

Title	Spray dried oleanolic acid powder for pulmonary delivery
Author(s)	Chen, S; Tong, HHY; Kwok, PCL
Citation	The 2013 Conference of Inhalation ASIA, Hong Kong, 26-28 June 2013. In Conference Abstracts Book, 2013, abstract no. 13PS51
Issued Date	2013
URL	http://hdl.handle.net/10722/190143
Rights	2013@Inhalation Asia.

Spray dried oleanolic acid powder for pulmonary delivery

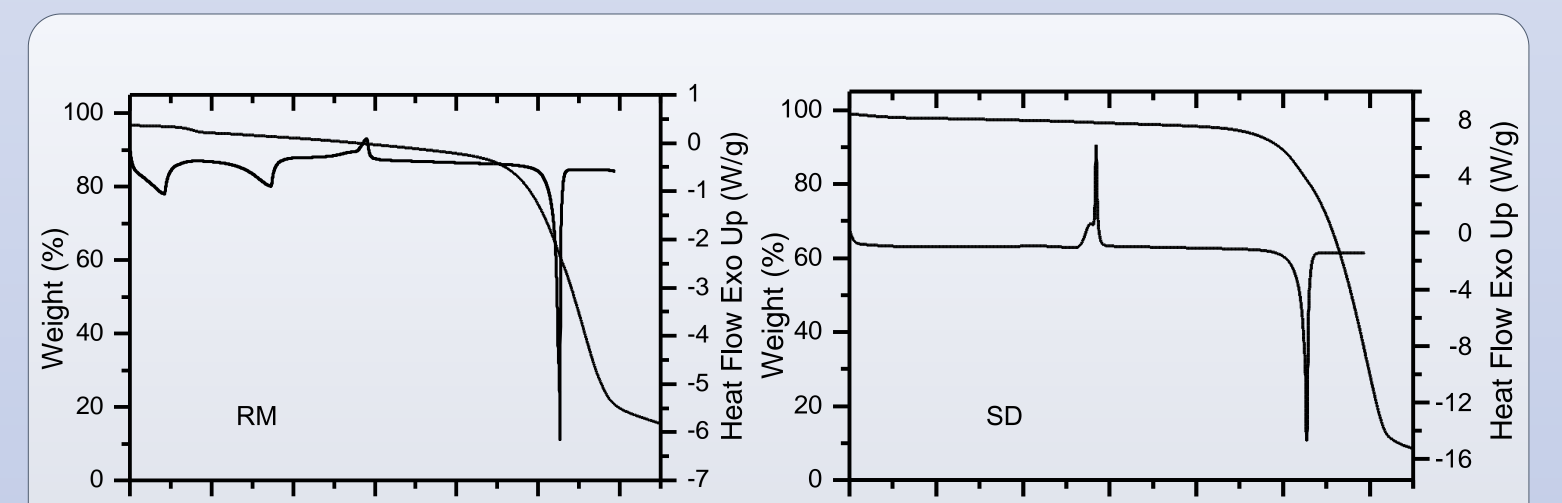


Shuangning Chen¹, Henry Hoi Yee Tong², Philip Chi Lip Kwok^{1*}

¹Department of Pharmacology and Pharmacy, The University of Hong Kong, 21 Sassoon Road, Hong Kong ²School of Health Sciences, Macao Polytechnic Institute, Macao

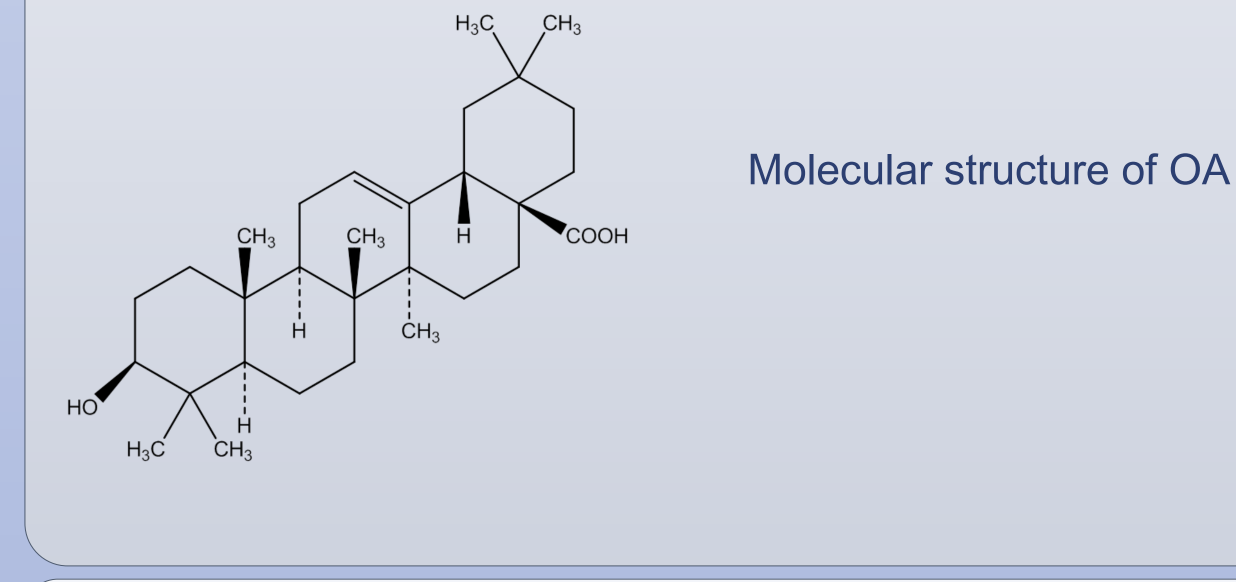
INTRODUCTION

 \succ Oleanolic acid (OA), well known for its hepatoprotective effect ¹, has been shown in vitro to be cytotoxic in A549 human non-small-cell lung cancer cell line². Thus it may be potentially useful for lung cancer treatment. Being a BCS Class IV drug, it has low oral bioavailability³. Therefore,



inhalation is the preferred route of administration for local delivery.

 \succ The aim of this study is to develop an inhalable oleanolic acid dry powder formulation.



Methods

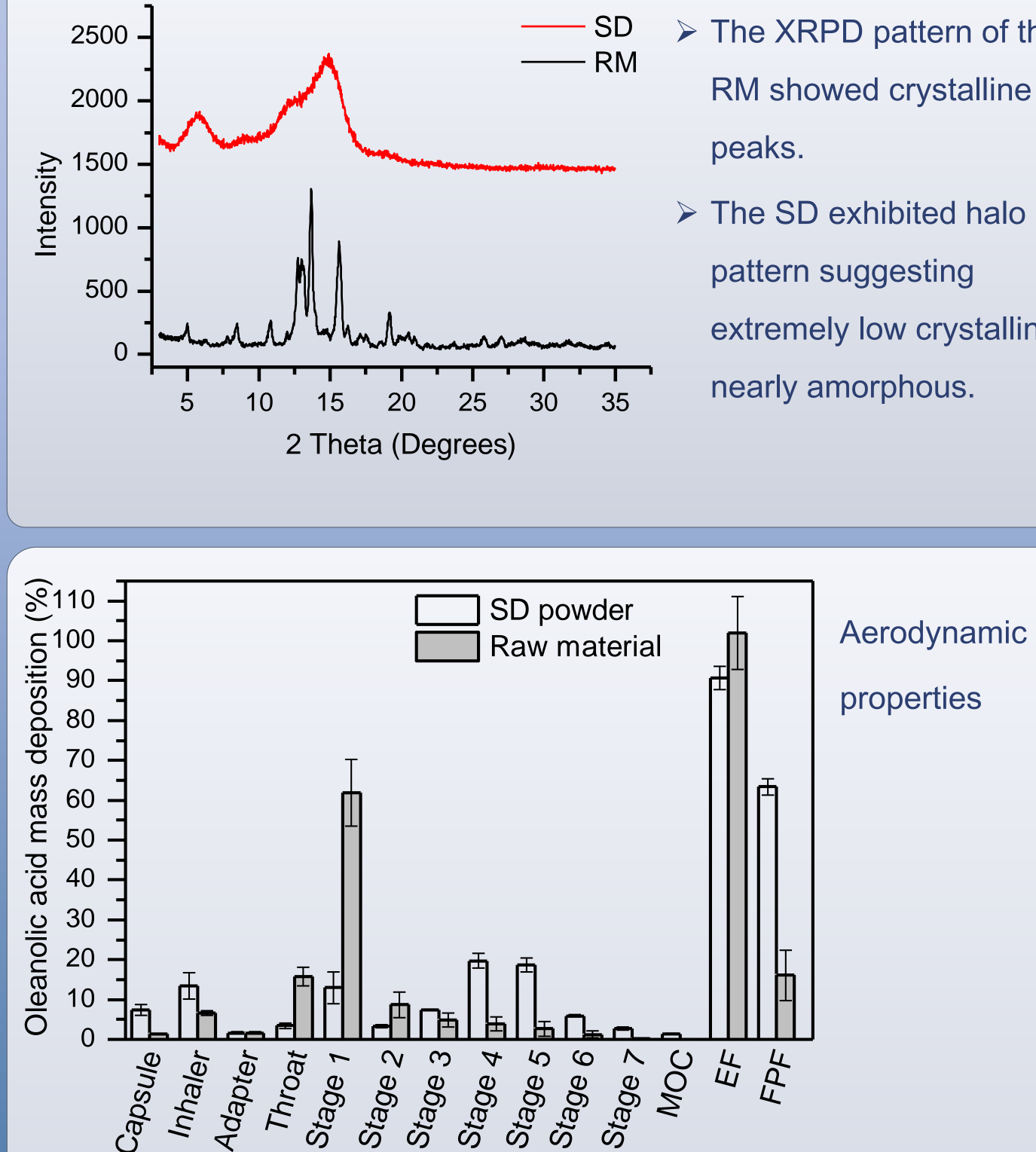
> OA was spray dried from an acetone solution using a Büchi B-290 Mini Spray Dryer. The spray dried powder was characterized and compared with raw OA.

100 150 200 250 300 50 350 Temperature (°C)

300 350 100 200 250 50 150 Temperature (°C)

Thermal analysis

 \succ For the SD, the exothermic process was observed at around 190 °C followed by endothermic process at around 310 °C with concomitant weight loss.

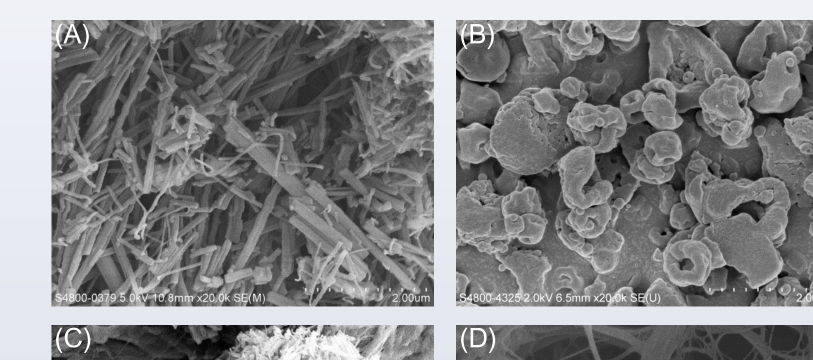


> The XRPD pattern of the RM showed crystalline

The SD exhibited halo pattern suggesting extremely low crystallinity nearly amorphous.

- Particle morphology was observed by scanning electron microscopy (SEM), whereas aerodynamic performance was measured by dispersion from an Osmohaler[™] into a Next Generation Impactor (NGI).
- > The solid state of dry powders was studied by thermal analysis and X-ray powder diffraction.

RESULTS



SEM pictures of raw (A, C) and spray dried (B, D) OA particles before (A, B) and after (C, D) dispersion.

The spray dried formulation exhibits a significantly higher fine particle fraction (FPF) (63.4 \pm 2.1%) than that of the raw material (16.1 \pm 6.3%), indicating an



Raw OA particles were needle-like, while the spray dried ones were corrugated spherical of 0.5–3 µm in diameter.

> After dispersion, spray dried OA could be dispersed into primary particles while the raw material seriously agglomerates.

enhanced dispersion efficiency.



An OA dry powder formulation was successfully prepared by spray drying. It showed excellent aerosol performance (63% FPF) and may be useful for pulmonary delivery.

References

(1) Liu, J. Journal of ethnopharmacology 1995, 49, 57. (2) Liu, Q.; Liu, H.; Zhang, L.; Guo, T.; Wang, P.; Geng, M.; Li, Y. European Journal of Medicinal Chemistry 2013, 64, 1. (3) Tong, H. H.; Wu, H. B.; Zheng, Y.; Xi, J.; Chow, A. H.; Chan, C. K. International journal of pharmaceutics 2008, 355, 195.