Supporting Information

for

Conformational Sensitivity of Surface Selection Rules for Quantitative Raman Identification of Small Molecules in Biofluid

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Figure S1. Diameter statistics of PS beads in Figure 1a.

Figure S2. SEM images of polymer films formed under different experimental parameters. (a) PS/PMMA=1:9. (b) PS/PMMA=2:8. (c) PS/PMMA=3:7. Spin-coating speed at (d) 3500rpm, (e) 4000rpm and (f) 4500rpm.
Figure S3. Diameter statistics of PS beads in Figure S2c.

Figure S4. Diameter chart of the distribution holes in the PMMA film in Figure 1b.
Figure S5. Ag structural growth conditions optimization with different soaking time in 3 mM AgNO₃ solution: (a) 5 s, (b) 10 s, (c) 15 s, (d) 2 mM AgNO₃ for 20s.

Figure S6. Confocal images of evaporation traces of R6G solution.
Figure S7. The Raman shift at 230 cm\(^{-1}\) represents the Ag-S bond.

Figure S8. SERS signal of 10\(^{-4}\) M DA solution on superhydrophobic substrate.
Figure S9. SERS reproducibility of MPBA on as-prepared platform via statistics analysis of a ratiometric indicator of relative peak intensity, I_{1582/1071}.

Figure S10. (a) Averaged SERS spectrum of MPBA-DA complexing at 30 min produced by triplicate measurements with standard deviations at each wavenumber. (b) Time-course variations (Δr) of r_{1170/1071}, r_{1380/1071}, and r_{1582/1071} in Figure 2d. (d) Time-dependent values of r_{1170/1071}, r_{1380/1071}, and r_{1582/1071} in Figure 2d.
**Figure S11.** The Raman spectra of MPBA-DA and MPBA-DA bonded to an Ag\textsubscript{4} cluster.

**Figure S12.** MPBA vibration mode at 1050 cm\textsuperscript{-1} and 1558 cm\textsuperscript{-1} in simulated Raman.

**Figure S13.** MPBA vibration mode at 1050 cm\textsuperscript{-1} and 1060 cm\textsuperscript{-1} in simulated Raman spectra of MPBA and MPBA-DA.
Figure S14. DFT calculated Raman spectrum of MPBA-DA and contributions of transition bands.