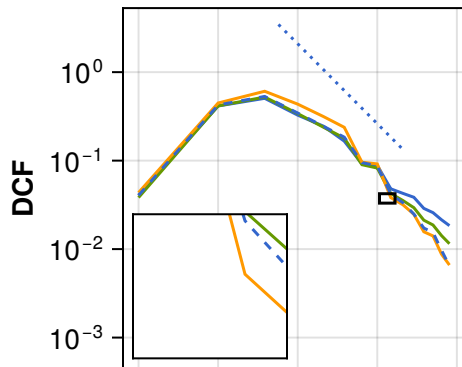
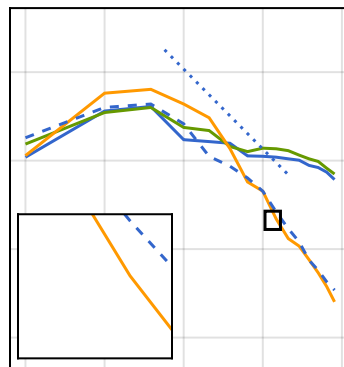


# Energy spectra (face-average, n = 32)

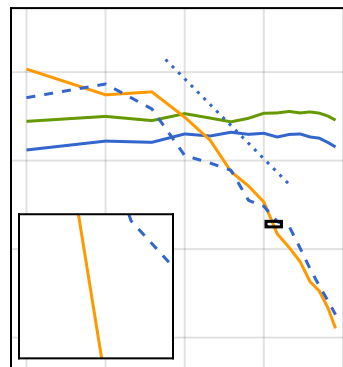
t = 0.1



t = 0.5



t = 1.0



t = 2.0

