

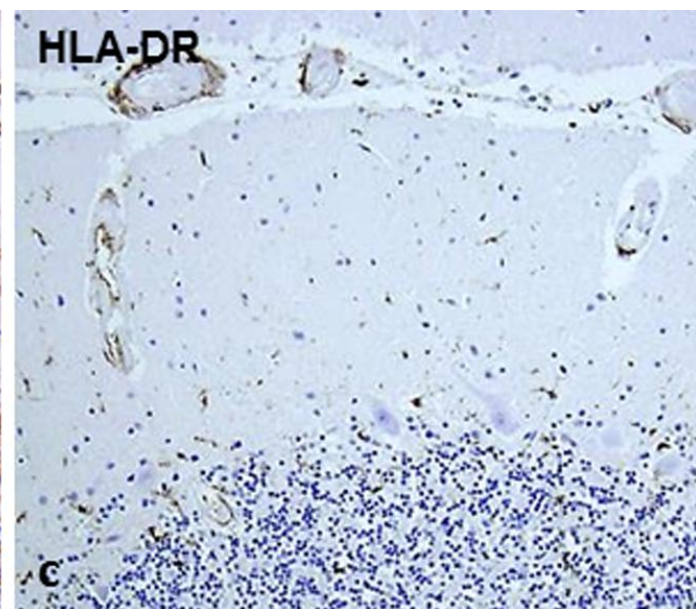
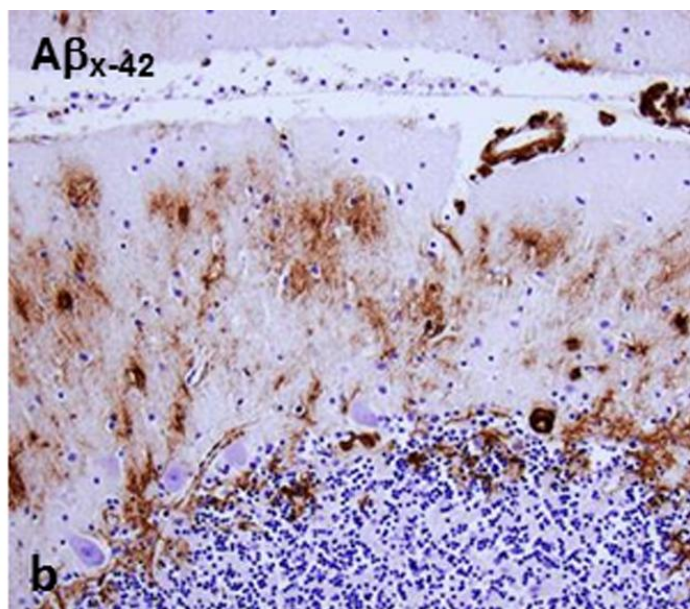
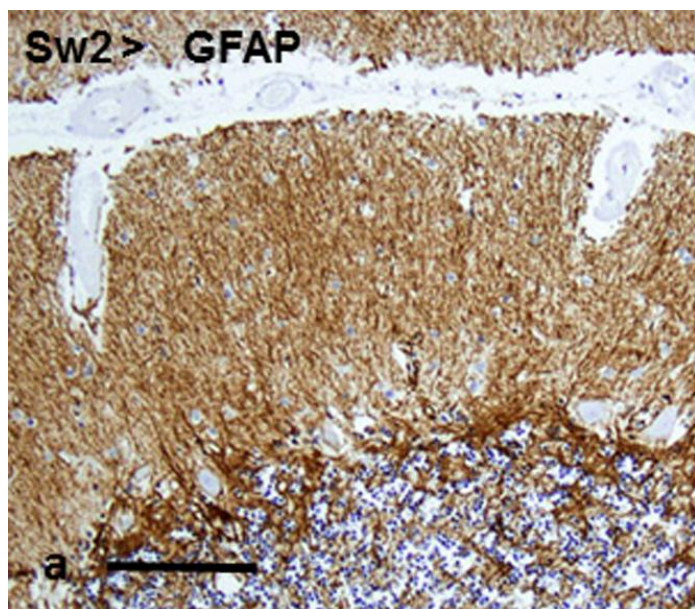
Title: The *Arctic APP* mutation leads to Alzheimer's disease pathology with highly variable topographic deposition of differentially truncated A β

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Suppl. Fig. 8 Sw2 patient's cerebellum. **a:** The density of GFAP-positive network is prominent, especially in the molecular layer. **b:** The density of the network corresponds to the deposition of A β in the Purkinje cell layer, but it differs from the perivascular perivascular accentuation of A β in the molecular layer. **c:** The microglial reaction to A β is minimal (*bar* in **a** 200 μ m for all panels)