D$_2$ as the reaction gas to improve robustness of K isotope analysis by collision cell MC-ICP-MS

SHICHAO AN, JIAYANG CHEN, SAMUELE BOSCHI AND WEIQIANG LI

Nanjing University
Presenting Author: asch@nju.edu.cn

Stable K isotope is a newly emerging research tool for various research areas. K isotope investigations have mushroomed since five to ten years ago due to high-precision K isotope ratio analytical methods developed intensively. K isotope ratio measurements using collision cell multi-collector ICP mass spectrometers (CC-MC-ICP-MS), emerging in the last 2-3 years, have boosted the K isotope investigation due to its high signal sensitivity and convenience during analysis. However, previous studies have reported that trace-level Ca contaminants can induce severe K isotope offset, which shows a significant influence of Ca contamination on the robustness of K isotope analysis by CC-MC-ICP-MS. In this study, we demonstrate that the cause why Ca contamination lays the influence on K isotope analysis by CC-MC-ICP-MS is the formation of positively charged calcium-hydride molecules in the collision cell. Usage of D$_2$ other than H$_2$ generates $^{40}$CaD$^+$ that does not interfere with $^{41}$K$^+$ as $^{40}$CaH$^+$ does, so the influence can be dramatically reduced using D$_2$ instead of H$_2$ (the default gas) as the reaction gas that goes into the collision cell, which makes the robustness of K isotope analysis by CC-MC-ICP-MS significantly enhanced. K isotopic results of seven geo-standards by this method agree well with the literature data, which confirms the validity of the method. With the improved method, we have performed K isotope studies on a 17 mg lunar basalt fragment from China’s Chang’e-5 lunar returned samples at sub-microgram K consumption. Six mineral separates (plagioclase and pyroxene, three each) picked up from the lunar basalt fragment, weighted low down to 0.5 mg and 400 ng K mass, have shown varied K isotopic compositions with $\delta^{41}$K of plagioclase from -0.60‰ to +0.03‰, and pyroxene from -0.37‰ to -0.18‰, which firstly reveals significant millimeter-scale K isotope variability in lunar samples worldwide.