



Halogen activation from NO_x: Field observations of nitryl chloride (ClNO₂) production from uptake of N₂O₅ to chloride-containing aerosol

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Nitryl chloride, ClNO₂, is produced from the heterogeneous uptake of N₂O₅ to sea salt and other chloride-containing aerosol. Until recently, the impact of this process on the atmospheric halogen budget was thought to be modest. New analytical techniques for measurement of ClNO₂ based on chemical ionization mass spectrometry (CIMS) have quantified its atmospheric abundance during recent field campaigns. The observations show unexpectedly large ClNO₂ mixing ratios, in excess of 1 ppbv within marine NO_x plumes. Production of these levels requires heterogeneous uptake of N₂O₅ both to sea salt aerosol and to chloride-containing submicron aerosol, indicating that the production of ClNO₂ from NO_x pollution sources is more efficient than commonly recognized in atmospheric models. This presentation will describe the implications of these measurements for air quality in polluted, coastal areas as well as the possible magnitude of ClNO₂ as a global halogen source.