

Structural investigation of platinum solubility in silicate glasses

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ABSTRACT

The coordination environment of 20–200 ppm Pt in yellowish glasses from the CaO–Al₂O₃–SiO₂ (CAS) ternary was studied using X-ray absorption fine structure spectroscopy at the Pt-L_{III} edge. Analysis of the Pt-L_{III} edge region suggests that Pt in these glasses is mainly tetravalent and sixfold-coordinated by O (with a mean Pt–O distance of 2.08 ± 0.02 Å). No evidence for Pt²⁺ or Pt⁶⁺ was found in any of the glasses studied, suggesting that one can not derive valence information easily from solubility data. No second-neighbor contribution was observed around Pt⁴⁺O₆ polyhedra. However, bond-valence modeling suggest that these polyhedra are likely to bond mostly to ^{IV}Ca²⁺, which should promote high positional disorder of second-neighbor cations around Pt. This particular bonding arrangement may explain the relatively high solubility of Pt in these relatively depolymerized melts, as CaPtO₃-type units.