

IR Pump-Probe Spectroscopy of Nanoconfined Water Structure in Reverse Micelle

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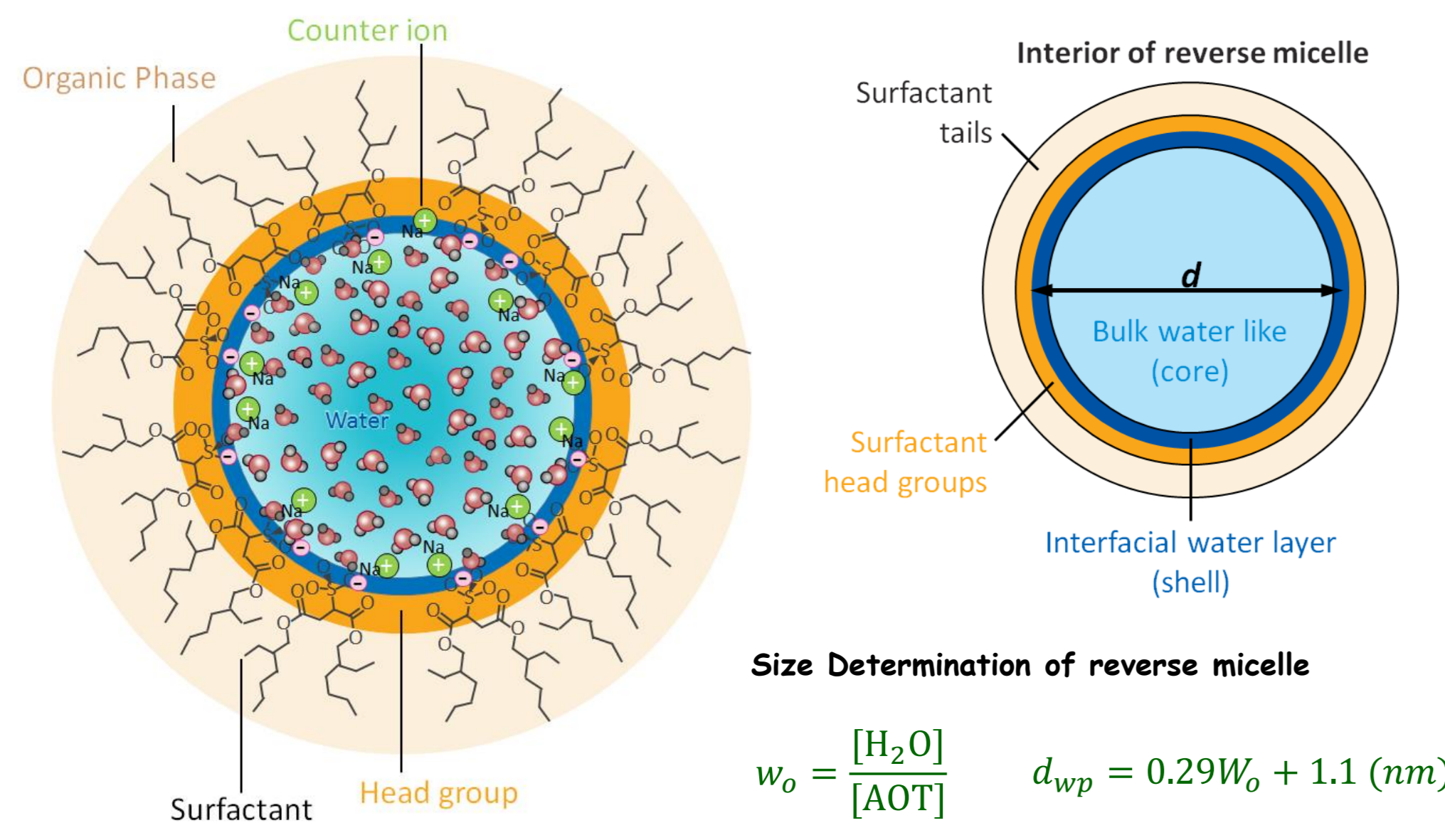
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Interior of Reverse Micelle

(M. D. Fayer et al. *Annu. Rev. Anal. Chem.* 2010, 3, 89-107)

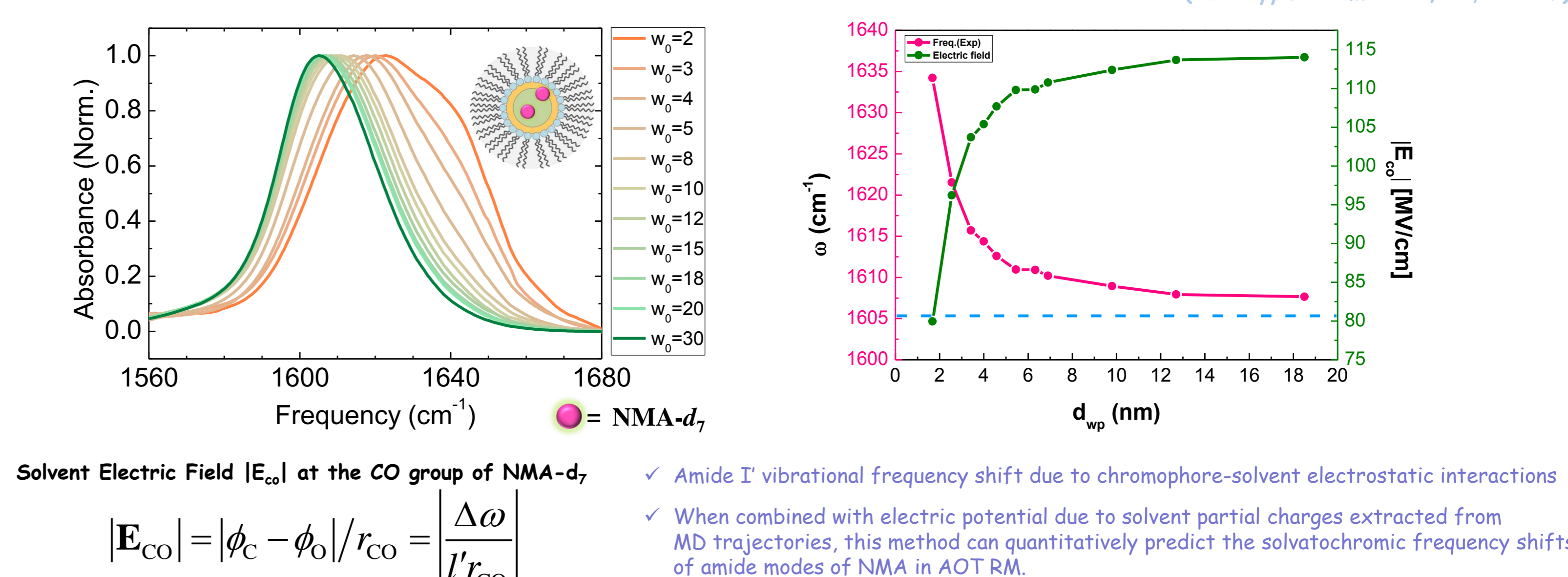


w_0	d_{wp} (nm)	V_{wp} (10^{-27})	# of H ₂ O	# of Solute
2	1.68	2.482713	82.99008	0.747545
5	2.55	8.681988	290.2144	2.614146
8	3.42	20.94483	700.1268	6.30649
10	4	33.51032	1120.156	10.08996
12	4.58	50.30314	1681.492	15.14627
15	5.45	84.75945	2833.269	25.52107
18	6.32	132.1752	4418.243	39.79794
20	6.9	172.0069	5749.706	51.79128
30	9.8	492.807	16473.15	148.3842
40	12.7	1072.531	35851.68	322.939

* Solute : ¹NMA-d₇, ²HN₃

Electrostatic Environment in RMs

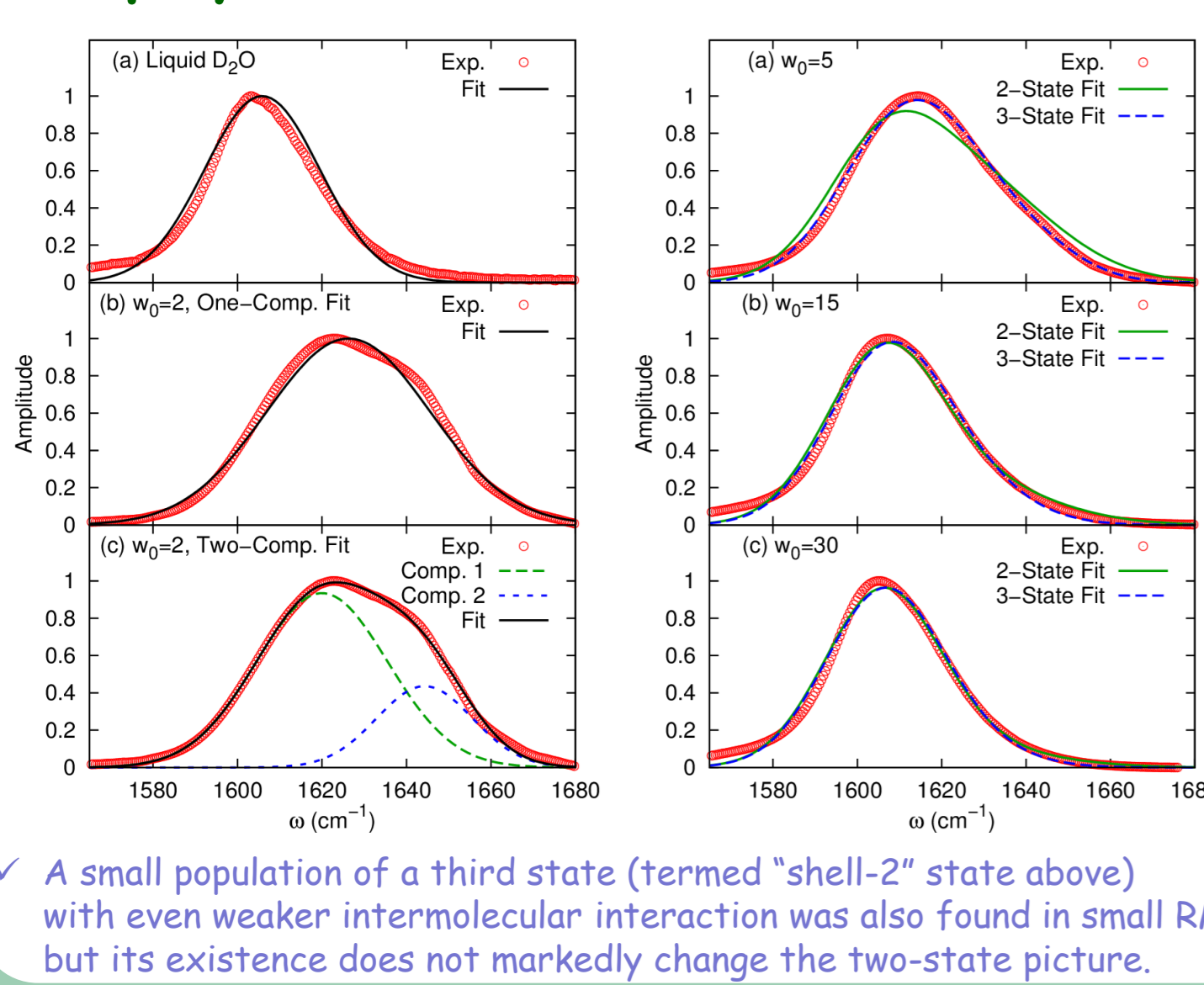
(*Pure Appl. Chem.* 2014, 86, 135-149)



Theoretical Background of Two-/Three-Ensemble model

(J. Lee et al. *Pure Appl. Chem.* 2014, 86, 135-149)

Employment of Subensemble Model



MD Simulations

Molecular mechanics (MM) force field

- Isocane: TraPPE united atom force field (No explicit hydrogen)
- AOT anion: Hybrid TraPPE (carbon tail) and CGenFF 2b7 (head group)
- Water: Rigid TIP4P model
- Na⁺: CHARMM or Jensen-Jorgensen
- NMA-d₇: CHARMM

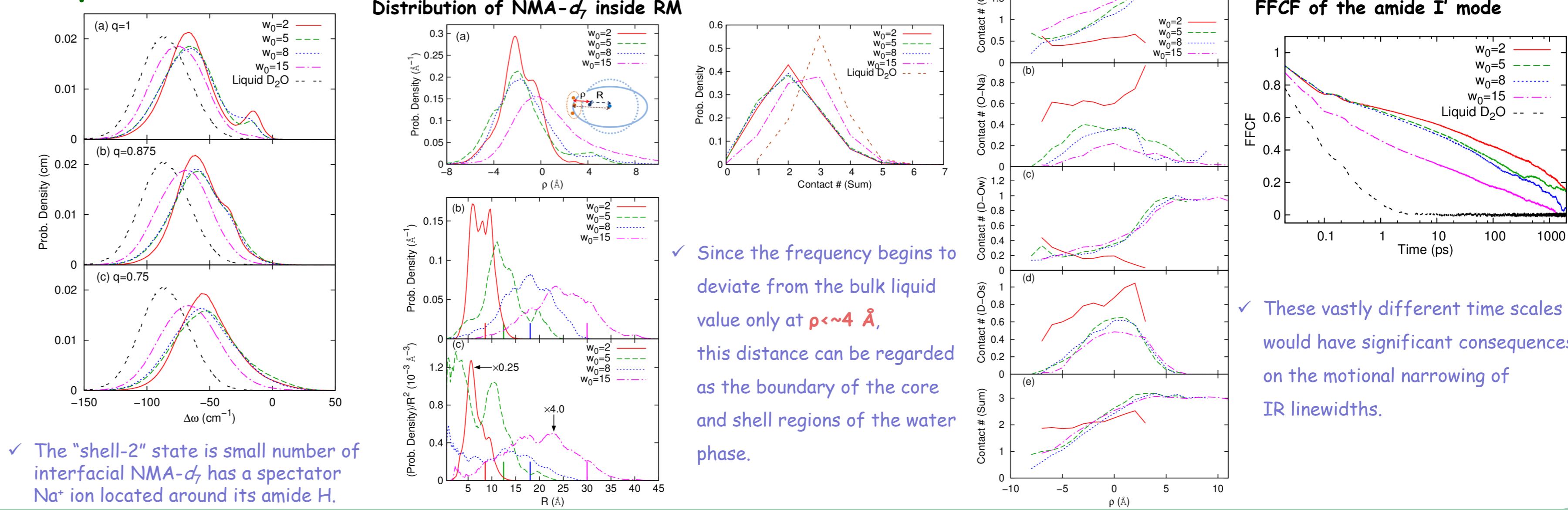
CHARMM and NAMD packages are used for system build-up and NAMD is used for MD.

All bonds involving H/D are constrained.
2 fs time step size
Periodic boundary condition and particle mesh Ewald (PME) electrostatics.
Lennard-Jones interactions are cut off at 12 Å.

To maintain spherical micelle structure, harmonic restraints are applied on surfactant sulfur atoms during equilibration of more than 10 ns.
In production runs of 1.5-2 ns, only water and Na⁺ are weakly restrained within a sphere.

Configurations were saved every 0.05 ps for FFCF calculation.

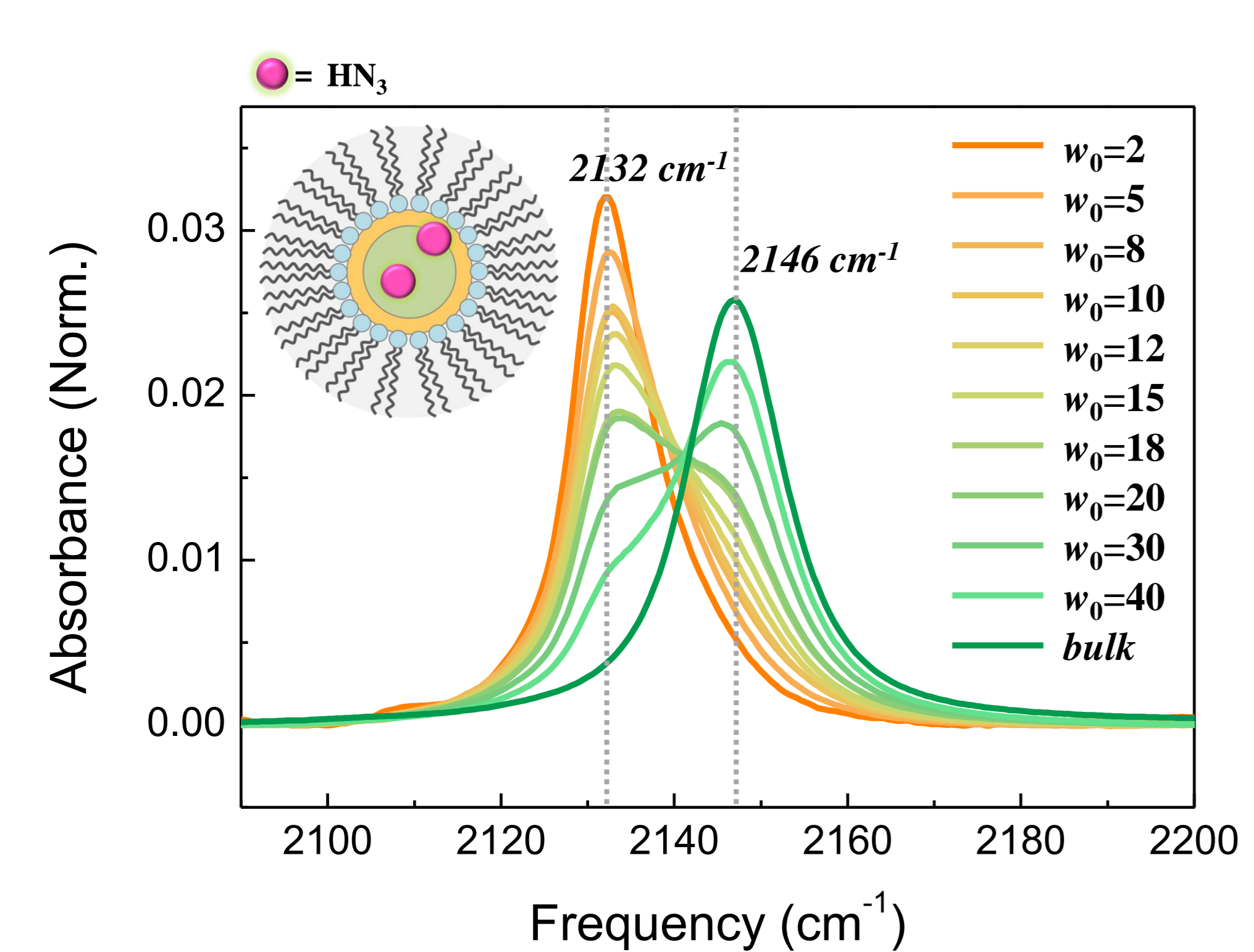
Computational Results



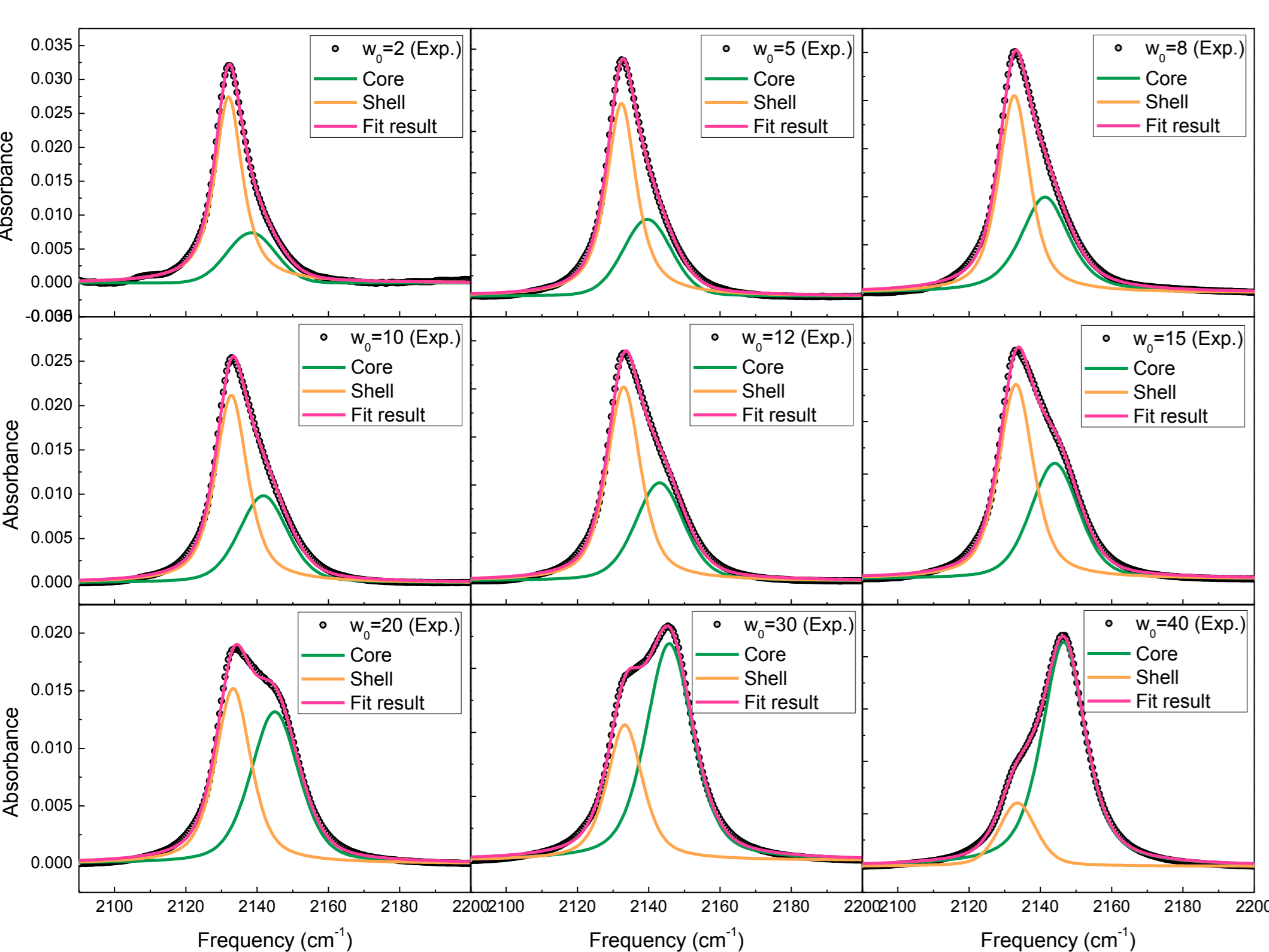
Simple Two-Ensemble model

(J. Lee et al. *J. Phys. Chem. Lett.* 2014, 5, 3404-3407)

FT-IR Spectroscopy



$$A_{total}(\omega) = A_{core}(\omega) + A_{shell}(\omega)$$

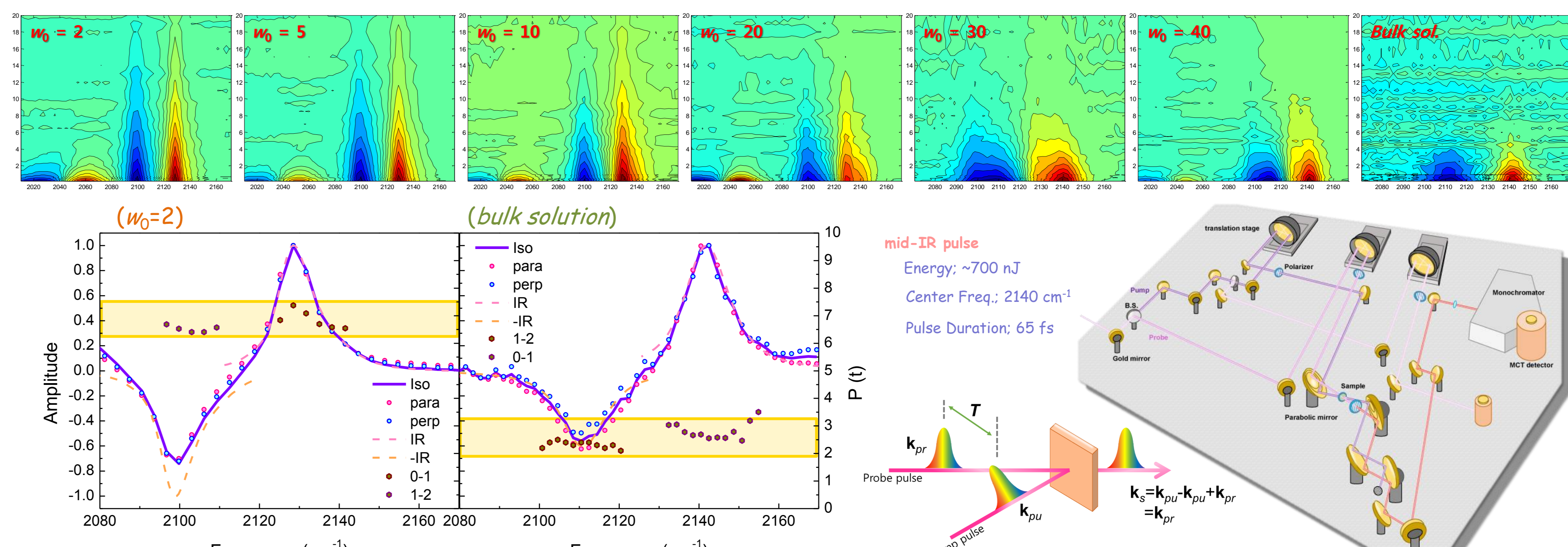


- The hydrazoic acid has been found to possess most of important characteristics of good IR probes for studying water structures in nanoconfined environment.
- The spectral components commonly referred to as core and shell are well-separated and easily identified in the infrared absorption spectra.

Expansion of Two-Ensemble(Core/Shell) Model

(J. Lee et al. *J. Phys. Chem. Lett.* 2014, 5, 3404-3407)

IR Pump-Probe Spectroscopy



$$C(t) = A_{core}e^{-t/T_{core}} + A_{shell}e^{-t/T_{shell}}$$

