

1
2
3
4
5
6
7
8
9
10
11 **Analysis of high-precision vanadium isotope ratios by**
12 **medium resolution MC-ICP-MS**
13
14
15
16
17
18
19
20
21
22

23 Sune G. Nielsen^{a,b,*}, Jeremy D. Owens^{a,b,1} and Tristan J. Horner^{a,c}
24
25
26
27
28
29
30
31
32
33
34

35 ^aNIRVANA laboratories, Woods Hole Oceanographic Institution, Woods Hole, MA, USA
36

37 ^bDepartment of Geology and Geophysics, Woods Hole Oceanographic Institution, Woods Hole,
38 MA, USA
39
40
41

42 ^cDepartment of Marine Chemistry and Geochemistry, Woods Hole Oceanographic Institution,
43 Woods Hole, MA, USA
44
45
46
47
48

49 *Corresponding author
50

51 ¹Now at: Department of Earth, Ocean and Atmospheric Science, Florida State University,
52 Tallahassee, FL 32306, USA
53
54
55
56
57
58
59
60

1
2
3 **Abstract** - We present and verify a new method to measure vanadium isotope ratios using a
4 Thermo Scientific Neptune multi-collector inductively-coupled plasma mass spectrometer (MC-
5 ICP-MS) operated in medium mass resolution mode. We collect masses 48 through 53
6 simultaneously using the L2, L1, Center, H1, H2 and H3 collectors. The Center cup is equipped
7 with a $10^{12} \Omega$ resistor, H1 is equipped with a $10^{10} \Omega$ resistor, while the rest of the collectors have
8 standard $10^{11} \Omega$ resistors. Unlike previous low-resolution methods, the use of medium mass
9 resolution ($\Delta M/M \sim 4,000$) permits separation of V, Ti and Cr isotopes from all interfering
10 molecular species representing combinations of C, N, O, S, Cl, and Ar. We show that the
11 external reproducibility follows a power law function with respect to the number of V^+ ions
12 collected and achieve an external reproducibility of $\pm 0.15 \%$ with total V^+ ion beam intensities
13 of ~ 1 nA. The separation of interfering molecular species from the V mass spectrum reduces the
14 V requirement for precise isotope data to as little as 200-300 ng V per analysis — a reduction of
15 $\sim 90\%$ compared with previous methods — making several low-V matrices amenable to V
16 isotope analysis.
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

1. Introduction

At chemical equilibrium, the stable isotope fractionation between reactants and products depends on the bond strengths of the molecules on both sides of the reaction¹. Equilibrium stable isotope fractionation is often greatest for chemical reactions that involve several oxidation states of an element because the bond strengths tend to vary systematically as a function of oxidation state. Therefore, vanadium (V) isotope measurements have great potential as tracers of a wide range of reduction-oxidation (redox) processes because of the four naturally occurring valence states (+2, +3, +4, and +5) in different environments on Earth, and potentially other bodies in the solar system²⁻⁶. Vanadium has two stable isotopes (⁵⁰V and ⁵¹V) with abundances of 0.24 % and 99.76%, respectively. Theoretical calculations predict that V isotope fractionation between molecules of V in different oxidation states should be up to several permil at Earth surface temperatures⁷, but only recently was a method developed that is sufficiently precise and accurate to resolve the predicted natural V isotope variation^{8,9}. Subsequently it has been shown that even rocks formed at temperatures in excess of 1000K display resolvable V isotope differences¹⁰, which underlines the potential of V isotopes to track redox processes both in high and low temperature environments.

As is common to other metal isotope systems¹¹, measuring V isotope ratios with high precision and accuracy requires 1) efficient and quantitative separation of the element of interest from sample matrices be obtained and 2) sufficient ion beam signal is produced for the minor (⁵⁰V) isotope in order to achieve acceptable counting statistics for collection of high precision isotope data. Previously described methods for high precision V isotope measurements meet both of these criteria^{8,9} using liquid ion-exchange chromatography to remove isobaric Ti and Cr from the sample matrix whilst ensuring quantitative V recovery⁸. The mass spectrometric protocols of

1
2
3 the original V isotope measurement method⁸, utilizes a $10^9 \Omega$ (Ohm) resistor on the faraday
4 detector that collects ^{51}V , which allows larger amounts of V to be introduced into the multi-
5 collector inductively-coupled plasma mass spectrometer (MC-ICP-MS). The larger quantities of
6 V yield sufficient ^{50}V signals on a standard $10^{11} \Omega$ resistor for collection of high precision data.
7
8
9

10
11
12 However, there are two major drawbacks to the method that uses 10^9 and 10^{11} resistors,
13 which can affect the sample diversity that can be investigated and potentially the data quality.
14 First, due to the use of the $10^9 \Omega$ resistor, it was found that at least $2 \mu\text{g}$ of V was consumed for
15 each V isotope analysis⁸. Although many rocks and sediments on Earth contain several hundred
16 $\mu\text{g/g}$, there are also many environments that are characterized by much lower V abundances,
17 which makes separation of sufficient amounts of V for isotope analysis impractical or impossible
18 with current chemical separation protocols. Secondly, the mass spectrum collected for vanadium
19 isotope measurements contains a large array of interfering molecular species (Table 1), which
20 cannot be individually monitored and corrected for in low resolution (LR, low resolution $\Delta\text{M}/\text{M}$
21 ~ 300). The molecules listed in Table 1 require a mass resolving power of $\Delta\text{M}/\text{M} = 1900\text{-}3200$
22 and, therefore, it has to be assumed that these molecular interferences are either so small that
23 they do not affect V isotope measurements, or that they are effectively constant between
24 measuring samples and standards. While these assumptions are likely correct for many types of
25 sample, it is exceedingly difficult to identify small (i.e. sub permil) effects from molecular
26 interferences because there is no independent way to monitor them.
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47

48 Here, we present a new mass spectrometric method using medium resolution mode (MR,
49 $\Delta\text{M}/\text{M} \sim 4,000$) of a Thermo Scientific Neptune MC-ICP-MS. The setup presented is able to
50 resolve all interfering molecular species in the V mass spectrum, thus removing the uncertainty
51 of minor isotope effects from combinations of polyatomic C, N, O, S, Ar and Cl. The method
52
53
54
55
56
57
58
59
60

1
2
3 utilizes high-transmission interface cones that significantly lower the amount of V required per
4 analysis to roughly 200-300ng, or almost an order of magnitude less than previous methods.
5
6
7
8
9

10 **2. Mass spectrometry protocols**

11
12 Vanadium isotope compositions are measured using a Thermo Scientific Neptune MC-ICP-
13 MS. The front-end is configured with a 'Jet' sampler cone and an 'X'-type skimmer cone in order
14 to obtain the highest possible V transmission efficiency. Vanadium is introduced into the mass
15 spectrometer as a dry aerosol by passing the sample through a CETAC Aridus II desolvator,
16 which is fed by an ESI perfluoroalkoxy alkane (PFA) nebulizer that aspirates the sample solution
17 at ~120 μ l/min. This configuration routinely produces a vanadium ion beam of ~2.5 nA for a
18 solution containing 1 μ g/g vanadium when the mass spectrometer is operated in MR, equivalent
19 to a total ion transmission of ~0.7%.
20
21
22
23
24
25
26
27
28
29
30

31 We collect masses 48 through 53 simultaneously using the L2, L1, Center, H1, H2 and H3
32 collectors (Table 2). The Center cup is equipped with a 10^{12} Ω resistor, H1 is equipped with a
33 10^{10} Ω resistor, while the rest of the collectors are connected to standard 10^{11} Ω resistors. The
34 setup produces a mass spectrum as shown in Fig. 1a.
35
36
37
38
39
40

41 Vanadium isotope measurements are otherwise performed according to the protocols
42 outlined by Nielsen et al.⁸. Each sample analysis consists of 30s background that is measured by
43 defocusing and deflecting the ion beam away from the detectors in the electrostatic analyzer,
44 followed by 40 cycles of 4.194 s integrations. Correction for interferences on ^{50}V from ^{50}Ti and
45 ^{50}Cr that cannot be resolved in MR (nor high resolution) mode are performed as outlined in
46 Nielsen et al.⁸. Each sample is bracketed by four (two on each side) measurements of the Alfa
47 Aesar (AA-V) standard that has been defined as $\delta^{51}\text{V} \equiv 0$ (where $\delta^{51}\text{V} = 1000 \times [(^{51}\text{V}/^{50}\text{V})_{\text{sample}} -$
48
49
50
51
52
53
54
55
56
57
58
59
60

1
2
3 $^{51}\text{V}/^{50}\text{V}_{\text{AA-V}})/^{51}\text{V}/^{50}\text{V}_{\text{AA-V}}]$). In addition, each sample measurement is bracketed by two
4
5 measurements of the BDH vanadium solution that has an isotope composition of $\delta^{51}\text{V} = -1.19 \pm$
6
7 0.12‰ (2sd)⁸.
8
9

10 We use an autosampler to perform long sequences of V isotope measurements, which serves
11
12 to minimize uncertainty that may arise from systematic drift in the instrumental mass bias. Each
13
14 sample and standard measurement are always spaced by the exact same amount of time, enabling
15
16 a time-dependent correction of instrumental mass bias.
17
18
19

20 21 22 **3. Tests of mass spectrometric method performance** 23

24 In order to investigate the accuracy and precision of the MR protocols we designed a series
25
26 of tests to determine the sensitivity of measured $^{51}\text{V}/^{50}\text{V}$ ratios to several instrumental
27
28 parameters, which are addressed in the following sections:
29
30

- 31 1. The effect of measuring at different m/z to assess the proportion of the peak flat that
32 is free from interferences
33
- 34 2. The effect of (mis)-matching V concentrations between samples and standard
35 solutions
36
- 37 3. The addition of sulfur to samples and artifacts from S-O molecular interferences
38
- 39 4. The optimal V concentration—in terms of precision—at which to perform V isotope
40 measurements to obtain data with precision comparable to that of previously
41 described LR methods
42
43
44
45
46
47
48
49

50 It should be noted that these tests were not intended to investigate effects resulting from
51
52 inadequate V separation from sample matrix during liquid ion exchange chromatography, as
53
54 these are discussed in detail elsewhere⁹.
55
56
57
58
59
60

3.1. Width of peak flat and magnet stability

The width of the peak flat obtained in MR is approximately 0.01 atomic mass units (amu) (Fig. 1a). We position the magnet approximately 1/3 of the way from the left shoulder and keep this mass setting throughout each run sequence. The magnet is positioned towards lower m/z to avoid any potential effects from tailing of interference peaks¹² that all possess higher m/z . After each sequence is completed we scan across the mass spectrum to verify that the magnet has not drifted during the run. So far we have not observed the magnet to drift by more than 0.002 amu (20 % of peak width) during sequences of at least 12 hours. We are thus confident that magnet stability is sufficient to avoid effects from magnet drift. As expected, our tests of the peak flat (Fig. 1b and 1c) also showed that tailing from interferences on ^{50}V that cause a systematic decrease in the measured $^{51}\text{V}/^{50}\text{V}$ ratios are the most significant sources of deviation from peak flatness on the high m/z side of the peak flat. However, these interference peak tails appear to be similar for standards and samples because no change in the relative isotopic difference between BDH and AA-V was recorded over the entire peak flat (Fig. 1c).

3.2. Effects from variations in sample and standard V concentrations

As outlined above, a host of molecular interferences are collected together with the main isotopes of Ti, V and Cr during LR measurements (Table 1). Assuming that these molecular interferences are constant for samples and standards, it was determined that the V concentrations of samples and bracketing standards had to be matched to better than 15%^{8, 9} in order to normalize away the molecular interferences. However, all molecular interferences across the V mass spectrum are efficiently resolved in MR, suggesting that V isotope ratios should be

1
2
3 insensitive to this issue as long as V is effectively isolated from the sample matrix. To test if this
4
5 was the case, we performed two tests using (i) matrix-free AA-V standards of differing
6
7 concentrations and (ii) sample seawater matrix-doped AA-V that was subsequently processed
8
9 through previously published ion exchange chromatographic procedures⁸. The latter tests were
10
11 conducted to ascertain if any of the molecular interferences were dependent on residual matrix
12
13 from geological samples.
14
15

16
17 The results of the tests with a seawater matrix are shown in Figure 2, illustrating a small but
18
19 measureable effect for large mismatches in concentration between samples and standards. The
20
21 effect correlates with the ratio between the V ion beam intensities of samples and standards ($R^2 =$
22
23 0.71) and shows that >30% mismatch results in a systematic offset in $\delta^{51}\text{V}$ of $\sim 0.15\%$. This
24
25 effect is also apparent for standards that have not been doped with sample matrix (Fig. 1c). As
26
27 such, this effect most likely relates to minor peak tails from interferences on the high m/z side of
28
29 the peak flat that can affect measured $^{51}\text{V}/^{50}\text{V}$ ratios, rather than molecular interferences that are
30
31 present in residual sample matrix. Though these effects are probably smaller than those observed
32
33 for the LR method⁸, we recommend that samples and standards continue to be matched to within
34
35 $\pm 15\%$ in order to eliminate these effects. In addition, it can be seen that the effect is enhanced at
36
37 lower ion beam intensities (Fig. 1c), which suggests that the interference tail(s) originate from
38
39 polyatomic molecular species containing Ar, O, and N, rather than sample matrix. However, the
40
41 effect may still be enhanced by interferences originating from residual sample matrix (e.g. most
42
43 likely S, C, and Cl), but based on our present tests (Fig. 2) there is no indication that these
44
45 contributions are significant. However, it is likely that residual sample matrix left over from V
46
47 separation depends strongly on sample type processed, such that we encourage future studies to
48
49 perform V isotope verification for each type of sample matrix encountered (e.g. seawater,
50
51
52
53
54
55
56
57
58
59
60

1
2
3 organic rich sediments, basalts, granites) that may introduce small, but significant inaccuracies in
4
5 V isotope data.
6
7
8
9

10 3.3. Sulfur addition to standard solutions

11
12 Previous V isotope measurement techniques documented that significant quantities of S in
13 sample solutions greatly modify measured V isotope compositions. This effect is due to the
14 apparent increase in Ti and V ion beams that results from S-O molecular interferences (Table 1).
15
16 When operating the Neptune MC-ICP-MS at Woods Hole Oceanographic Institution (WHOI) in
17 LR, we observed much larger effects on measured V isotope ratios from S addition than was
18 previously observed for the Nu Plasma (Fig. 3). Given that oxide production rates for the Nu
19 Plasma and Neptune MC-ICP-MS instruments are similar¹³, the higher S-O interferences
20 observed for the Neptune is likely due to the higher transmission for the Neptune in the low mass
21 end of the periodic table than the Nu Plasma. Thus, for a solution with a fixed S:V ratio,
22 significantly higher S/V is measured on the Neptune MC-ICP-MS compared to the Nu Plasma.
23
24 In turn, this relationship results in higher abundances of S-O interferences on the Neptune (Fig.
25 3). The published V separation protocols may result in non-trivial amounts of S in the isolated V
26 because no specific part of the chemical separation procedure removes S quantitatively. We
27 therefore tested the effect of S addition to BDH solutions with known $\delta^{51}\text{V}$ of $-1.19 \pm 0.12\text{‰}$
28 (2sd) by adding enough sulfur to obtain $\text{S}/\text{V} \sim 2$. Vanadium isotope measurements in LR with
29 half the S/V ratio revealed offsets from the true value of up to several permil (Fig. 3). However,
30 the results from this test in MR show that - even at low V ion beam intensities - there is no
31 detectable effect on measured V isotope composition (Fig. 4b).
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

3.4. Optimal V concentration for samples and standards

Several of the mass spectrometric tests that use very low V ion beam intensities (Fig. 1 and 2) display precision that is slightly inferior ($\pm 0.2\%$ vs. $\pm 0.12\%$) to that previously achieved for $5\mu\text{g/g}$ solutions in LR mode with a Nu Plasma MC-ICP-MS⁸. We have tested the uncertainty on the BDH standard measurements as a function of ^{51}V ion beam intensity and find that individual sequences produce uncertainties on repeat BDH measurements better than or equal to $\pm 0.12\%$ only when ^{51}V ion beam intensity is greater than ~ 1.5 nA (Fig. 4a). In addition, the uncertainty generally follows a power law whereby higher ion beam intensities are associated with lower uncertainties. However, the slope of the power law we derive from measurements of BDH is approximately -0.9 (Fig. 4a), which is significantly steeper than the slope of -0.5 expected if all uncertainties were explained by counting statistics¹⁴. Similar to effects observed for iron isotope measurements using standard-sample bracketing¹⁴, we conclude that uncertainties on V isotope measurements have a significant component from other sources of uncertainty, such as instability of the plasma, that are most important at low ion beam intensities. These uncertainties only apply to pure vanadium standard solutions and are likely slightly larger for samples, as has been previously reported^{9, 10}. However, with our routine running conditions where we obtain V ion currents of ~ 1.5 nA total V for solutions containing $0.6\mu\text{g/g}$ vanadium, we are able to achieve $\pm 0.12\%$ (Fig. 4), which consumes only $\sim 260\text{ng}$ of V per analysis (30 sec background, 20 sec idle time, 168 sec analysis time). This amount of V is almost an order of magnitude less than required by the previous LR method using a Nu Plasma MC-ICP-MS. Two of the major reasons for this reduction in sample-derived V is that (i) previous methods utilized low-transmission sampler and skimmer cones to minimize the impact of molecular interferences across the vanadium mass spectrum⁸ and (ii) previous methods used tuning parameters that were set to

1
2
3 minimize molecular interferences, by employing low Ar gas flows across the desolvating
4 membrane, and utilizing high RF power to break down molecular interferences. Both of these
5 tuning parameters result in variably lower V transmission than if tuning for optimum V
6 transmission efficiency as we do here.
7
8
9
10
11

12 13 14 15 **4. Conclusions**

16
17 We have developed and verified a new medium resolution mass spectrometric protocol to
18 measure high precision V isotope ratios using a Thermo Scientific Neptune MC-ICP-MS. The
19 method achieves a comparable precision comparable to previous LR methods of $\pm 0.12\%$, while
20 consuming as little as 260 ng of V—almost an order of magnitude less V per analysis. Given the
21 practical limits of ion-exchange chromatographic separation procedures that cannot more than
22 100mg of matrix per resin column, this ~90 % reduction in V requirements by using MR will
23 significantly aid future studies in obtaining precise V isotope data for samples with low V
24 concentrations that were otherwise difficult or impractical to measure. For example, this new
25 method makes many types of previously inaccessible samples with low V contents amenable to
26 V isotope analysis, such as seawater, depleted peridotites, iron meteorites, carbonates, and
27 biomass, which are all of significant interest across the fields of Earth and Environmental
28 Sciences.
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47

48 49 **5. Acknowledgements**

50 We thank Dr. Jerzy Blusztajn for his tireless help in the WHOI plasma mass spectrometer
51 facility. This study was funded by a WHOI IR&D grant to SGN and NSF OCE grant 1434785 to
52 JDO and SGN and Agouron Geobiology Fellowships to JDO and TJH.
53
54
55
56
57
58
59
60

6. References

1. H. C. Urey, *J. Chem. Soc.*, 1947, 562-581.
2. J. L. Morford and S. Emerson, *Geochim. Cosmochim. Acta.*, 1999, **63**, 1735-1750.
3. B. J. Wood, J. Wade and M. R. Kilburn, *Geochim. Cosmochim. Acta.*, 2008, **72**, 1415-1426.
4. K. Righter, S. R. Sutton, M. Newville, L. Lei, C. S. Schwandt, H. Uchida, B. Lavina and R. T. Downs, *American Mineralogist*, 2006, **91**, 1643-1656.
5. B. Wehrli and W. Stumm, *Geochim. Cosmochim. Acta.*, 1989, **53**, 69-77.
6. M. D. Lewan and J. B. Maynard, *Geochim. Cosmochim. Acta.*, 1982, **46**, 2547-2560.
7. F. Wu, T. Qin, X. F. Li, Y. Liu, J. H. Huang, Z. Q. Wu and F. Huang, *Earth Planet. Sci. Lett.*, 2015, **426**, 216-224.
8. S. G. Nielsen, J. Prytulak and A. N. Halliday, *Geostand. Geoanal. Res.*, 2011, **35**, 293-306.
9. J. Prytulak, S. G. Nielsen and A. N. Halliday, *Geostand. Geoanal. Res.*, 2011, **35**, 307-318.
10. J. Prytulak, S. G. Nielsen, D. A. Ionov, A. N. Halliday, J. Harvey, K. A. Kelley, Y. Niu, D. W. Peate, K. Shimizu and K. W. W. Sims, *Earth Planet. Sci. Lett.*, 2013, **365**, 177-189.
11. C. M. Johnson, B. L. Beard and F. Albarede, *Rev Mineral Geochem*, 2004, **55**, 1-24.
12. S. Weyer and J. Schwieters, *Int. J. Mass Spectrom.*, 2003, **226**, 355-368.
13. J. L. Payne, N. J. Pearson, K. J. Grant and G. P. Halverson, *J. Anal. At. Spectrom.*, 2013, **28**, 1068-1079.
14. S. G. John and J. F. Adkins, *Mar. Chem.*, 2010, **119**, 65-76.

Figure captions

Figure 1: a) Mass scan of the AA-V standard in medium resolution mode across the interference free portion of the mass spectrum measured. Note that $m/z = 49$ and $m/z = 53$ are not shown because they are so small for AA-V that the ion beams are primarily noise. b) Vanadium isotope measurements of the AA-V standard across the portion of the mass spectrum in a) (from 49.92 to 49.93 amu) that visually appears free of interferences. Measurements were performed using four different ion beam intensities. Peak tails of molecular interferences on ^{50}V cause the lower $^{51}\text{V}/^{50}\text{V}$ recorded towards the right side of the mass spectrum. c) AA-V standard solutions of different vanadium ion beam intensities measured against an AA-V solution that yielded ~ 0.67 nA ^{51}V across the portion of the mass spectrum in a) (from 49.92 to 49.93 amu) that visually appears free of interferences. Also shown are measurements of the BDH vanadium solution against AA-V across the same portion of the mass spectrum. The ion beam intensity of the BDH and bracketing AA-V solutions were ~ 0.67 nA and were matched to within 4%.

Figure 2: Vanadium isotope compositions measured for four separate seawater matrix-doped splits of AA-V plotted against the ratio of the ion beam intensity for the processed AA-V and the bracketing unprocessed AA-V standards. Shading denotes the areas where samples and standards are matched to better than $\pm 30\%$ and V isotope compositions are within $\pm 0.15\%$ of the true value ($\equiv 0 \text{ ‰}$). The seawater matrix was removed from each AA-V split using ion exchange chromatography. Different colored symbols denote different splits of AA-V that were put through the column chemistry separation procedure. When sample ion beam intensities are more than 30% different to the bracketing standards, V isotope compositions recorded deviate from the

1
2
3 true value by more than $\pm 0.15\%$. This effect is similar to that observed for pure standards (Fig.
4
5
6 1c) suggesting that the V isotope artifact does not originate from residual sample matrix.

7
8 Figure 3: Vanadium isotope compositions measured for BDH vanadium solutions doped
9
10 with variable amounts of sulfur plotted against the recorded ratio of $m/z - 49/ m/z - 51$.
11
12 Measurements performed in both low and medium resolution on the Neptune MC-ICP-MS are
13
14 displayed. Also shown are similar tests performed on the Nu Plasma⁸ albeit with significantly
15
16 greater amounts of sulfur added. Grey bars denote the value with 2sd error envelope of undoped
17
18 BDH vanadium solutions.
19
20

21
22 Figure 4: a) Two standard deviation uncertainties on repeat measurements of BDH
23
24 vanadium solutions in medium resolution as a function of the vanadium ion beam intensity. Each
25
26 data point represents between 26 and 43 individual analyses of BDH. b) Vanadium isotope
27
28 composition of six repeat analyses of BDH vanadium solution doped with sulfur (red circle; S/V
29
30 ~ 2). Error bars are 2sd. Also shown are the undoped BDH vanadium measurements plotted in
31
32 a). The grey envelope denotes the fitted uncertainty from a) associated with measurements of
33
34 undoped BDH vanadium solutions in medium resolution.
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

Table 1: Molecular species resolved in medium resolution

Molecular species*	Species abundance	Reference isotope	$\Delta M/M$
$^{32}\text{S}-^{16}\text{O}_6$	0.94793	^{48}Ti	2518
$^{36}\text{Ar}-^{12}\text{C}$	0.00333	^{48}Ti	2446
$^{36}\text{S}-^{12}\text{C}$	0.00019	^{48}Ti	2506
$^{34}\text{S}-^{14}\text{N}$	0.04194	^{48}Ti	2085
$^{33}\text{S}-^{15}\text{N}$	0.00002	^{48}Ti	2030
$^{35}\text{Cl}-^{13}\text{C}$	0.00833	^{48}Ti	1976
$^{37}\text{Cl}^{12}\text{C}$	0.23954	^{49}Ti	2715
$^{33}\text{S}^{16}\text{O}$	0.00748	^{49}Ti	2646
$^{34}\text{S}^{15}\text{N}$	0.00015	^{49}Ti	2435
$^{36}\text{Ar}^{13}\text{C}$	0.00004	^{49}Ti	2125
$^{32}\text{S}^{17}\text{O}$	0.00036	^{49}Ti	2098
$^{35}\text{Cl}^{14}\text{N}$	0.75503	^{49}Ti	2035
$^{38}\text{Ar}^{12}\text{C}$	0.00062	^{50}V	3208
$^{34}\text{S}^{16}\text{O}$	0.04200	^{50}V	3198
$^{35}\text{Cl}^{15}\text{N}$	0.00278	^{50}V	2291
$^{37}\text{Cl}^{13}\text{C}$	0.00266	^{50}V	2261
$^{36}\text{S}^{14}\text{N}$	0.00020	^{50}V	2172
$^{36}\text{Ar}^{14}\text{N}$	0.00336	^{50}V	2129
$^{32}\text{S}^{18}\text{O}$	0.00190	^{50}V	2075
$^{35}\text{Cl}^{16}\text{O}$	0.75600	^{51}V	2572
$^{34}\text{S}^{17}\text{O}$	0.00002	^{51}V	2212
$^{36}\text{Ar}^{15}\text{N}$	0.00001	^{51}V	2150
$^{37}\text{Cl}^{14}\text{N}$	0.24131	^{51}V	2037
$^{33}\text{S}^{18}\text{O}$	0.00002	^{51}V	1911
$^{36}\text{S}^{16}\text{O}$	0.00020	^{52}Cr	2418
$^{40}\text{Ar}^{12}\text{C}$	0.98504	^{52}Cr	2375
$^{36}\text{Ar}^{16}\text{O}$	0.00336	^{52}Cr	2366
$^{38}\text{Ar}^{14}\text{N}$	0.00063	^{52}Cr	2053
$^{37}\text{Cl}^{15}\text{N}$	0.00089	^{52}Cr	2037
$^{34}\text{S}^{18}\text{O}$	0.00008	^{52}Cr	1959
$^{35}\text{Cl}^{17}\text{O}$	0.00029	^{52}Cr	1891
$^{37}\text{Cl}^{16}\text{O}$	0.24162	^{53}Cr	2626
$^{40}\text{Ar}^{13}\text{C}$	0.01096	^{53}Cr	2111
$^{35}\text{Cl}^{18}\text{O}$	0.00152	^{53}Cr	1935

*Only molecular species with an abundance of more than 0.001% of the total species considered

Table 2: Isotope abundances in % of measured Ti, V and Cr isotopes

Mass (amu)	48	49	50	51	52	53
Ti	73.8	5.51	5.34			
V			0.24	99.76		
Cr			4.31		83.76	9.55
Resistor (Ω)	10^{11}	10^{11}	10^{12}	10^{10}	10^{11}	10^{11}







