Supplementary Data:

ANU Argon Facility Technical Report: ANU07-2021 40Ar/39Ar Analysis for National Argon Map By Marnie Forster and Davood Vasegh

NAM Proposal 06:

New Ar-Ar geochronology age constraints on the Lord Howe Seamount Chain

Methods and procedures

Sample selection and mineral separation:

The samples in this study were provided by Geological Survey of New South Wales and the mineral separation procedures were undertaken in rock crushing and mineral separation laboratories at The Australian National University (Table 1) No chemical or leaching treatments were used during separation.

Mineral separation begins by choosing the most pristine sections with no evidence of weathering or staining. For samples with a targeted microstructure, the rock is first sliced into thin slabs using a trim saw, the selected area was then cut from the rock using a band saw. Once the selected area was separated, it was then crushed, milled and de-slimed as many times as was necessary to clean the grains and finally washed in deionised water.

Whole rock crush procedure:

For whole rock the crushed grains are sieved into size fractions: 250-420µm. A final check and handpicking of the best quality aggregates was done in the Argon Preparation Laboratory.

Sample ID	Target Mineral	Mass (mg)	Grain Size (μm)	Treatment / Comment	Picture
NSWSPJG0075	Basalt (WR)	144.1	250-420	Dark black basalt	
N3W3FJG0075	Dasait (WK)	299.7		grains, clean fraction	

Mineral separation details:

Table 1: Mineral separation details

Sample irradiation details:

Irradiation of samples for ⁴⁰Ar/³⁹Ar analysis was undertaken at the University of California Davis McClellan Nuclear Research Centre, CA, US in Central Facility position of TRIGA reactor without rotation, with 1.0mm of Cadmium shielding as ANU CAN #37 for 8.00 hours on 12 December 2020.

The calculated amounts of grains were weighed and recorded and then wrapped in labelled aluminium packets in preparation for irradiation. The sample filled foils were placed into a quartz irradiation canister together with aliquots of the flux monitor Biotite GA1550. The foil packets of GA1550 standards were dispersed 6-8mm apart throughout the irradiation canister, between the unknown age samples. In addition, packets containing K₂SO₄ and CaF₂ were placed in the middle of the canister to monitor argon isotope production from potassium and other interfering elements. Irradiated samples were unwrapped upon their return to the Australian National University, and then

rewrapped in tin foils in preparation for analysis under vacuum in the furnace. Tin foil is used because the melting temperature of tin is lower than the experiment starting point in the furnace and gasses from tin can be pumped away prior to the sample analysis.

⁴⁰Ar/³⁹Ar procedures and analysis information

Methodology:

Temperature-controlled resistance furnace step-heating experiments is the technique that is used in the ANU Argon laboratory to extract argon isotopes from the samples to ensure 100% release of ³⁹Ar. A sample is dropped into a cleaned furnace and heated to 400°C to melt the tin foil and then left in the furnace at 350°C for 8-12 hours to pump away unwanted gases. This cleaning procedure has proven to significantly improve the quality of the resultant data. The step-heating experiment then starts at 450°C, and each incremental heating step is heated at a constant temperature for 15 minutes. The heating process involves rapid heating to the setpoint temperature with no overshoot, then temperature is maintained for 15 minutes followed by rapid cooling; this procedure produces a square wave in temperature for each heating step. The heating step schedule for biotite and muscovite rises by 30°C increments (except for the last a few steps), with 30 steps per sample, while K-feldspar is analysed in more than 40 steps, including numerous isothermal steps. Diffusion experiments, as conducted in the ANU Argon laboratory, are designed to retrieve diffusion parameters which can be used in quantitative temperature-time modelling. The heating schedules are recorded in the Excel Tables for each sample.

Cleaning of the furnace between samples is vital in this method. The furnace is degassed four times at 1,450°C for 15 minutes and the gas pumped away prior to the loading of the subsequent sample. Blanks are measured to monitor the cleaning process. The flux monitor crystals are fused using a CO_2 continuous-wave laser. Gas released from either the flux monitors or each step of the sample analyses are exposed to three Zr-Al getters; two AP10 (Cold and hot) and one CP50, each for 10 minutes, to remove active gases. The purified extracted gasses are then isotopically analysed in the Argus VI mass spectrometer. The 40 Ar/ 39 Ar dating technique is adapted from McDougall and Harrison (1999) and described in Forster and Lister (2009).

Background levels are measured and subtracted from all analyses, from flux monitors and samples. The nuclear interfering values for the correction factors for the isotopes are listed below (Tetley et al 1980). These are measured for the reactions and uncertainties of $({}^{36}\text{Ar}/{}^{37}\text{Ar})_{Ca}$, $({}^{39}\text{Ar}/{}^{37}\text{Ar})_{Ca}$, $({}^{40}\text{Ar}/{}^{39}\text{Ar})_{K}$, $({}^{38}\text{Ar}/{}^{39}\text{Ar})_{K}$ and $({}^{38}\text{Ar})_{Cl}/({}^{39}\text{Ar})_{K}$, and were calculated prior to sample analyses.

Mass spectrometer setup and procedures

Samples and standards were analysed in the Argon Laboratory at the Research School of Earth Science, The Australian National University, Canberra, Australia using a *Thermo Fisher ARGUS-VI* multi-collector mass spectrometer (Table 2).

Mass Spectrometer:		TI	Thermo Fisher Argus VI			
Detector Type:		Fa	Faraday Cups only x5			
Calibrations:		3	3 levels (Zero Offset, Gain and Cross Calibration)			
Peak Centring:		0	Once for every measurement @H2 (⁴⁰ Ar)			
Measurement Cycles:		5	51 cycles on all detectors			
Extrapolation Method:		E	Exponential extrapolation and uncertainty			
	Name	UEC Offset [f]]		Gain	Cross Calibration Factor	

Na	me	UFC Offset [fA]	Gain	Cross Calibration Factor
Þ	H2	-4.9761469	0.9871203	1
	H1	-2.2071069	0.9671459	1.007184188
	AX	-7.6814703	0.9769602	1.017518151
	L1	-2.3979322	0.9706487	1.030604297
	L2	-3.1329948	0.9676338	1.047244337

Table 2: Detector Calibration Values

The calculation parameters:	
Lambda ⁴⁰ K (Renne et al 2011)	5.5305E-10
Lambda ³⁷ Ar (Kondev et al 2017)	1.9798E-02
Lambda ³⁹ Ar (Kondev et al 2017)	7.0548E-06
Lambda ³⁶ Cl (Kondev et al 2017)	6.2985E-09
Flux Monitor	GA1550 @ 99.18 ± 0.14 Ma
Total irradiation power	8.00 MW
Irradiation Date	12 Dec, 2020
Irradiation shielding	Cadmium 1.0mm
Interfering isotope production ratios:	
(³⁶ Ar/ ³⁷ Ar) _{Ca} correction factor	1.39373E-04
(³⁹ Ar/ ³⁷ Ar) _{Ca} correction factor	1.28113E-03
$(^{40}Ar/^{39}Ar)_{K}$ correction factor	3.68304E-02
$({}^{38}Ar/{}^{39}Ar)_{K}$ correction factor	1.15198E-02
(³⁸ Ar) _{Cl} /(³⁹ Ar) _K correction factor	8.04118E-02
Ca/K conversion factor	1.90
Atmospheric Argon correction ratio:	
⁴⁰ Ar/ ³⁶ Ar (Lee et al 2006)	298.57
⁴⁰ Ar/ ³⁸ Ar (Lee et al 2006)	1,583.52
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Representative air shot and blanks measurements:

The discrimination factor was calculated by analysing five air shots analysis on either side of sample analysis and the calculation of the 1amu was used for the discrimination factor. Table 3 shows an example of the analysed air shots.

Date	⁴⁰ Ar ± %	err	³⁸ Ar ± %err		³⁶ Ar ± %err		1amu ± %err		Reported Value
07-Apr-2021	1,902.294	0.011	1.188	1.412	6.439	0.276	1.00265	0.148	
07-Apr-2021	1,898.336	0.011	1.151	1.631	6.397	0.272	1.00152	0.146	1.0021836 ±
07-Apr-2021	1,896.060	0.010	1.205	1.325	6.419	0.313	1.00269	0.165	0.096%
08-Apr-2021	1,896.448	0.009	1.201	1.479	6.395	0.279	1.00170	0.149	0.09078
08-Apr-2021	1,893.733	0.008	1.167	1.580	6.402	0.247	1.00235	0.134	

Table 3: Air Shots and Mass Discrimination Factor

The blank measurements are undertaken with different temperature schedules between 300°C and 1450°C, depending on the degassing behaviour and previous blank measurement results. The

degassing and blank measurement procedure continues until the ratios of ⁴⁰Ar, ³⁸Ar and ³⁶Ar drop to atmospheric ratios, and ³⁹Ar and ³⁷Ar drop below detectable levels. The entire procedure of degassing and blank measurements is repeated at the end of a set of samples. Blanks will be done in-between samples that belong to a set, with reduced steps at 300°C, 1300°C and 1450°C to check isotope levels. In addition, the mass of each sample is calculated so that the volume of gas released from each step overwhelms the volume of gas that may occur in the blank. The table 4 is a representative sequence of measured blank values recorded during a monitoring process.

		0	01			
Temperature	⁴⁰ Ar	³⁹ Ar	³⁸ Ar	³⁷ Ar	³⁶ Ar	⁴⁰ Ar∕ ³⁶ Ar
300	390.4368171	0.905537608	0.225735172	*	1.270363042	307.3427
500	395.5969863	0.774176443	0.200701385	*	1.290189995	306.6192
700	397.2960946	0.708335676	0.243386524	*	1.260730764	315.1316
900	436.6219355	0.649146429	0.246339898	*	1.426302145	306.1216
1100	597.095862	0.641103058	0.371303411	*	1.909715988	312.6621
1300	649.6245349	0.607193377	0.402933796	*	2.099636135	309.3986
1450	1374.433483	0.664379683	0.800095274	*	4.461063725	308.0955

Table 4: Example of the blanks measurements during a sequence of blanks where isotopes were being monitored prior to sample analysis (* => Not Detectable). Temperature is °C.

Data reduction software:

The calculations were done with an adapted version of *Noble* Software (2022, developed and adapted by the Australian National University Argon Laboratory) and all interpretations have been undertaken with *eArgon* (developed and adapted for ANU Argon Laboratory by G.S. Lister).

Reported Data:

The reported data have been corrected for system backgrounds, mass discrimination, fluence gradients and atmospheric contamination. GA1550 standards were analysed, and an exponential best fit was then used for the calculation of the J-factor and J-factor uncertainty (Table 5).

Sample Name	J-Factor ± %unc	J-Factor ± %uncertainty		Mass Discrimination Factor ± %uncertainty		
NSWSPJG0075 (G1)	1.53061E-3	0.2203	1.00218	0.096	11-Apr-2021	
NSWSPJG0075 (G2-3)	1.53126E-3	0.2202	1.00218	0.096	14-Apr-2021	

Samples J-Factor, Mass Discrimination, and uncertainties:

Table 5: Sample analysis and calculation details

⁴⁰Ar/³⁹Ar isotopic data of the samples are supplied in the Excel Data Tables, which include details on the heating schedule, Argon isotopes abundances and their uncertainty levels, %Ar*, ⁴⁰Ar*/³⁹Ar(K), Cumulative ³⁹Ar%, calculated age and its uncertainty, Ca/K, Cl/K, J-Factor and its uncertainty. Noting that all the reported uncertainties are at one sigma level and the fractional uncertainties are shown as % in the headings of the appropriate columns of data tables. The components involved in the calculation of the uncertainties are listed in Table 6.

Uncertainty of:	Components involved in the calculation
Isotono Abundancos	Uncertainty of isotope measurement
Isotope Abundances	Uncertainty of Mass Discrimination Factor (except for ³⁹ Ar)
	Uncertainty of ⁴⁰ K Decay Constant
J-Factor	Uncertainty of Age of the Flux monitor
	Uncertainty of Flux monitor isotopes abundances

	Uncertainty of Isotopes Abundances J-Factor value and uncertainty of J-Factor
Calculated Age	⁴⁰ K Decay Constant value and uncertainty
	of ⁴⁰ K Decay Constant

Table 6: Components involved in the calculation of each uncertainty

References:

- Forster, M.A. and Lister, G.S. 2004. The interpretation of 40Ar/39Ar apparent age spectra produced by mixing: application of the method of asymptotes and limits. Journal of Structural Geology 26, 287–305
- Forster, M.A. and Lister, G.S. 2009. Core-complex-related extension of the Aegean lithosphere initiated at the Eocene-Oligocene transition. Journal Geophysical Research, 114, B02401.
- Kondev, F.G. and Naimi, S. 2017. The NUBASE2016 evaluation of nuclear properties. Chinese physics C, 41(3), p.030001.
- Lee, J.Y., Marti, K., Severinghaus, J.P., Kawamura, K., Yoo, H.S., Lee, J.B. and Kim, J.S. 2006. A redetermination of the isotopic abundances of atmospheric Ar. Geochimica et Cosmochimica Acta, 70(17), 4507-4512.
- McDougall, I. and Harrison, T.M. (Eds.). 1999. Geochronology and Thermochronology by the 40Ar/39Ar Method, 2nd ed., 269 pp. Oxford Univ. Press, New York.
- Renne, P.R., Balco, G., Ludwig, K.R., Mundil, R. and Min, K., 2011. Response to the comment by WH Schwarz et al. on "Joint determination of 40K decay constants and 40Ar*/40K for the Fish Canyon sanidine standard, and improved accuracy for 40Ar/39Ar geochronology" by PR Renne et al.(2010). Geochimica et Cosmochimica Acta, 75(17), pp.5097-5100.
- Spell, T. L. and I. McDougall. 2003. Characterization and calibration of 40Ar/39Ar dating standards. Chemical Geology, 198, 189–211.
- Tetley, N., McDougall, I. & Heydegger, H.R. 1980. Thermal neutron interferences in the 40Ar/39Ar dating technique. Journal Geophysical Research, 85, 7201–7205.