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Water contents of natural zircons are controlled by their Y+REE contents

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The water and trace element contents of non-metamict igneous zircons were determined to constrain the H incorporation mechanism and to evaluate the use of zircon to constrain water contents of melts [1]. Zircons from Fe-Ti oxide gabbros from the Vema Fracture Zone [2] contain up to 980 ppm H_2O_2 , 1.4 wt% Y₂O₃ and 0.6 wt% P₂O₅ and are generally strongly zoned. Y+REE are partially charge-balanced by P $(Y^{3+}+P^{5+}=Zr^{4+}+Si^{4+})$ but a large Y excess is present (Fig. 1A). On an atomic basis, this excess closely matches the amount of H present in the zircons (Fig. 1B). We therefore conclude that H is incorporated by a charge-balance mechanism $(H^{\scriptscriptstyle +}\!+\!Y^{\scriptscriptstyle 3+}\!\!=\!\!Zr^{\scriptscriptstyle 4+}).$ This is supported by FTIR data, which show a strongly polarised absorption band at 3100 cm⁻¹ similar to experimental Lu-doped hydrous zircons. No other absorption bands are visible, excluding a hydrogrossular-type exchange mechanism. Because of charge-balanced uptake of H, P and Y+REE in zircon, the partitioning of these elements into zircon is dependent on each of their concentrations. Hence, using H in zircon to determine water contents of melts is challenging. As Ce⁴⁺ partitioning is not affected, Ce anomalies may depend on H₂O and P₂O₅ contents of the melt, in addition to its oxidation state [3].

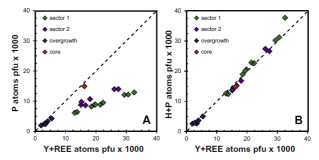


Figure 1: (A) P vs Y+REE (B) H+P vs Y+REE

