

1 **Design, construction, automation, and first-principles calibration of a**  
2 **low volume He measurement line**

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7 **Abstract.** A noble-gas analysis line capable of accurate and precise measurements of small absolute amounts of <sup>4</sup>He released  
8 from crystals is a key analytical step in the production of (U-Th)/He chronologic data. He analysis lines that are custom-built  
9 in-house can be optimized for specific lab needs and facilitate continued maintenance, repair, and upgrades. However, there is  
10 little information in the published literature about the methods and approaches for building a He line. Here, we describe the  
11 design, construction, automation, and metrological calibration of a custom <sup>4</sup>He extraction and analysis line as part of  
12 establishing laser-ablation (U-Th)/He methods in the University of Colorado Thermochronology Research and Instrumentation  
13 Lab (CU TRaIL). The line, called the Jimbochron, is designed to precisely measure very small (~fmol) amounts of <sup>4</sup>He while  
14 being fully automated and easily modifiable in the future. These goals are achieved by minimizing the line volume, adopting  
15 a unique double-hexagonal manifold configuration, installing a high-sensitivity quadrupole mass spectrometer, and developing  
16 LabView code for instrument communication and automation that is open and straightforward to update. We also explain the  
17 steps used to calibrate the Jimbochron metrologically from first principles with a new in-house calibration volume to ensure  
18 high-accuracy He measurements. The Jimbochron accurately and precisely measures small gas volumes (<0.1 fmol) and now  
19 routinely generates accurate and precise He data for laser-ablation (U-Th)/He applications.

20 **1 Introduction**

21 (U-Th)/He chronology is a versatile radiometric dating method based on the accumulation of He in geologic materials produced  
22 by the alpha decay of U, Th, and Sm. Calculating a (U-Th)/He date requires measuring the absolute amounts of both the  
23 radiogenic decay isotope (<sup>4</sup>He) and the radioactive parent isotopes (<sup>238</sup>U, <sup>235</sup>U, <sup>232</sup>Th, and <sup>147</sup>Sm) present in the material. The  
24 precision and accuracy of these measurements are first-order controls on the precision and accuracy of the resulting date. U,  
25 Th, and Sm measurements are often performed using inductively-coupled plasma mass spectrometry, which is incapable of  
26 measuring noble gases. Therefore, <sup>4</sup>He measurements are typically performed using noble gas mass spectrometry, where a gas  
27 source mass spectrometer is attached to a vacuum sample extraction line.

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29 Although high-quality noble gas analysis lines are available commercially and can be a good choice for labs seeking to  
30 efficiently implement well-established gas measurement methods, these instruments may not be ideal for researchers interested

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37 in developing new and emerging techniques. Commercial machines with control software that is proprietary or otherwise  
38 inaccessible are not easily modified, customized, and optimized. These limitations can make it difficult and expensive for  
39 researchers to innovate and refine analytical approaches as techniques evolve. This restricted flexibility also hinders repairing  
40 and replacing hardware components as they fail and complicates updating software and operating systems to maintain  
41 compliance with institutional security requirements.

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43 These challenges can be circumvented with He gas analysis lines that are designed and constructed in-house. In 2018, we  
44 decided to build a custom He measurement line specifically optimized for laser-ablation, in situ He analyses as part of  
45 expanding the analytical capabilities of the University of Colorado Thermochemistry Research and Instrumentation Lab (CU  
46 TRaIL) to include laser-ablation (U-Th)/He chronology. Theoretically, the He analyses could have been performed on our  
47 existing ASI (Australian Scientific Instruments) Alphachron<sup>®</sup> He analysis line, which has been continuously and reliably  
48 producing conventional whole-grain He data for over a decade. However, we do not have the free machine time on the  
49 Alphachron<sup>®</sup> for the experiments required to fully develop new analytical protocols for laser-ablation He analysis. Moreover,  
50 the Alphachron's<sup>®</sup> inaccessible control software makes it difficult to implement new analytical routines with new components  
51 (e.g., an excimer laser). Therefore, we designed and constructed a new He analysis line optimized for high-precision, low  
52 volume, laser-ablation He analyses, automated the line using custom LabView<sup>®</sup> software, and calibrated it metrologically  
53 using a new in-house calibration volume. To distinguish this He line from the Alphachron<sup>®</sup>, the new custom line was named  
54 the Jimbochron.

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56 Although some labs have personnel with the expertise and skills required to build and commission a He analysis line, little  
57 easily accessible information on this topic is available. This creates a situation where valuable information is both difficult to  
58 learn and important institutional knowledge can be lost when experienced scientists retire. The geochronology,  
59 thermochronology, and mass spectrometry communities benefit from sharing the diverse approaches and methods that can be  
60 used to design and calibrate analysis lines, as done in important previous contributions (e.g. Gautheron et al., 2021; Lanphere  
61 and Dalrymple, 2000; Lesnek et al., 2024; Morgan et al., 2011; Ritter et al., 2021; Yakubovitch et al., 2024). Here, we provide  
62 an overview of the steps along our path from deciding to build the Jimbochron to its routine generation of precise and accurate  
63 He measurements in the CU TRaIL. Our goal is to share the design principles (Sect. 2), construction information and parts  
64 suppliers (Sect. 3), software (Sect. 4), and calibration methods (Sect. 5) necessary to build and commission a He analysis  
65 line.

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68 **2 Design principles and considerations**

69 **2.1 Design**

70 The primary motivation for the overall physical design of the Jimbochron is to minimize the internal volume of the vacuum  
71 line in order to maximize the partial pressure of the small amounts of <sup>4</sup>He evolved from laser ablation pits (often ~1 fmol).  
72 The gas evolved from a sample will expand into the available volume, so a smaller expansion volume means a higher He  
73 concentration in the line, and therefore a larger He signal at the mass spectrometer. Larger signals are more stable and easier  
74 to measure on mass spectrometers. Additionally, smaller volumes are easier to evacuate than larger volumes, which means  
75 lower He blanks in the analysis line can be attained more quickly. The arrangement of the main line skeleton influences the  
76 total volume that sample gas must expand into because it dictates the total length of [steel](#) tubing required to connect all of the  
77 necessary components.

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79 A second important design motivation is a He analysis line that can be modified as easily as possible as the needs of the lab  
80 evolve. Such a system allows for future alterations, additions, and improvements without significant deconstruction,  
81 recalibration, or re-design of the instrument.

82  
83 To achieve the goals of an adaptable line with minimized internal volume, we designed a unique configuration in which  
84 pneumatic valves are arranged around two hexagonal manifolds (vacuum chambers with multiple openings; Fig. 1). This  
85 design allows a large number of valves, and therefore line components, to be arranged closely together using a relatively small  
86 total length of [steel](#) tubing, therefore minimizing the total internal volume. The hexagonal manifolds also allow for easy  
87 rearrangement of hardware, and have unused and available ports that don't significantly increase internal volumes but are  
88 available for more components (e.g., diode laser, vacuum pumps, additional getters) to be added in the future.

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90 We also prioritized selecting a mass spectrometer with high sensitivity. He measurements on the Jimbochron are performed  
91 by quadrupole mass spectrometry because many commercially available quadrupole mass spectrometers (QMSs) are easy to  
92 install, simple to maintain, affordable, and more than sensitive enough to accurately and reproducibly measure the gases  
93 necessary for (U-Th)/He analyses. Although magnetic sector mass spectrometers have considerably more resolving power,  
94 their cost and more difficult maintenance have made QMSs more common in (U-Th)/He labs. As explained in more detail in  
95 Sect. 3.6, the Jimbochron uses a QMS that is approximately an order of magnitude more sensitive than the QMS on our existing  
96 Alphachron® He system.

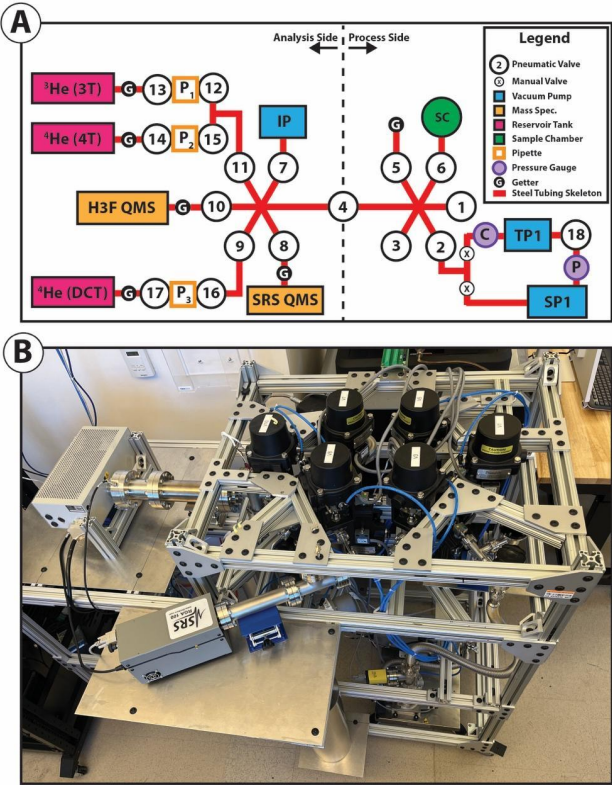
97  
98 In addition, developing flexible and accessible software for instrument control and automation was a central focus during  
99 Jimbochron development. Code that can be easily modified is crucial for replacing parts on the line, for updating the operating  
100 system, and for modifying protocols as our understanding of methods evolve. In addition, full automation of He analysis

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104 routines is required for efficient and reproducible noble gas analysis. For example, creating He maps of mineral grains similar  
105 to those of Danišik et al., (2017) is time consuming, and if done at any significant scale realistically requires automation.



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107 Figure 1: (A) Schematic diagram of the Jimbochron highlighting key components and connections. The line is conceptually divided  
108 into a "Process Side" which is exposed to relatively high pressures of atmospheric gas during sample exchange, and the "Analysis  
109 Side" which only regularly sees gas that has been gettered and cleaned. The three vacuum pumps (in blue) are the Ion Pump (IP),  
110 Turbo Pump (TP), and Scroll Pump (SP). The three reservoir tanks (in pink) are the 3T tank that contains  $^3\text{He}$ , and the 4T tank and  
111 DCT (depletion-correction tank) that contain  $^4\text{He}$ . The tanks each have a getter and are attached to three separate pipettes labeled  
112 P1, P2, and P3. The two quadrupole mass spectrometers (QMS, solid orange boxes) are the Hidden 3F<sup>®</sup> (labelled H3F) and the  
113 Stanford Research Systems RGA100<sup>®</sup> (labelled SRS). The purple circles are the Pirani (P) and Cold Cathode (C) pressure gauges.  
114 (B) Picture of the Jimbochron in a similar orientation to the schematic diagram in (A). Note many of the valves are mounted with  
115 the actuators facing down and not visible in the picture.

116 2.2 Achieving and maintaining ultra-high vacuum conditions

117 Noble gas analysis typically requires measurement of sample gas in an ultra-high vacuum (UHV;  $<1 \times 10^{-8}$  torr) environment  
118 to minimize contamination from atmospheric gases. Although atmospheric He is not abundant (~5 ppm in dry air), the amount  
119 of He contained in only ~0.01 uL of air is roughly equivalent to that contained in a typical apatite crystal. It therefore is  
120 necessary to evacuate an analysis line to as low a pressure as possible. Lines typically achieve and maintain ultra-high vacuum  
121 conditions using multiple vacuum pumps, getters, and all-metal bakeable components.

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123 Conceptually, the Jimbochron can be divided into the process and analysis sides of the line (Fig. 1). The purpose of segregating  
124 the line in this way is to keep the line as clean and at as low a vacuum as possible. The process side is where samples are  
125 loaded and initially put under vacuum, so this side is regularly exposed to atmosphere. It is good practice to minimize the line  
126 volume exposed to atmosphere during sample loading to preclude adsorption of atmospheric gas to internal surfaces. The  
127 analysis side of the line, which includes the mass spectrometer, is only exposed to higher pressures on rare occasions when  
128 significant maintenance (e.g. changing a mass spectrometer filament) is required. Only small volumes of conditioned (see next  
129 section) sample gas are released into the analysis side of the line. The process side of the line is primarily pumped by the turbo  
130 and scroll pumps, whereas the analysis side of the line is primarily pumped by the ion pump (see Sect. 3.5). Pressures measured  
131 by the mass spectrometer in the analysis chamber of the Jimbochron are typically  $<5 \times 10^{-10}$  torr.

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133 The attainment of ultra-high vacuum conditions is enhanced by “baking out” the system at elevated temperatures in order to  
134 release and pump away adsorbed gas. All of the tubing, flanges, and components are safe to bake out to at least 225°C, however  
135 we found 200°C sufficient to reduce pressures to  $<1 \times 10^{-9}$  torr. After construction the Jimbochron was baked out at 200°C  
136 for ~48 hours using heating tapes controlled by an Amptek BT-15<sup>®</sup> benchtop temperature controller and monitored with an  
137 integrated type K thermocouple.

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140 2.3 Gas handling: Liberating, conditioning, and analyzing the gas

141 He lines have three main tasks required to analyze the radiogenic He trapped in geologic materials: gas must be liberated,  
142 conditioned, and analyzed. A common way to liberate the gas trapped in a crystal is with lasers. Long-wavelength lasers heat  
143 the sample to diffuse the gas out of the crystal. Short-wavelength lasers ablate a small part of the sample to break apart the  
144 crystal lattice and release the trapped gas. Vacuum resistance furnaces, induction furnaces, or projector-bulb style diffusion  
145 cells (Farley et al., 1999) can also be used to heat the sample. However, lasers are generally preferred over these other methods  
146 because long-wavelength lasers heat smaller volumes and produce lower He blank levels during whole grain heating (e.g.  
147 House et al., 2000), and short-wavelength lasers can ablate small regions of the sample to measure gas *in situ*. The

152 Jimbochron’s primary approach for liberating the gas from crystals is through laser-ablation with a short-wavelength excimer  
153 laser (an ESL NWR193UC® laser), although a long-wavelength diode laser will be added in the future to enable conventional  
154 whole crystal heating and continual ramped heating techniques.

155

156 After the gas has been liberated, it is expanded into the analysis line and conditioned. Although conditioning can involve a  
157 variety of processes, on the Jimbochron it means that the gas is cleaned and spiked. The Jimbochron cleans the liberated gas  
158 using multiple getters (see Sect. 3.5). Getters are devices that contain highly reactive and porous compounds, often zirconium  
159 alloys or powders, that bond with or trap non-noble gases and effectively sequester them from the rest of the line. The <sup>4</sup>He  
160 liberated from the sample is also spiked with <sup>3</sup>He (see Sect. 5.1). The combined gas is given time to mix and interact with the  
161 getters prior to analysis.

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163 Finally, the <sup>4</sup>He and <sup>3</sup>He are analyzed using a quadrupole noble gas mass spectrometer (Sect. 3.6). The Jimbochron analyzes  
164 the gas in static mode, in which the analyzed gas is isolated from the vacuum pumps during measurement (Reynolds, 1956),  
165 allowing for repeated measurements of very small gas volumes. After analysis, the valves to the vacuum pumps are opened  
166 and the gas is pumped away in preparation for the next sample.

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**Table 1:** Summary of parts and suppliers

| Purpose                          | Item                                                   |
|----------------------------------|--------------------------------------------------------|
| Framing                          | 1.5" Aluminum T-slotted stock                          |
| Roughing pump                    | Agilent IDP3 Dry Scroll Pump®                          |
| Turbo Pump                       | Agilent Twistorr 74 FS®                                |
| Ion Pump                         | Agilent VacIon Plus 20®                                |
| Line Getters                     | SAES Capacitorr CF16-MK2®                              |
| Tank Getters                     | SAES ST172®                                            |
| <u>Hidden QMS Chamber Getter</u> | <u>SAES Sorb-AC®</u>                                   |
| Pipettes                         | Becker Systems® ~0.2 cc all metal                      |
| Reservoir Tanks                  | Custom cylindrical 10L made by NorCal®                 |
| Hexagonal manifolds              | Custom fabricated by MDC®                              |
| Tubing connectors                | Custom fabricated by MDC®                              |
| Pirani pressure gauge            | MKS 925®                                               |
| Cold cathode pressure gauge      | Instrutech WASP WGM701®                                |
| Mass Spectrometer 1              | Hidden HAL21 3F®                                       |
| Mass Spectrometer 2              | SRS RGA100/123®                                        |
| Valves                           | Swagelok SS-4BG®                                       |
| Compressed air manifold          | Festo <u>VTUG-10-MSDR-BIT-25V20-Q8-U-QH 6S-10K+M1®</u> |
| Battery Back-up System           | TrippLite SMART1500LCD®                                |

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| Valve control relay board         | Grayhill 70MRCK24-HL® PCB                                                                      |
| Excimer Laser control DI/O device | National Instruments USB-6501®                                                                 |
| Valve control DI/O device         | National Instruments USB-6509® with a R100M-50F-50F Ribbon Cable and a CB-50LP connector block |
| Bakeout Controller                | Amptek BT-15®                                                                                  |

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3 Construction

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The main physical components of the Jimbochron are the framing (physical structure that holds the skeleton and components), skeleton (tubing and valves), reservoir tanks, pumps and getters, sample chamber, and quadrupole mass spectrometer. Electricity and compressed air are also required for the Jimbochron’s operation. Table 1 lists the main parts and suppliers. Figure 1 illustrates their layout on the Jimbochron.

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3.1 Framing

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The Jimbochron is constructed on a metal frame of 1.5” wide T-slotted aluminum bars to ensure that the components and tubing of the line are arranged and supported properly. T-slotted aluminum framing systems are inexpensive and versatile, with multiple vendors offering a wide variety of connectors, brackets, mounts, and other accessories. The framing lengths can be cut easily using a non-ferrous metal blade on a standard miter or cutoff/chop saw, so all sizing and finishing can be done in-house, allowing for future modifications and additions. It also is straightforward to tap threaded holes into the framing, further simplifying customization. Additionally, the Swagelok® pneumatic valve bodies (see Sect. 3.2) can be attached directly to the aluminum framing using the two standard mounting holes present on the Swagelok® 5 series actuator bodies, negating the need for additional valve supports or holders.

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The aluminum framing is light enough, even when holding all of the line equipment, that it can be placed on wheels and moved around easily. Mobility is an important consideration for our design to allow the line to be as close as possible to the excimer laser during use, but conveniently moved out of the way for maintenance or other laser applications. Although the rigidity of aluminum can transmit vibrations from turbo or roughing pumps, these vibrational effects can be avoided (see Sect. 3.5). While welded steel frames can offer more strength and stability, aluminum frames are lighter, more adaptable, and require little skill, equipment, or experience to assemble.

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The aluminum frame was constructed so that the midpoint height of the Jimbochron matches the height of the sample chamber of our ESL NWR193UC® laser (~110 cm above the ground). This design requires only a short (several cm) flex hose for the connection between the two, therefore minimizing the total line volume. Excimer lasers are relatively tall compared to other laboratory equipment, so a “standard”-height analysis line would require longer connector tubing.

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199 **3.2 Skeleton**

200 The term “skeleton” refers to the central network of tubing and valves to which various components can be attached (Fig. 1).  
201 The main skeleton of the Jimbochron consists of ¼” outer diameter (OD), electro-polished, stainless-steel tubing and  
202 manifolds. The components are attached to all-metal Swagelok pneumatic valves with normally closed actuators and ConFlat®  
203 or VCR® style all-metal, bakeable, ultra-high vacuum compatible seals.

204

205 ConFlat® seals use a copper gasket pressed between two steel flanges with sharp, “knife edge” ridges to form the seal. ConFlat®  
206 flanges are the preferred vacuum seals for the main line components because they are stable and sealing them does not introduce  
207 any torque into the line. Although vertical ConFlat® flanges can be challenging to seal properly because of difficulties in  
208 properly aligning the gasket, we found the method of McCabe and Utz (1998) to be highly effective.

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210 VCR® seals use copper, nickel, or steel gaskets pressed between two highly polished and rounded surfaces. Components made  
211 with multiple VCR® flanges can be more difficult to seal than ConFlat® flanges, owing to the torque introduced while  
212 tightening the seals. Therefore, VCR® seals are only used in a few instances on the Jimbochron and avoided when possible.

213 **3.3 Reservoir tanks**

214 Our method for calibrating the mass spectrometer requires three isotopically pure reservoir tanks (one tank with <sup>3</sup>He and two  
215 tanks with <sup>4</sup>He), each attached to their own ~0.2 cc all-metal gas pipette (Sect. 5; Fig. 1). These tanks are referred to as the  
216 working <sup>4</sup>He tank (4T), the <sup>3</sup>He spike tank (3T), and the <sup>4</sup>He depletion correction tank (DCT). The pipettes are made by Becker  
217 Systems® using all-metal Swagelok BG Series 3® pneumatic valves so that they can be controlled using the same pneumatic  
218 air system as the other valves on the line. The reservoir tanks are made from stainless steel with an electropolished interior,  
219 are designed to hold ~10 L, and were custom-made by NorCal Products®. Each tank has one 1.33” ConFlat® port where a  
220 stainless steel ConFlat® tee connects the tank to its pipette and holds a small SAES ST172® donut getter. Each getter is mounted  
221 on two-pin ConFlat® feedthroughs, is run cold, and helps maintain the He purity in each tank.

222 **3.4 Sample chamber**

223 The sample chamber of the laser was provided by ESL® and is made of two 4.5” ConFlat® flanges. The bottom flange has a  
224 depression to hold the sample and a port that can be connected to the vacuum line. The top flange contains a deep UV sapphire  
225 glass viewport. Sapphire glass and fused silica viewports are both compatible with 193 nm excimer lasers. While sapphire  
226 viewports tend to have lower leak rates and lower He blanks than fused silica, fused silica viewports have significantly higher  
227 transmissivities. We have tested both materials and decided to use the sapphire viewport because the He blanks with the fused

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237 silica viewport were far too high. The chamber attaches to the ESL<sup>®</sup> laser sample stage which holds the sample at the  
238 appropriate focal length for the laser to focus on the sample surface.

239 **3.5 Pumps, getters, and gauges**

240 The UHV conditions ( $P_{\text{total}} < 1 \times 10^{-8}$  torr) required for low blank and high-precision analyses on the Jimbochron are  
241 maintained using three vacuum pumps (Fig. 1). The line's workhorse pumps are an Agilent IDP-3<sup>®</sup> dry scroll pump backing  
242 an Agilent TwisTorr 74<sup>®</sup> turbo pump. Although other turbo pumps have higher quoted He compression ratios and pumping  
243 speeds, in our experience the actual difference in pumping efficiency between turbo pump brands isn't that significant, perhaps  
244 indicating that the line conductance is the dominant factor. Agilent pumps were chosen because of their price (CU Boulder has  
245 favorable purchase agreements with Agilent), our experience with Agilent pump reliability, and the ease with which we can  
246 communicate with the pump controllers. The standard arrangement is for the scroll pump to work as a backing pump for the  
247 turbo. However, large volumes of gas can be pumped using only the scroll pump by isolating the turbo pump to avoid damaging  
248 the turbo rotors. Both the scroll and turbo pumps create significant vibrations, and during initial development we observed that  
249 these vibrations transmit through the aluminum framing of the line. To avoid problems associated with persistent vibration,  
250 both pumps are connected to the line only through flexible stainless steel bellows tubing, and reside on a wheeled cart that is  
251 not directly attached to the main line. Although future designs will shorten the tubing connecting the pumps to the line, in  
252 practice pump down times are acceptable as is. The third pump is an Agilent VacIon 20 StarCell<sup>®</sup> ion pump, which is only  
253 used to pump volumes when the pressures are already low ( $P_{\text{total}} < 1 \times 10^{-7}$  torr). Future expansion plans include the addition  
254 of more pumps to allow for greater flexibility in gas handling.

255  
256 In addition to the vacuum pumps, the Jimbochron employs multiple getters to both reduce the total line pressure and remove  
257 non-noble gases (see also Sect. 2.3). We currently use a combination of CapaciTorr<sup>®</sup>, Sorb-AC<sup>®</sup>, and ST172 SAES<sup>®</sup> getters,  
258 run both hot and at room temperature (Fig. 1; [Table 1](#)).

259  
260 The Jimbochron measures the vacuum with three different gauges (Fig. 4). Between the turbo pump and the roughing pump is  
261 a MKS 925<sup>®</sup> Pirani gauge capable of measuring pressures from atmospheric to  $\sim 1 \times 10^{-5}$  torr. The Pirani gauge is mainly used  
262 during the initial evacuation of the line to know when it is safe to engage the turbo pump. Between the turbo pump and Valve  
263 2 is an Instrutech Wasp<sup>®</sup> wide-range combination cold cathode and pirani gauge that can measure pressures from atmospheric  
264 pressure to  $7.6 \times 10^{-10}$  torr. This gauge is used to monitor the vacuum in the process side of the line. The ion pump measures  
265 pressure on the analysis side of the line but is only used when pressures are below  $\sim 1 \times 10^{-7}$  torr. The QMS can also be used  
266 to measure the pressure of the analysis chamber, and in practice gives readings that are similar to the ion pump.

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273 3.6 Quadrupole mass spectrometers

274 The Jimbochron has two QMSs. The main QMS is a Hiden 3F<sup>®</sup> equipped with both a faraday cup and a single channel electron  
275 multiplier, customized to only analyze a 20 amu range in order to maximize the resolution of the He peaks. We chose the  
276 Hiden 3F<sup>®</sup> primarily because the manufacturer provided data indicate that it is both approximately an order of magnitude more  
277 sensitive than other QMSs commonly used in (U-Th)/He labs, and its triple-quadrupole filters separate mass peaks more  
278 effectively than single-quadrupole filter machines. Additionally, Hiden<sup>®</sup> provides extensive and free LabView<sup>®</sup> drivers for  
279 their mass spectrometers which makes automation straightforward (see Sect. 5).

281 The Jimbochron also has a Stanford Research Systems (SRS) RGA100 QMS<sup>®</sup> (RGA-residual gas analyzer). This QMS is less  
282 expensive and less sensitive than the Hiden<sup>®</sup>, but has a larger mass range. It will be used for future continual ramped-heating  
283 (CRH) experiments, which require a mass spectrometer to be exposed directly to the sample chamber during heating and  
284 degassing (Idlemann et al., 2018). The CRH method can result in large gas volumes being introduced into the mass  
285 spectrometer chamber, and we decided it would be beneficial to allow the laser ablation side of the line (the Hiden<sup>®</sup>) to remain  
286 as clean as possible during experimentation and development. The CRH method is beyond the focus of this contribution, so  
287 this second QMS will not be discussed further.

290 3.7 Electricity and compressed air

291 Operation of the analysis line requires both electricity to operate various components and compressed air to open and close the  
292 pneumatic valves. Historically, the lab has suffered outages and interruptions of both power and compressed air, so the  
293 Jimbochron design includes backup systems for both.

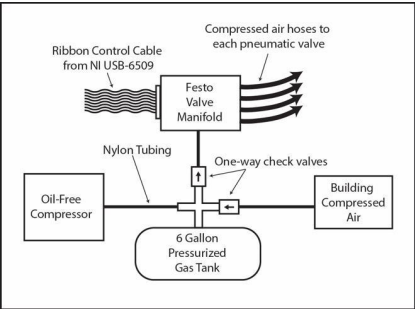
295 The electrical components are backed up by a single, rack-mounted, 120V uninterruptible power supply (UPS). During  
296 operation, the entire line only requires ~250 W of power, so even an inexpensive UPS backup system can provide >20 minutes  
297 of run time. In practice this system has worked successfully during all but the longest (>5 minutes) power outages that the line  
298 has experienced in the last few years. An integrated lab-wide UPS system would be ideal, but the cost of installation and  
299 maintenance, coupled with the rarity of long-duration power outages, has led us to not prioritize adding a lab-wide system.

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**Figure 2:** Schematic diagram of the compressed air backup system. A 6-gallon steel tank is connected through a tee-connector to a pressure-activated oil-free compressor, the building compressed air supply, and the Festo® valve manifold. One-way check valves ensure that if building air pressure is lost, the tank and compressor volumes will maintain their air pressure (direction of allowed flow shown by the arrows).

The compressed air system which supplies air to the Festo valve manifold is backed up by a reservoir tank that can be filled by either the building compressed air (primary source), or a pressure-activated, external, oil-free air compressor that is also on a UPS system (Fig. 2). The compressed air backup system is important not only for outages but also for low-pressure events caused by leaks, broken equipment, and building events that affect the compressed air supply. Although Swagelok® claims that the pneumatic valve actuators will work with pressures as low as 50 psi, in our experience the response is uneven below about 90 psi with some valves opening and others not.

Importantly, the Jimbochron is designed to fail safely during an extended power outage. The default configuration is for all of the valves to be closed. When power is lost, the valve manifold will depressurize even if the air pressure is sufficient for operation. The scroll pump also has an isolation valve on the inlet that automatically closes when power is lost, preventing air from rushing into the turbo pump. In practice when the Jimbochron loses power the interior of the line, including both the sample chamber and mass spectrometer, do not experience sudden increases in pressure.

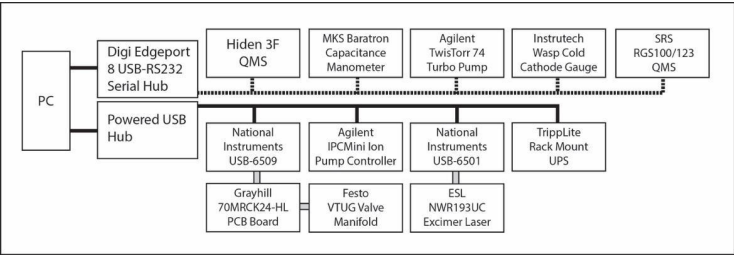
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## 4 Communication and automation

### 4.1 Communication

Efficient and reproducible noble gas analysis requires that all of the critical components are automated, which involves important hardware and software components. Most modern lab equipment has communication and control capabilities included as standard features, typically connecting to a PC via serial communication protocols.

344 The PC used to control the hardware has a limited number of serial ports and is far enough from the electronics rack holding  
345 the Jimbochron controllers that hubs and converters are leveraged to reduce the number of cables crossing the lab. USB cables  
346 from the PC lead to two separate hubs, a USB to DB9 serial hub with 8 ports, and a powered USB 3.0 hub with 6 ports. Figure  
347 3 is a schematic illustration of the connections and devices used for communication. Many of the devices, including both mass  
348 spectrometers, ion and turbo pump controllers, cold cathode gauge, and capacitance manometer, are connected directly to these  
349 hubs.



350  
351 **Figure 3: Schematic diagram of the communication connections used to control the Jimbochron. The solid black lines are USB**  
352 **cables, the dashed lines are RS232 DB9 serial cables, and the thicker grey lines are miscellaneous cables (ribbon or BNC).**

353 Two devices require additional communications hardware. First, the valve controls are routed through a USB to ribbon  
354 cable converter ([National Instruments USB-6509](#)), which operates relays on a [PC board](#). This board in turn powers solenoid valves  
355 in a Festo® pneumatic valve manifold that regulates compressed air delivery to the valves. The electrical connections on the  
356 [PC board](#) are also connected to manual 3-way switches so that each pneumatic valve can either be controlled remotely via the  
357 PC, or manually using switches mounted on the front panel of an electronics rack. Second, excimer laser automation is  
358 accomplished with a 24-channel USB digital I/O device ([National Instruments USB-6501](#)), connected to an external triggering  
359 port on the laser, which is a standard feature on ESL NWR193UC® excimer lasers.

360 **4.2 Automation**

361 Control and automation of the Jimbochron is done with custom software written in the [National Instruments LabView®](#)  
362 programming environment. Below, we explain the choice of Labview® and the key features of the software. However, in lieu  
363 of in-depth descriptions of the software architecture, access to the key components [is provided through an online repository](#).

365 LabView® was chosen [because of our in-house expertise with the platform, and because LabView® makes it straightforward](#)  
366 [to automate and control multiple pieces of equipment, perform simultaneous tasks, record data, and develop intuitive graphical](#)  
367 [user interfaces and controls](#). Many manufacturers provide LabView® drivers for their devices. LabView® also features tools to  
368 facilitate direct communication with devices that lack such drivers. Additionally, National Instruments®, the company that  
369 makes LabView®, sells a variety of [easy-to-use](#) hardware items for digital and analog I/O with almost any device, and LabView

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379 is compatible with many third-party digital and analog I/O products. Moreover, the University of Colorado provides discounted  
380 site licenses for LabView®, which makes this programming environment affordable. National Instruments also provides a fully  
381 functional version of LabView® called the “LabView Community Edition®”, which is free for non-commercial users and  
382 available for multiple operating systems.

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384 We prioritized developing four key software features based on our collective experience with analytical equipment. First, the  
385 code is editable and flexible. All key activities are written as their own subroutines, which allows equipment and software to  
386 be changed, repaired, replaced, or updated smoothly and quickly without modifying the entire project. It also allows new  
387 routines to be developed and tested offline and then added to the main code once complete. In contrast, code that locks the  
388 user into specific pieces of hardware will become outdated quickly, undermining both the longevity and efficiency of  
389 equipment.

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390  
391 Second, the software evaluates the line status before proceeding with a measurement routine. Although rare, lines that do not  
392 perform these checks can experience sample and data loss. Before ablating a sample, for example, the software will confirm  
393 that the mass spectrometer is still functioning (e.g. the filament hasn’t died or communication hasn’t been lost), that pressures  
394 are below a certain threshold, and that <sup>4</sup>He and <sup>3</sup>He readings are at the appropriate “blank” levels. This information allows the  
395 software to adjust in real time to changing measurement conditions, and ensures consistent analytical conditions for each  
396 sample. Additional safety checks or measurements can easily be added in the future.

397  
398 Third, every action taken either manually or during automation is logged and time-stamped. This feature was originally  
399 installed for debugging purposes, but has proven to be a useful record for quality control and diagnosing irregular behavior.

400  
401 Finally, all inputs, including individual analytical routines and sequences of multiple analyses, can be input as text files. This  
402 both creates a permanent record of all analyses and allows very large sequences (>100) of analyses to be loaded easily, which  
403 is important for tasks such as automated He mapping. This strategy also facilitates consistent analytical routines and sample  
404 sequences by enabling text files to be duplicated and updated for the next analytical run, removing the human error that can  
405 be introduced when re-entering complex analytical and sample sequences.

## 406 5 Determining volumes and calibrating depletion corrections from first principles

### 407 5.1 Absolute He amounts, volumes, and depletion corrections

408 (U-Th)/He chronology requires determining the absolute amount of <sup>4</sup>He in a sample. Mass spectrometers, however, only  
409 measure the magnitude of the beam of ionized particles collected on a detector (often in amps or volts depending on the  
410 detectors and amplifiers used). While the magnitude of this beam is proportional to the abundance of a particular mass,

413 converting a measured beam into moles or some absolute amount of He requires calibration. The Jimbochron uses a variation  
414 of the isotope dilution method, described below, to convert the mass spectrometer signals into absolute amounts of <sup>4</sup>He.

415  
416 Since the 1950's isotope dilution has been the preferred method for precise and accurate isotopic measurement of elements  
417 that have more than one isotope (e.g. Stracke et al., 2014). In isotope dilution, the sample is mixed with a known amount of  
418 another solution, called a "spike", which has a known concentration and is enriched in one or more of the isotopes of the  
419 element of interest in the sample. All of the isotopes of interest in the combined sample and spike mixture are then measured  
420 on a mass spectrometer. These data allow the amounts of isotopes in the unknown sample to be calculated (Webster, 1959).

421  
422 In many ways, <sup>4</sup>He is an ideal candidate for isotope-dilution analysis because natural samples contain essentially no <sup>3</sup>He and  
423 isotopically pure <sup>3</sup>He is available from multiple suppliers such as Sigma-Aldrich® and AirLiquide®. In geochronology it is  
424 common for spikes to be liquid solutions where the precise amount of spike added to a sample can be determined by weighing  
425 each aliquot (e.g. Stracke et al., 2014). Gas spikes, however, cannot be easily weighed. Instead, the absolute amount of gas in  
426 a spike aliquot is calculated using the ideal gas law, the pressure and temperature of the gas, and volume of the pipette. While  
427 pressure and temperature can be measured directly, precise and accurate volume measurements are more complicated.

428  
429 For gas analysis, the spike is delivered to the sample gas as an aliquot, or shot, of gas from a pipette attached to a reservoir  
430 tank. With each shot the pressure in the reservoir tank decreases, which means that the concentration of <sup>3</sup>He in each successive  
431 shot also decreases. This phenomenon is called "tank depletion" and can be corrected for as described below. Although this  
432 correction is small, any inaccuracy in the tank depletion rate accumulates over time. For example, in a typical year, a He line  
433 may make 5000 measurements of blanks and samples, each of which is spiked with a shot of <sup>3</sup>He. If the reservoir tank and its  
434 pipette have volumes of 10 L and 0.2 cc, respectively, these values yield a depletion rate of 0.99998, such that the 5000th shot  
435 would have ~10% less <sup>3</sup>He than the first shot. Thus, even a small inaccuracy in the amount of <sup>3</sup>He in the spike shot could  
436 become significant. Owing to regular use of the 3T tank (every analysis is spiked), this tank must be refilled and recalibrated  
437 when the <sup>3</sup>He signal in the spike shot becomes too small for precise measurement.

438  
439 There are two main ways to determine the tank depletion rate: it can be calculated or it can be measured directly. Calculating  
440 the tank depletion rate requires knowledge of only the volume of the pipette and the tank.

441  
442 
$$D = \frac{V_r}{V_r + V_p} \tag{1}$$

443  
444 Where D is the calculated depletion rate, V<sub>R</sub> is the volume of the reservoir tank, and V<sub>P</sub> is the volume of the pipette. Measuring  
445 the tank depletion rate requires an additional tank and pipette of <sup>4</sup>He, called the depletion correction tank (DCT), to regularly

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456 evaluate and refine the actual depletion rate of the <sup>3</sup>He tank (3T) and the <sup>4</sup>He tank (4T) and minimize compounding  
457 uncertainties. Directly measuring tank depletion is done by comparing the <sup>4</sup>He/<sup>3</sup>He of standards (using a shot of <sup>4</sup>He from the  
458 4T and a shot of <sup>3</sup>He from the 3T) to <sup>4</sup>He/<sup>3</sup>He of depletion correction measurements (using a shot of <sup>4</sup>He from the DCT and a  
459 shot of <sup>3</sup>He from the 3T) over time. Measuring the tank depletion correction directly on a regular basis is important because  
460 these measurements reduce the uncertainty in the depletion rate as the tank is used. Otherwise, this uncertainty would  
461 compound exponentially with tank use.

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462 **5.2 Standard-shot calibration method to determine the absolute amount of <sup>4</sup>He in a sample**

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463 The isotope-dilution variation used by the Jimbochron is the standard-shot method. This approach measures shots of a known  
464 amount of <sup>4</sup>He (from the 4T tank) spiked with <sup>3</sup>He (from the 3T tank) as standards, typically before, during, and after a batch  
465 of unknown analyses. This method assumes that the concentration in each <sup>3</sup>He shot is constant between these standard  
466 measurements, which is justifiable because the tank depletion over the course of a batch of measurements is much less than  
467 the sensitivity of the mass spectrometer (~0.04% for 20 shots). The ratio of the known amount of <sup>4</sup>He from the standards and  
468 the unknown amount of <sup>4</sup>He from the samples to the “constant” <sup>3</sup>He amounts can then be used to calculate the unknown <sup>4</sup>He  
469 amounts. This method has several advantages: the <sup>4</sup>He standard shots are used much less often than the <sup>3</sup>He spike shots, so  
470 compounding uncertainties from errors in the depletion correction take longer to gain significance, and the absolute amount of  
471 <sup>3</sup>He becomes irrelevant such that refilling the 3T tank requires no additional calibration.

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473 The standard-shot calibration method requires precise knowledge of only the amount of <sup>4</sup>He in the initial “calibration” shot  
474 from the 4T and the depletion rate of the 4T. The amount of <sup>4</sup>He in the calibration shot can be calculated directly from the tank  
475 and pipette volumes and the pressure the tank was filled with, or can be determined using an external standard.

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477 The volumes of all three tanks and pipettes (~10 L and ~0.2 cc, respectively) were chosen to minimize the tank depletion rate  
478 and therefore minimize compounding uncertainty in absolute <sup>4</sup>He determination. The pipette volume was also selected so that  
479 a shot of gas at the pressures the tanks could be reliably filled with would approximate the amount of <sup>4</sup>He in typical samples.  
480 Significantly larger pipettes would have required lower pressures in each tank to deliver the desired He amount than could be  
481 reliably measured with our capacitance manometer. Significantly smaller volume pipettes can be problematic to fabricate  
482 because the flexibility of the metal and the valve seat can become a significant proportion of the pipette volume and lead to  
483 inconsistent gas delivery (T. Becker, person. comm).

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485 As described in more detail below and shown in Fig. 4, our steps for accurately and precisely quantifying the pipette and tank  
486 volumes were: 1) metrologically determining the volumes of standard solid objects, 2) using the standard objects in expansion  
487 experiments to calibrate the volume of a vacuum chamber (thereby establishing a “known volume”), and 3) using the known  
488 volume in expansion experiments to calibrate the pipette and tank volumes. All expansion experiments were conducted using

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dry, ultra-high purity Ar gas. Ar was chosen because it is not affected by the getters, is easy to obtain, and any residual Ar left in the system will not impact He measurements. We checked our metrologically-determined volumes with volumes that we constrained using our existing calibrated He line. Following these volume measurements, the tanks were filled with isotopically pure <sup>3</sup>He or <sup>4</sup>He to the target tank pressures.

An advantage of determining internal volumes and calibrating depletion corrections from first principles during the construction of a noble gas line is that uncertainties can be determined and propagated at each step. Uncertainty on the Jimbochron is accounted for in three ways. Uncertainty of the densities of the standard solid objects (Al and Si, see next section) are from the literature. Uncertainties on measured values are determined through repeated measurement (e.g. the uncertainty on the mass of the Al rods was determined by computing the standard error of 20 separate measurements). Uncertainties for calculated values, like the volume of the Al rod, are propagated from the density and mass values used as inputs. Figure 4 is a flow chart showing how the measured and calculated values combine to determine the absolute amount and uncertainty of <sup>4</sup>He in a sample.

5.3 Determining standard volumes

Calibration of volumes in the vacuum line first requires determining the volume of one or more standard solid objects, V<sub>s</sub>. Our standard solid objects are the right size to fit in the vacuum chamber and use readily available materials with well characterized densities so that volumes can be determined metrologically: 1) a ~12 mm diameter, 99.9999% pure rod of Aluminum that was cut into two pieces (density of 2.6984 +/- 0.0003 g/cc; Sverdlin, 2003), and 2) three 99.999% pure ~1 cm<sup>3</sup> Silicon metal cubes (density of 2.329002 +/- 0.000007 g/cc; Henins, 1964; Table 2). While Morgan et al (2011) used steel spheres in their calibrations, steel density can vary significantly, necessitating an independent determination of either volume or density. Volumes for the Al rods and Si metal cubes in our study were determined using the above densities and masses measured repeatedly on a calibrated balance. Each mass was weighed at least 10 times, yielding 1<sub>s</sub> standard errors of < 0.002%. We chose to use multiple solid objects with different densities, rather than a single object, to check the reproducibility of our volume calculations and calibration procedures.

Table 2: Summary of items used for density and volume calibration.

| Item      | Mass (g) <sup>1</sup> | ± 1s SE | Density (g/cc) <sup>1,2</sup> | ± 1s SD <sup>2,3</sup> | Volume (V <sub>s</sub> ) (cc) <sup>1</sup> | ± 1s SD |
|-----------|-----------------------|---------|-------------------------------|------------------------|--------------------------------------------|---------|
| Al rod 1  | 20.00166              | 0.00003 | 2.6984                        | 0.0003                 | 7.4124                                     | 0.0001  |
| Al rod 2  | 9.95863               | 0.00004 | 2.6984                        | 0.0003                 | 3.6906                                     | 0.0001  |
| Si cube 1 | 2.24177               | 0.00003 | 2.329002                      | 0.000007               | 0.96255                                    | 0.00005 |
| Si cube 2 | 2.23193               | 0.00005 | 2.329002                      | 0.000007               | 0.95832                                    | 0.00006 |

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|--------------|---------|---------|----------|----------|---------|---------|
| Si cube 3    | 2.27550 | 0.00004 | 2.329002 | 0.000007 | 0.97703 | 0.00005 |
| All Si cubes | 6.74920 | 0.00007 | 2.329002 | 0.000007 | 2.8979  | 0.0001  |

<sup>1</sup> Each mass was weighed at least 10 times.

<sup>2</sup> Aluminum density from Sverdlin, A., 2003, Properties of Pure Aluminum, in Handbook of Aluminum, New York, Marcel Dekker, Inc.,1296 pages.

<sup>3</sup> Silicon density from Henins, I., 1964, Precision density measurement of silicon: Journal of Research of the National Bureau of Standards Section A: Physics and Chemistry, v. 68A, p. 529, doi:10.6028/jres.068A.050.

<sup>4</sup> The volumes reported here are the Vs values used in the calibration procedure and associated calculations. These are shown in Figure 4 and discussed in section 5.3. Volumes were not corrected for thermal expansion because the estimated corrections based on ambient temperature were ~1 X 10<sup>-15</sup> cc, much smaller than our uncertainties.

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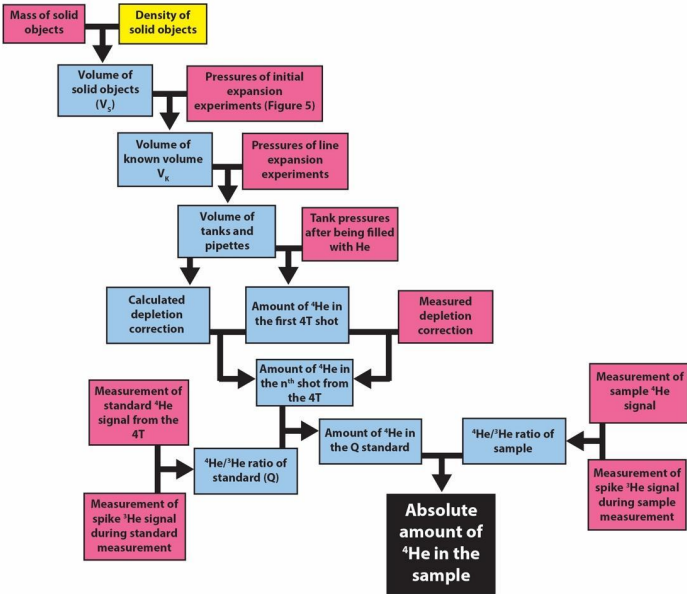


Figure 4: Flow chart showing how the absolute amount and uncertainty of <sup>4</sup>He in a sample (black box) are determined. The colored boxes are measured values (pink), calculated values (blue), and values obtained from the literature (yellow). The black lines and arrows represent calculations where uncertainties are propagated.

#### 5.4 Determining a known volume

The standard solid objects were used to calibrate a known volume,  $V_K$ , that could then be employed to calibrate the other volumes in the line. Theoretically we could have used any chamber of the Jimbochron for this purpose, but decided to instead

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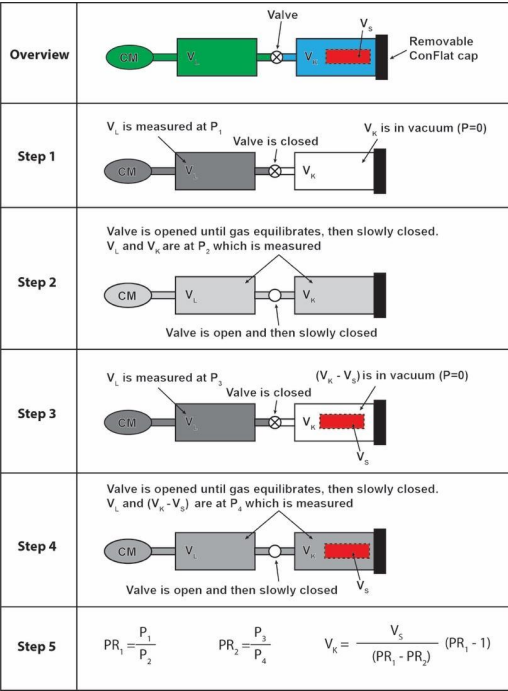
544 create a separate, removable vacuum chamber to use as  $V_K$  in case we wanted to calibrate other vacuum lines. Additionally,  
545 the sample chamber, which is the only chamber large enough on the Jimbochron to accommodate the known solid objects, had  
546 not yet been delivered at the time of calibration. The vacuum chamber used for  $V_K$  is a 3" long, 1.33" ConFlat® nipple attached  
547 to a manually actuated all metal Swagelok® valve with a standard ConFlat® flange. The nipple can be opened from the end  
548 opposite the valve, which is important for this method, and resealed in a repeatable and consistent manner. We anticipate  
549 changes in the internal volume due to slight differences in the position of the ConFlat® flange when it is re-sealed between  
550 experiments to be negligible, consistent with Morgan et al., (2011).

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552 Our method of determining  $V_K$  involves measuring the pressure of a gas with an absolute capacitance manometer, expanding  
553 the gas into an evacuated  $V_K$ , and measuring the resultant pressure drop (Fig. 5 steps 1 and 2). This process is repeated with a  
554 standard solid object ( $V_s$ ) inserted into  $V_K$  (Fig. 5 steps 3 and 4; see Table 3 for an example of these pressure measurements).  
555 The difference in pressure drops between these two experiments is recorded and used to calculate  $V_K$  (Fig. 5 step 5). This  
556 method is similar to that described by Morgan et al. (2011), but instead of comparing depletion curves uses expansion ratios  
557 to calculate internal volumes (Fig. 5). We recorded pressures using a 1 torr maximum range MKS 690A Baratron® absolute  
558 capacitance manometer connected to an MKS 670B® signal conditioner. Theoretically, a high-quality mass spectrometer  
559 should measure the same relative drops in partial pressure and allow for the measurement of the internal volume of noble gas  
560 lines without purchasing an expensive absolute capacitance manometer or breaking vacuum to install one.

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**Figure 5: Illustration of the process to calibrate a known volume ( $V_K$  denoted in blue in the Overview) using the metrologically-determined volume of one or more standard solid objects ( $V_S$ , denoted in red in the Overview). Pressures are measured with a capacitance manometer (CP), but could theoretically be measured instead with a mass spectrometer. Step 1: The pressure of a gas  $P_1$  is measured in  $V_L$  (in green, the volume of the line connecting the capacitance manometer and  $V_K$ ). Step 2: The gas in  $V_L$  is expanded into an evacuated  $V_K$ , the valve is closed, and the pressure of the gas  $P_2$  in both  $V_L$  and  $V_K$  are measured. Steps 3 and 4: Steps 1 and 2 are repeated for measurements of  $P_3$  and  $P_4$ , but with one or more standard solid objects with metrologically determined volumes inserted. Step 5 shows the equations used to calculate  $V_K$  given the difference in pressure ratios (PRs) between the two expansion experiments ( $PR_1$  and  $PR_2$ ). When a volume is described as being “in vacuum”, it is well below the minimum pressure measurable by the capacitance manometer, which in our experiments was  $\sim 1 \times 10^{-7}$  torr.**

The volume of  $V_K$  was measured ~~multiple~~ times for different combinations of Si cubes and Al rods. Pressures measured every 0.5 seconds were recorded using custom LabView software for processing. The pressure measurements and corresponding uncertainties were corrected for both baseline and thermal transpiration (Poulter et al., 1983). Thermal transpiration is relevant because our capacitance manometer operates at 45°C whereas the volumes in the line were at ambient laboratory temperature (19-22°C). During measurements, laboratory temperatures were monitored to account for volume changes from thermal expansion, but these volume change estimates were well below the detection limit of the instrumentation ( $\sim 1$  fcc, femto-cm<sup>3</sup>),

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581 and therefore are ignored. Table 3 shows the results of one set of expansion experiments (Fig. 5) using the larger Al rod as the  
582  $V_K$ . The value for  $V_K$  reported in Table 4 is an average of multiple expansion experiments using combinations of Al rods and  
583 Si cubes.

**Table 3:** Example Volume Calibration Measurements using the Aluminum Rod<sup>1,2</sup>

*PR1 Measurements (Expansion into empty  $V_K$ )*

| Attempt            | P1 (mtorr) | ±    | P2 (mtorr) | ±    | PR1            | ±              |
|--------------------|------------|------|------------|------|----------------|----------------|
| 1                  | 700.38     | 0.03 | 589.34     | 0.01 | 1.18841        | 0.00005        |
| 2                  | 680.76     | 0.02 | 572.88     | 0.01 | 1.18831        | 0.00004        |
| 3                  | 734.68     | 0.01 | 618.13     | 0.01 | 1.18855        | 0.00002        |
| 4                  | 731.04     | 0.01 | 615.13     | 0.01 | 1.18844        | 0.00003        |
| 5                  | 732.25     | 0.02 | 616.10     | 0.01 | 1.18853        | 0.00003        |
| 6                  | 731.88     | 0.01 | 615.80     | 0.02 | 1.18851        | 0.00003        |
| 7                  | 807.67     | 0.03 | 679.53     | 0.01 | 1.18857        | 0.00004        |
| 8                  | 599.53     | 0.01 | 504.48     | 0.01 | 1.18841        | 0.00003        |
| 9                  | 596.76     | 0.01 | 502.17     | 0.01 | 1.18837        | 0.00002        |
| 10                 | 800.09     | 0.01 | 673.17     | 0.01 | 1.18855        | 0.00002        |
| 11                 | 594.13     | 0.01 | 499.91     | 0.01 | 1.18848        | 0.00003        |
| 12                 | 795.16     | 0.02 | 669.03     | 0.02 | 1.18852        | 0.00004        |
| 13                 | 899.58     | 0.01 | 757.04     | 0.01 | 1.18828        | 0.00002        |
| <i>PR1 Average</i> |            |      |            |      | <i>1.18846</i> | <i>0.00009</i> |

*PR2 Measurements (Expansion into  $V_K$  with Al rod inside)*

| Attempt            | P3 (mtorr) | ±    | P4 (mtorr) | ±    | PR2            | ±              |
|--------------------|------------|------|------------|------|----------------|----------------|
| 1                  | 787.75     | 0.02 | 715.01     | 0.02 | 1.10172        | 0.00004        |
| 2                  | 787.63     | 0.03 | 714.92     | 0.01 | 1.10170        | 0.00004        |
| 3                  | 787.35     | 0.03 | 714.66     | 0.01 | 1.10171        | 0.00004        |
| 4                  | 786.39     | 0.02 | 713.80     | 0.01 | 1.10169        | 0.00003        |
| 5                  | 786.73     | 0.01 | 714.11     | 0.02 | 1.10171        | 0.00003        |
| 6                  | 787.91     | 0.02 | 715.17     | 0.01 | 1.10171        | 0.00003        |
| 7                  | 788.00     | 0.02 | 715.27     | 0.01 | 1.10169        | 0.00003        |
| 8                  | 787.15     | 0.02 | 714.43     | 0.02 | 1.10180        | 0.00003        |
| 9                  | 787.40     | 0.02 | 714.72     | 0.01 | 1.10169        | 0.00003        |
| 10                 | 860.72     | 0.03 | 781.30     | 0.02 | 1.10165        | 0.00004        |
| 11                 | 856.99     | 0.01 | 777.84     | 0.01 | 1.10176        | 0.00002        |
| <i>PR2 Average</i> |            |      |            |      | <i>1.10171</i> | <i>0.00004</i> |

<sup>1</sup> This experiment uses  $V_S = 7.4124 \pm 0.0001$  cc for the Aluminum rod (see Table 2).

<sup>2</sup> This set of experiments yields a value for  $V_K = 16.104 \pm 0.003$  cc. This value differs from that reported in Table 4 which is an average of multiple experiments using the different solid objects listed in Table 2.

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587 **5.5 Determining tank and pipette volumes**

588 Once we were confident in the value of  $V_K$ , volumes for the rest of the vacuum line were determined sequentially using Boyle's  
589 law and the capacitance manometer. Multiple gas expansion experiments were run on each volume to ensure reproducibility  
590 and determine measurement uncertainties. The volume of the pipettes, for example, were measured 3-6 times using different  
591 loading pressures, with the resultant 1 $\sigma$  SE on volume determinations  $\leq 0.5\%$ . Table 4 reports all measured volumes. Figure 4  
592 illustrates how uncertainties were propagated for these values.

**Table 4:** Summary of measured internal volumes.

| Name                   | Volume (cc) | $\pm$ (1 $\sigma$ SE) | Description                                                                                                  |
|------------------------|-------------|-----------------------|--------------------------------------------------------------------------------------------------------------|
| $V_K$                  | 16.11       | 0.07                  | This is the known volume described in Sect. 5.3, used to calibrate the rest of the line.                     |
| $V_{QCalibration}$     | 47.93       | 0.05                  | Volume of the interior of the line with only valves 4 and 11 open.                                           |
| $V_{DTCalibration}$    | 56.60       | 0.38                  | Volume of the interior of the line with valves 4, 9, and 11 open.                                            |
| $V_{Analysis}$         | 60.11       | 0.03                  | Volume during an analysis with valves 4, 6, and 11 open.                                                     |
| $V_{Pipette\_1}$ (3T)  | 0.2111      | 0.0003                | Volume of the gas pipette used for $^3\text{He}$ from the 3T tank.                                           |
| $V_{Pipette\_2}$ (4T)  | 0.2150      | 0.0003                | Volume of the gas pipette used for $^4\text{He}$ from the 4T tank used for regular calibrations.             |
| $V_{Pipette\_3}$ (DCT) | 0.2169      | 0.0005                | Volume of the gas pipette used for $^4\text{He}$ from the DCT tank used for tank depletion experiments only. |
| $V_{Tank\_1}$ (3T)     | 10341       | 12                    | Reservoir tank of pure $^3\text{He}$ known as the 3T tank.                                                   |
| $V_{Tank\_2}$ (4T)     | 10344       | 12                    | Reservoir tank of pure $^4\text{He}$ known as the 4T tank, used for regular calibrations.                    |
| $V_{Tank\_3}$ (DCT)    | 10368       | 12                    | Reservoir tank of pure $^4\text{He}$ known as the DCT tank, used for tank depletion experiments.             |

593

594 The volumes of the three reservoir tanks were more difficult to measure using this approach than the other line volumes for  
595 several reasons. First, each tank volume is much larger than any other volume, so when the higher pressure in the relatively  
596 small line volume ( $\sim 50$  cc) was expanded into the much larger tank volume ( $\sim 10,000$  cc), the final tank pressure of  $\sim 5$  mtorr  
597 was near the minimum sensitivity of the capacitance manometer. Second, the gas can only access the tanks through the pipettes,  
598 which do not have high conductance, such that it took a long time ( $\sim 15$  minutes) for pressures to equilibrate once expanded.  
599 During gas expansion the volume being measured is not pumped, so with very long equilibration times comes corresponding  
600 slow increases in pressure. These pressure increases are irrelevant for the higher pressure measurements on the other, smaller  
601 line volumes. However, because the tank expansions yield much lower pressures at the limit of the capacitance manometer's  
602 measurement range, small pressure increases due to extended equilibration durations add uncertainty to the tank volume

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612 measurements. Third, impurities (which are slowly removed by gettering) and the slow gas accumulations that happen when  
613 volumes aren't pumped can make it difficult to determine when the pressure has actually equilibrated. Finally, because of the  
614 low conductance through the pipettes, pumping and properly evacuating the tanks after an expansion experiment was time  
615 consuming, limiting the number of full volume measurements that were feasible. There are other, more direct ways to measure  
616 tank volumes, notably filling the tank with water and determining the mass, however our tank design includes permanent  
617 getters which would be damaged by water.

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619 To circumvent the above challenges, an improved approach would be to fill the tank and interior of the line to ~1000 mtorr,  
620 close off the tank, evacuate the line, and then expand the gas trapped in the tank into the rest of the line. This strategy would  
621 require measuring pressures of ~1000 mtorr and ~995 mtorr, which are well within the calibration range of the manometer.  
622 This evacuation and expansion routine can be efficiently repeated multiple times, because the same gas load in the tank can be  
623 used rather than fully evacuating the tank before repeating the measurements.

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625 Our existing, calibrated He line enabled us to check our volume measurements (Table S) by introducing a known amount of  
626 <sup>4</sup>He into the Jimbochron for analysis, and comparing the results to those predicted from the volume calibrations. Because the  
627 pipette volumes determined metrologically as described above were reproducible and did not experience the same problems  
628 described with the tank volume determinations, the small discrepancy observed in this experiment suggests that the tank  
629 volume measurements were slightly too small. This tank volume discrepancy yields an estimated depletion correction for the  
630 4T tank that differs by only 0.00004%. This difference in the depletion correction is not significant because the tank depletion  
631 rate will be continually measured and refined as described above. The difference in the volume measurement would create a  
632 small discrepancy in the calculated amount of <sup>4</sup>He in the initial shot, which would impart a small but systematic error in He  
633 determinations if not addressed. We expect that the small tank volume discrepancy would be eliminated by the improved tank  
634 measurement strategy described above.

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### 635 5.6 Tank filling and He pressures

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636 For measurements of unknown <sup>4</sup>He amounts during regular use of the line, the <sup>3</sup>He spike shots and <sup>4</sup>He calibration shots should  
637 ideally be similar in size to the <sup>4</sup>He yielded by a typical sample. To achieve this objective, the target pressure for each reservoir  
638 tank was ~0.01 mtorr. Given the measured volumes of the tanks and pipettes, this tank pressure would deliver ~100-200 fmol  
639 of gas for the <sup>3</sup>He spikes and <sup>4</sup>He calibration shots, which is close to the median expected sample size for unknowns.

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641 In preparation for tank filling, the tanks, pipettes, and line were first baked at ~200°C for ~48 hours while being pumped by  
642 both turbo and ion pumps to ensure as evacuated a volume and as low a pressure as possible (pressures based on ion pump  
643 currents were <1 X 10<sup>-9</sup> torr). The tanks are pumped through the pipettes, which have low conductance, necessitating the long  
644 baking and pumping time. The target tank pressures are significantly lower than can be directly measured by any commercially

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656 available absolute capacitance manometer. The proper pressure in each tank was therefore achieved by loading each pipette  
657 with ~1 mtorr of either <sup>3</sup>He or <sup>4</sup>He (gettered extensively to ensure purity), which was expanded into the evacuated tank. The  
658 volume of He in each shot, as well as the depletion correction of the tanks owing to successive shots, can then be calculated  
659 directly from the volumes of the tank and pipette and the starting pressure of pure gas in the tank (Table 5).

**Table 5:** Reservoir tank pressures and depletion corrections

| Name                      | Loading pressure (mtorr) <sup>1</sup> | ± 1s SE | Final Tank pressure (mTorr) <sup>2</sup> | ± 1s SE | Calculated Depletion Correction <sup>3</sup> | Measured Depletion Correction <sup>4</sup> |
|---------------------------|---------------------------------------|---------|------------------------------------------|---------|----------------------------------------------|--------------------------------------------|
| P <sub>Tank_1</sub> (3T)  | 493.17                                | 0.01    | 0.01007                                  | 0.00001 | 0.9999796                                    | N/A                                        |
| P <sub>Tank_2</sub> (4T)  | 916.43                                | 0.02    | 0.01905                                  | 0.00001 | 0.9999792                                    | 0.9999788                                  |
| P <sub>Tank_3</sub> (DCT) | 919.48                                | 0.02    | 0.01924                                  | 0.00002 | 0.9999791                                    | 0.9999731                                  |

<sup>1</sup> The pressure of the gas in the pipette before being expanded into the evacuated reservoir tank (see Sect. 5.5)

<sup>2</sup> The pressure of gas in the reservoir tank calculated using the volume of the tank and pipette (see Sect. 5.5)

<sup>3</sup> The depletion correction calculated by the measured volumes of the pipette and tank (see equation 1 and Sect. 5.1)

<sup>4</sup> The depletion correction measured by repeated tank depletion experiments (see Sect 5.1). Depletion on the 3T cannot be measured directly.

660 **6 Summary**

661 The new, custom He analysis line in the CU TRaIL facility is now fully operational and making automated, high-precision  
662 measurements of small volumes of He from sample materials. Designing and building the apparatus in-house, as well as  
663 developing custom software for communication and automation, means that the Jimbochron can be more easily repaired and  
664 upgraded than off-the-shelf instruments. Building a customized line can also be cost-effective; at the time of construction, the  
665 total cost of the components required to build the Jimbochron was less than \$160,000. Future expansion plans include the  
666 addition of a CRH setup (Idlemann et al., 2018) as well as more vacuum pumps to allow for more specific valve sequencing  
667 and to decrease the time required to evacuate the line between analyses. We also plan to replace both the sample and mass  
668 spectrometer chambers with smaller volume components.

670 The high sensitivity of the Jimbochron due to a relatively small internal line volume, more sensitive mass spectrometer, and  
671 optimized tuning enables precise measurements of small gas volumes (<0.1 fmol). This capability allows for smaller  
672 sampling volumes, more targeted He spot analyses, higher-precision measurements on younger samples, analysis of smaller  
673 grains, and higher spatial resolution mapping of individual crystals.

675 Measurements of He from secondary mineral standards, including Durango fluorapatite (e.g., McDowell et al., 2005) and Juina  
676 zircon (currently used as an in-house standard, Zeigler et al., 2025), have demonstrated the ability of the Jimbochron to  
677 accurately and precisely measure typical volumes of <sup>4</sup>He released during laser-ablation analysis (~0.1-1000 fmol). Full laser-  
678 ablation (U-Th)/He date calculations, which require additional measurements, are also consistent with expected values.

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Deleted: Our sensitivity estimates of the Jimbochron indicate that it is approximately 2 orders of magnitude more sensitive than the lab's Alphachron He line (~5 X 10<sup>-11</sup> V/fmol compared to ~3 X 10<sup>-13</sup> V/fmol).

Deleted: The minimum <sup>4</sup>He blanks measured in both machines are similar (~0.04 fmol). However, for the same gas volume, the Jimbochron yields a larger signal to noise ratio with lower uncertainties on repeated measurements and more consistent and reproducible blank measurements. This improved

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Automated analytical sequences have also successfully created He abundance maps of individual zircon crystals. We expect further performance improvements as we test and gain experience with a variety of analytical conditions and continue experimenting with grain mounting materials, sample preparation protocol, and analytical (QMS) settings.

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Author Contributions

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JRM and RMF conceptualized the project and acquired funding; JRM constructed the line and performed the calibration experiments; JRM and RMF prepared the manuscript.

Competing Interests

The contact author has declared that none of the authors have any competing interests.

Disclaimer

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