

## **LANtern® (Lupus Asian Network) at Hospital for Special Surgery is pleased to offer a free lupus support group for the Asian community in Cantonese (Virtual)**

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We meet quarterly on the 1st Saturday at 11AM-12:30PM Eastern (Feb, May, Aug, Nov)

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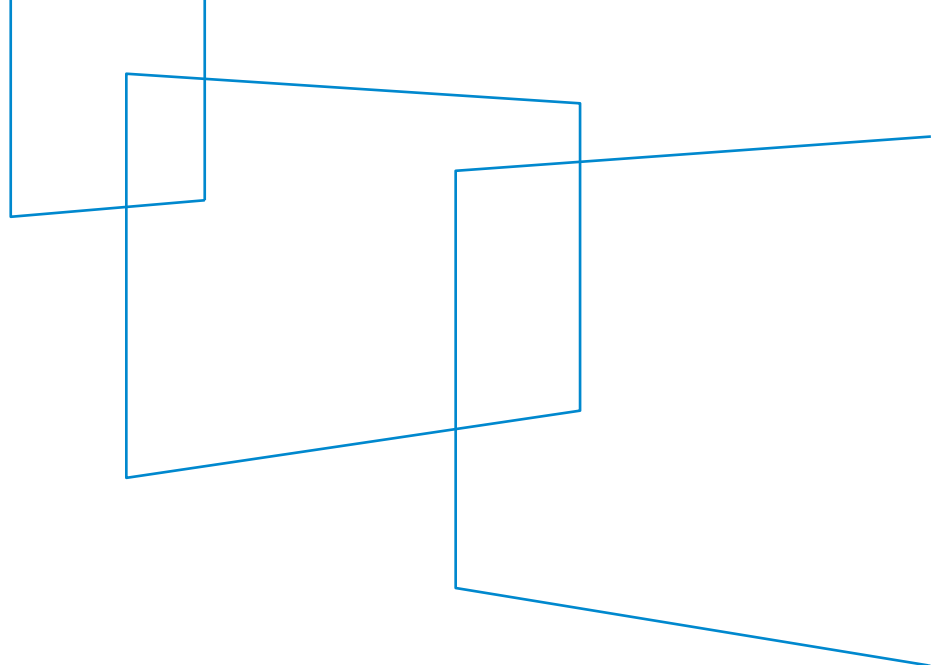
Eliza Ngan, LANtern® Supervisor, will facilitate the group discussion – learn about lupus, share coping strategies, and feel supported on your lupus journey.

For questions and registration, please call Eliza at (US) 646.797.8338 or email [ngandittgene@hss.edu](mailto:ngandittgene@hss.edu)

LANtern®(Lupus Asian Network) is a hospital-based bilingual support and education program dedicated to serving Asian Americans with lupus and their families.

### **Hospital for Special Surgery**

535 East 70th Street,  
New York, NY 10021  
hss.edu



## 紅斑狼瘡亞裔關懷聯盟設於 HSS 醫院為亞裔狼瘡患者和家人提供免費粵語 互助支持小組 (網上)

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會議於每季度月份第一個星期六  
11AM-12:30PM 東部時間舉辦 (2月, 5月, 8月, 11月)

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顏小源 (關懷聯盟主任) 帶領小組討論 - 在這裡您可以了解狼瘡，一起分享應對策略，並在狼瘡之旅中感受到支持。

查詢及報名請聯絡顏小姐 電話 (US)646.797.8338 電郵  
[ngandittgene@hss.edu](mailto:ngandittgene@hss.edu)

設於HSS醫院的紅斑狼瘡亞裔關懷聯盟, 致力為亞裔狼瘡患者及其家人  
提供雙語支援和教育服務

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