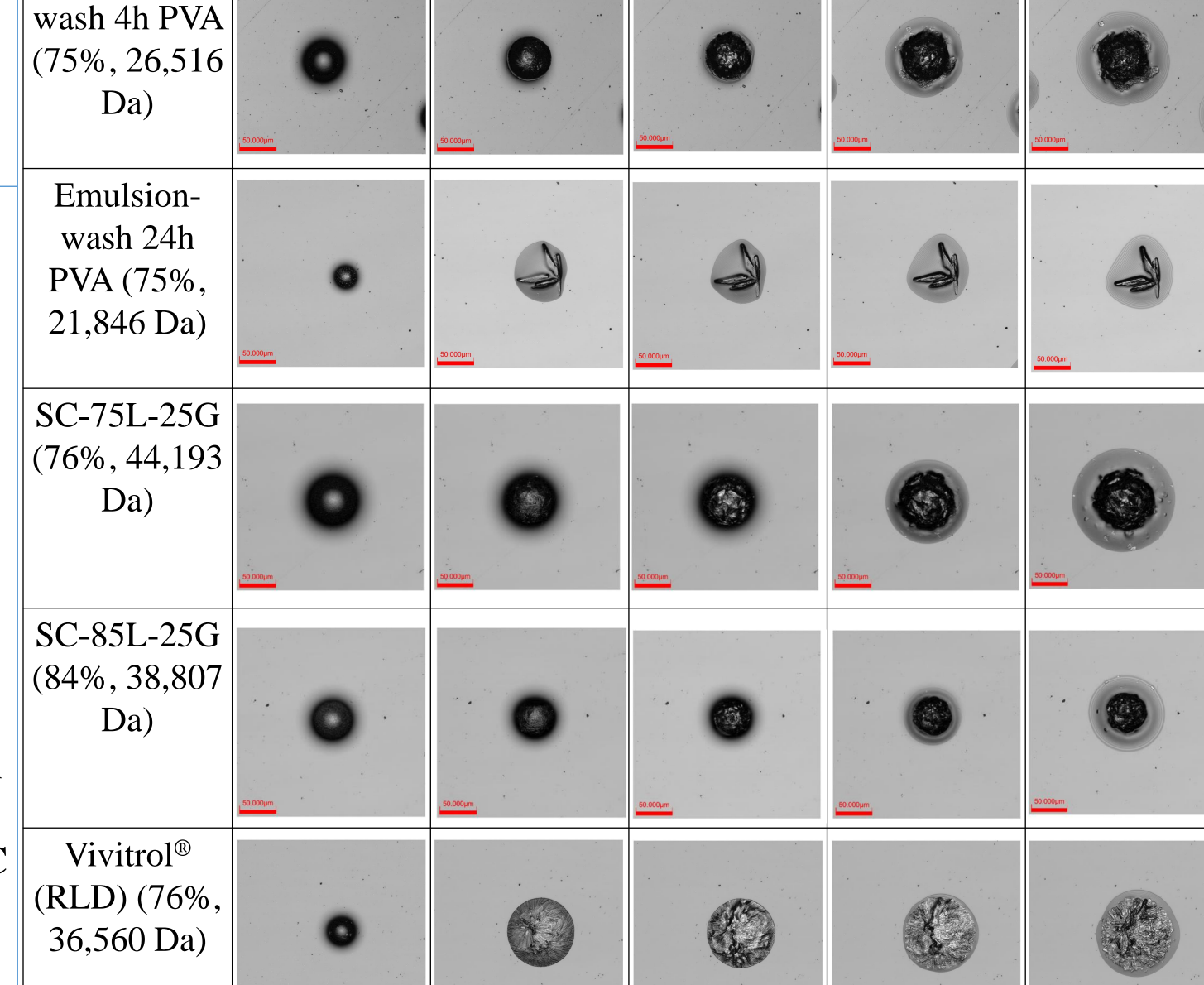
Table	<b>1.</b> Se	elected	paran	neters	from	samples	(average <u>+</u>
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standard deviation, N indicated for each sample)

PLGA (75L) blank (N = 29)								
Condition	Sq	Sk	Sku	S10Z	Ar			
Dry	$5.77 \pm 3.9$	$3.09 \pm 2.3$	$17.17 \pm 60.5$	$15.66 \pm 16.6$	$5.1\pm0.88$			
EI	$3.61 \pm 2.6$	$4.45 \pm 3.7$	$5.82 \pm 5.3$	$12.04\pm7.9$	$4.4 \pm 0.41$			
TOL	$0.72 \pm 0.5$	$0.56\pm0.5$	$9.80 \pm 5.9$	$6.77 \pm 3.7$	$1.4 \pm 0.14$			
2PE	$0.16 \pm 0.1$	$0.11 \pm 0.1$	$21.18\pm54.6$	$2.34 \pm 1.3$	$1.0 \pm 0.01$			
PA	$0.13 \pm 0.1$	$0.11 \pm 0.1$	$12.94 \pm 18.5$	$2.45 \pm 1.6$	$1.0 \pm 0.01$			
Emulsion-wash (4H PVA/8H ETOH) (N = 30)								
Dry	$4.17 \pm 1.1$	$4.80 \pm 2.7$	$2.37\pm0.9$	$13.41 \pm 8.6$	$5.1 \pm 0.41$			
EI	$4.22 \pm 1.2$	$4.25 \pm 1.4$	$3.61 \pm 3.3$	$27.30\pm10.8$	$4.6\pm0.74$			
TOL	$5.66 \pm 1.6$	$9.32 \pm 4.3$	$2.96 \pm 1.5$	$30.29\pm7.4$	$4.1 \pm 1.06$			
2PE	$4.48 \pm 1.7$	$8.11 \pm 5.2$	$3.29 \pm 0.9$	$25.79\pm8.5$	$2.2 \pm 0.45$			
PA	$4.16 \pm 1.9$	$8.16 \pm 5.6$	$3.92 \pm 1.2$	$22.86\pm8.3$	$1.8 \pm 0.23$			
Emulsion-wash (24H PVA/8H ETOH) (N = 40)								
Dry	$4.15 \pm 1.6$	$3.94 \pm 3.1$	$2.08\pm0.9$	$11.06 \pm 6.7$	$5.2\pm0.57$			
EI	$1.07\pm0.7$	$1.38 \pm 1.2$	$7.95\pm4.8$	$9.35\pm5.9$	$1.5 \pm 0.52$			
TOL	$1.22 \pm 0.7$	$1.63 \pm 1.3$	$9.23 \pm 15.1$	$10.48\pm5.9$	$1.5 \pm 0.49$			
2PE	$1.28 \pm 0.9$	$1.47 \pm 1.8$	$12.42 \pm 20.6$	$10.58 \pm 6.6$	$1.4 \pm 0.37$			
PA	$1.25 \pm 0.8$	$1.23 \pm 1.5$	$13.93 \pm 25.0$	$10.58 \pm 6.6$	$1.4 \pm 0.32$			
	Se	emi-Cont (75L,	<b>25G)</b> $(N = 30)$	)				
Dry	$5.26 \pm 1.6$	$4.23 \pm 2.2$	$2.88 \pm 1.1$	$10.15 \pm 5.7$	$4.2\pm0.18$			
EI	$6.80 \pm 1.4$	$7.17 \pm 2.4$	$3.34 \pm 1.0$	$23.40 \pm 11.5$	$5.0 \pm 0.32$			
TOL	$7.88 \pm 1.5$	$9.63 \pm 2.9$	$2.85 \pm 1.0$	$31.85 \pm 10.7$	$5.2 \pm 0.47$			
2PE	$6.15 \pm 1.9$	$13.35 \pm 3.9$	$2.75\pm0.5$	$29.15 \pm 9.4$	$2.3\pm0.50$			
PA	$4.90 \pm 2.3$	$10.74 \pm 5.6$	$3.27 \pm 0.8$	$25.35 \pm 9.3$	$1.9 \pm 0.39$			
Semi-Cont (85L, 25G) (N = 30)								
Dry	$4.90 \pm 1.6$	$5.14 \pm 2.7$	$2.84 \pm 1.3$	$13.83 \pm 10.0$	$4.1 \pm 0.61$			
EI	$4.66 \pm 2.4$	$5.72 \pm 2.5$	$3.23 \pm 1.2$	$21.85 \pm 14.4$	$3.9\pm1.07$			
TOL	$6.36 \pm 3.4$	$11.05 \pm 6.1$	$3.09 \pm 1.4$	$31.45 \pm 19.4$	$3.7 \pm 1.39$			
2PE	$3.14 \pm 2.8$	$4.05 \pm 6.5$	$7.46 \pm 4.0$	$20.46 \pm 13.7$	$1.5 \pm 0.44$			
PA	$2.68 \pm 2.4$	$2.49 \pm 5.5$	$9.30 \pm 4.7$	$19.05 \pm 11.4$	$1.4 \pm 0.31$			
Vivitrol (RLD) (N = 40)								
Dry	$5.97 \pm 2.2$	$6.47 \pm 2.6$	$2.80\pm0.9$	$18.95\pm7.9$	$5.2 \pm 0.93$			
EI	$3.43 \pm 3.2$	$4.91 \pm 5.1$	$5.70 \pm 3.0$	$21.47 \pm 14.3$	$3.2 \pm 1.67$			
TOL	$3.66 \pm 3.3$	$5.90 \pm 5.1$	$5.01 \pm 1.4$	$24.07 \pm 14.6$	$3.4 \pm 1.82$			
2PE	$4.56\pm4.9$	$9.12\pm9.5$	$4.25 \pm 1.6$	$26.65\pm21.6$	$2.6 \pm 1.61$			
PA	$4.51\pm4.7$	$9.62 \pm 10.4$	$4.53\pm2.2$	$24.41 \pm 18.8$	$2.2 \pm 1.12$			

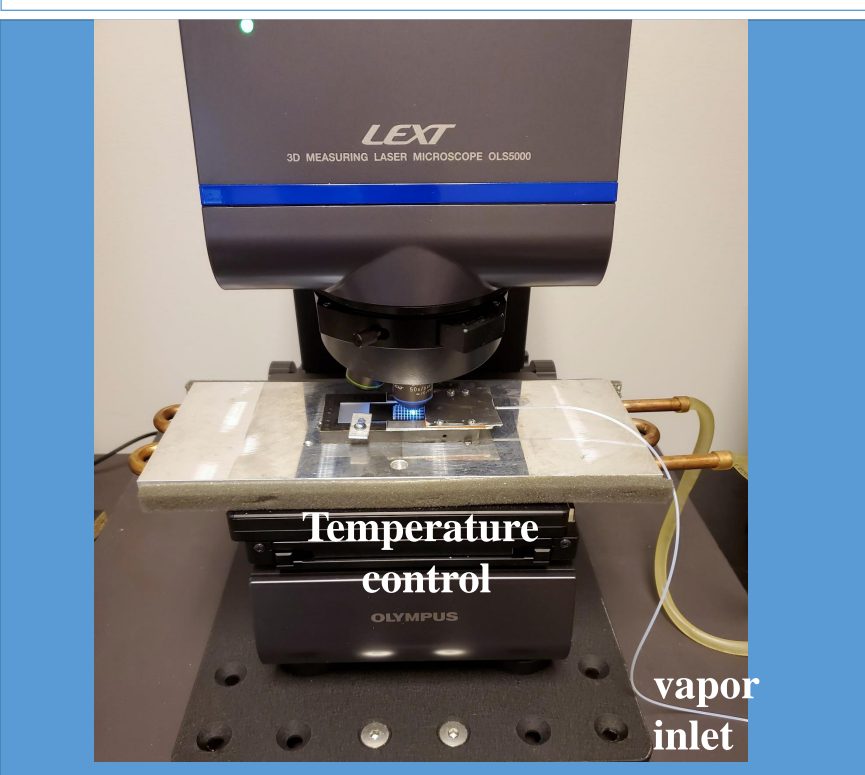
#### Methods

PLGA-NTX microparticles were manufactured by an emulsion method with controlled processing parameters. Additionally, Naltrexone XR Inj, the reference listed drug (RLD), Vivitrol<sup>®</sup>, and samples of particles generated by semicontinuous (SC) manufacturing [2] were also assayed for comparison. Each sample was spread across a microscope slide, and a series of particles were imaged using laser scanning confocal microscopy (LSCM) with particle locations programmed in for future imaging. The particles were warmed to 30°C and exposed to the horizontal flow of argon (40 mL/min) carrying 95±18 mg of a semisolvent for 10 minutes. Afterward, the particles were imaged, and then the particles were exposed to the next semi-solvent in the sequence. The semi-solvents used were ethyl isobutyrate (EI, PLGA solubility threshold of 85% lactide content, or 85L), toluene (Tol, 78L), 2-pentanone (2PE, 69L), and propyl acetate (PA, 63L) (**Fig 1**). Profilometry was performed using LEXT (Olympus) software.



# Conclusion

The SAVI method sequentially melts polymer layers away and allows for observation of differences in particle microstructure. It can be a valuable tool for testing long-acting injectable



# Particles were imaged (Fig 2), and LEXT software parameters were characterized for root mean square height (Sq, roughness), core height (Sk, bulk height variance), kurtosis (Sku, peak sharpness), 10 point height (S10z, tallest 5 peaks minus lowest 5 valleys), and area ratio (Ar, surface area/contact area) (Table 1). Reduced roughness (Sq) of PLGA blank after Tol exposure relative to all other samples reflects the smoother surface of the drug-free sample relative to drug-loaded samples. Area ratio (Ar) indicates the collapse of the particle, and the SC-85L-25G exhibited higher degree of collapse relative to SC-75L-25G, likely due to more soluble 85L PLGA. Additionally, Emulsion with 4 h in the PVA bath exhibited less collapse in EI and TOL relative to the emulsion with 24 h in the PVA bath. This correlated to the difference in measured number average molecular weight $(M_n)$ of the formulations as extra time in the emulsion bath led to increased hydrolysis of the sample with lower M<sub>n</sub> and more susceptibility to collapse. The Vivitrol<sup>®</sup> RLD exhibits significantly higher kurtosis (Sku) across EI, TOL, and 2PE relative to both Emulsion-4H and

formulations as part of research and quality control applications.

#### References

- [1] J. Garner, J. Hadar, S. Skidmore, F. Jessmon, H. Park, K. Park, Y. K. Jhon, B. Qin, Y. Wang. "Scanning Analysis of Semi-Solvent Impact (SASSI) assays of naltrexone microparticles manufactured using different solvents" Scientific poster presented at 2021 annual meeting of Controlled Release Society.
- [2] Sharifi, Farrokh, Andrew Otte, Gwangheum Yoon, and Kinam Park. "Continuous in-line homogenization process for scale-up production of naltrexone-loaded PLGA microparticles." Journal of Controlled Release 325 (2020): 347-358.

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FDA

# Figure 1. Confocal Microscope mounted with Scanning Analysis

Vapor Impact (SAVI) assembly on the

#### platform.

- SC-75L (P < 0.01). Kurtosis is related to peak sharpness and can
- be affected by the presence of NTX, which presents as poorly
- soluble cubic crystals that poke up from the melting polymer and
- provide a blocky structure. This exhibits a different Sku relative to
- the valleys and ravines structures observed for NTX in Vivitrol<sup>®</sup>.



