In Situ Sims, Micro-Baddeleyite U-Pb Dating Method for Mafic Rocks

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Mg and Si isotopes in the solar system
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Chemical heterogeneity in our solar system is well-documented. Silicate portions of the Earth, Moon and Mars are different from the chondritic meteorites in their volatile element budgets and in Mg/Si. Isotopic compositions also vary between Earth and other Solar System bodies. Chemical and isotopic differences between Earth and bulk meteorites can be broadly attributed to the following processes: (1) Terrestrial fractionation, (2) Fractionation and incomplete mixing in the solar nebula prior to accretion and (3) Late injection of pre-solar material and its inhomogeneous distribution.

It has been suggested [1] that the Earth and Moon have heavier Si-isotopic compositions compared to meteorites, all of which have indistinguishable Si-isotopes, resulting from core formation on Earth. Several efforts have been initiated to resolve the Mg isotopic compositions of the bulk Earth and meteorites [2, 3] but available results are contradictory.

We have initiated a combined Mg- and Si-isotopic study in terrestrial and planetary samples to address the following questions: (1) What are the BSE Mg- and Si-isotopic compositions? (2) Does the BSE and average chondrites have resolvable differences in Si and Mg-isotopic compositions and (3) if they are different, what is the most plausible explanation? So far our Mg-isotopic data suggest that terrestrial olivines from peridotites and whole-rock basalts show similar Mg-isotopic compositions which are slightly lighter than those of whole-rock chondrites. Olivines from several pallasites show heavier Mg isotopic compositions compared to both chondrites and terrestrial samples. This difference suggests that pallasites, likely derived from the core-mantle boundary of early differentiated planetesimals, sampled Mg-isotopic heterogeneity in the solar nebula.


In situ SIMS micro-baddeleyite U-Pb dating method for mafic rocks

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Baddeleyite (ZrO2) is a favorite target for U-Pb dating of mafic and ultramafic rocks. Although samples with grains that are large enough (100-200 micron long) for traditional separation and dating methods are scarce, micro-baddeleyite crystals (<20 microns long) are nearly ubiquitous. With refined methods, it is possible to separate 10 to 20 micron long grains for ID-TIMS analysis.

We are currently developing an in situ technique to date micro-baddeleyite crystals using the CAMECA ims 1270 ion microprobe, avoiding the separation and dissolution steps of ID-TIMS and permitting analysis of even smaller grains (2-5μm). The ion microscope optics of the ims 1270 can reduce the effective sampling diameter of a 20 μm primary ion beam to a few to 10 μm. An aperture blocks undesired secondary ions, in particular common Pb ions emitted from host phases surrounding micro-baddeleyite, and the target crystals can be dated in polished thin sections. CAMECA ims 1270 analytical conditions also involve oxygen-flooding the sample chamber (~105 Torr) to enhance Pb secondary ion yields for baddeleyite by approximately 10-fold, and a UO 2/U vs. U/Pb relative sensitivity calibration. Three baddeleyite standards analyzed under these conditions yielded U/Pb reproducibilities that only marginally exceeded those determined for NIST610 glass, suggesting that crystal orientation related bias in U-Pb sensitivity is minor to absent. Proterozoic-aged standards yielded 2 sigma precisions on 207Pb/206Pb dates from 1% to 0.2%. Preliminary results for micro-baddeleyite from Ordovician and Mesoproterozoic-aged mafic sills analyzed in situ yield U-Pb ages roughly equivalent in precision to those achieved on larger standard baddeleyite grains. Precisions on Phanerozoic 206Pb/238U dates are ~6%; Precambrian 207Pb/206Pb date precisions are 1% or less. The in situ technique is applicable to most unmetamorphosed to low-grade mafic rocks including dikes, sills, fine-grained volcanic rocks and extra-terrestrial rocks, greatly increasing the number of mafic samples that are dateable by U-Pb methods.