Experimental study on the heterogeneous reaction of gaseous OH radical with aqueous DMSO: Determination of the $CH_xSO_x^{-1}/SO_a^{-2}$ production ratio

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The oxidation of dimethyl sulfide (DMS) emitted from ocean (~ 45 Tg S per year) is a global source of cloud condensation nuclei. Hydrophobic DMS is mostly oxidized in the gas-phase into H₂SO₄(g) + DMSO(g) (dimethyl sulfoxide), whereas water-soluble DMSO is oxidized into $SO_4^{2-} + CH_3SO_3^{-}$ (methane sulfonate) on water surfaces. $R = CH_3SO_3^{-1}/SO_4^{-2}$ ratios therefore indicate the extent of DMSO heterogeneous oxidation if $R_{\text{het}} = \text{CH}_3\text{SO}_3^{-7}/\text{SO}_4^{-2-}$ for DMSO(aq) + \cdot OH(g) were determined. Here, products and intermediates of the oxidation of aqueous DMSO initiated by gas-phase hydroxyl radicals, OH(g), at the air-water interface were directly detected by mass spectrometry in a novel setup under various experimental conditions. Exposure of millimolar DMSO aqueous microjets to ~ 10 ns OH(g) pulses from the 266 nm laser flash photolysis of $O_3(g)/O_2(g)/H_2O(g)/N_2(g)$ mixtures yielded an array of interfacial intermediates/products, including $CH_3SO_3^-$ and HSO_4^- , that were unambiguously and simultaneously identified in situ by mass spectrometry. We determined $R_{\rm het}$ = 2.7 from the heterogeneous OH-oxidation of DMSO on aqueous aerosols for the first time. The nearly quantitative production of $H_2SO_4(g)$ (that leads to SO_4^{2}) in the oxidation of DMSO in the gas-phase versus the R $_{\text{het}} \sim 2.7$ value determined at the air-water interface means that $R = \text{CH}_3\text{SO}_3^-/\text{nss-SO}_4^{-2}$ variations in the aerosol, particularly in remote locations, should arise from the competition between the gas-phase versus the heterogeneous DMSO oxidation pathways. The present study reveals that interfacial OH-oxidation processes play a more significant role in the generation and growth of atmospheric aerosol over ocean than previously envisioned.

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