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**DiadFit: An Open Source Python3 Tool for Peak Fitting of Raman Data from silicate melts and volcanic fluids.**

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DiadFit: An Open-Source Python3 Tool for Peak fitting of Raman Data from silicate melts and volcanic fluids

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ABSTRACT

We present DiadFit - an open-source Python3 tool for efficient processing of Raman spectroscopy data collected from fluid inclusions, melt inclusions and silicate melts. DiadFit is optimized to fit the characteristic peaks from CO2 fluids (Fermi diads, hot bands, 13C), gas species such as SO2, N2, solid precipitates (e.g., carbonates), and Ne emission lines with easily tweakable background positions and peak shapes. DiadFit's peak fitting functions are used as part of a number of workflows optimized for quantification of CO2 in melt inclusion vapour bubbles and fluid inclusions. DiadFit can also convert between temperature, pressure and density using various CO2 and CO2-H2O equations of state (EOS), allowing calculation of fluid inclusion pressures (and depths in the crust), conversion of homogenization temperatures from microthermometry to CO2 density, and propagation of uncertainties associated with EOS calculations using Monte Carlo methods. There are also functions to quantify the area ratio of the silicate vs. H2O region of spectra collected on silicate glasses to determine H2O contents in glasses and melt inclusions.

KEYWORDS: Raman spectroscopy; Melt inclusions; Fluid inclusions; Volcanology; Open-source; Python.

1 INTRODUCTION

In recent years, Raman Spectroscopy has been increasingly used by igneous petrologists to perform non-destructive, in-situ measurements of melt and fluid compositions [Steele-Macinnis et al. 2011; Morizet et al. 2013; Hartley et al. 2014; Moore et al. 2015; Schiavi et al. 2018; Giordano et al. 2020; González-García et al. 2020; Schiavi et al. 2020]. Since 2014, there has been a growing body of literature using Raman Spectroscopy to measure the density of CO2-rich fluids in melt inclusion vapour bubbles to more accurately obtain the total CO2 content of the melt inclusion, and thus the magma storage depth [Hartley et al. 2014; Moore et al. 2015; Lamadrid et al. 2017; Allison et al. 2021; Wieser et al. 2021; DeVitre et al. 2023b]. Raman spectroscopy also shows enormous potential to quantify the densities of CO2-rich fluid inclusions [Wang et al. 2011; Kobayashi et al. 2012], allowing rapid and precise estimates of magma storage depths [Dujic et al. 2023; DeVitre and Wieser 2024; Zanon et al. 2024].

However, while there have been a large number of studies optimizing calibration and analysis protocols for CO2-rich fluids by Raman Spectroscopy (e.g., [Lamadrid et al. 2017; Bakker 2021; DeVitre et al. 2021]), there has been comparatively less focus on improving software tools for Raman peak fitting in volcanology (e.g. [Yuan and Maganovic 2017]). A number of highly flexible python packages for fitting Raman spectroscopy data are available (e.g., Ramp, [Le Losq 2018], RamanSpy, [Georgiev et al. 2023]). At the other end of the spectrum, van Gerve and Namur [2023] published a python-based graphic user interface for the specific use case of quantifying H2O contents of silicate glasses. However, there are no packages providing optimized workflows for the various procedures used to quantify CO2 in volcanological systems. Instead, many studies use proprietary peak fitting software from Raman instrument companies (e.g., [Moore et al. 2015]), which affects reproducibility because data is collected (and thus processed) on different Raman instruments (e.g., WITec, HORIBA, Bruker, Renishaw). Another popular approach is to use the open-source peak fitting GUI Fitk [Wojdor 2010]. While scripting is possible in this tool, most users fit curves by manually clicking to select peak and background positions (e.g. [DeVitre et al. 2021]). Other studies use OriginLab software which has an annual subscription fee of ~ 200 USD per license per year (e.g., [Hartley et al. 2014]), or have developed their own Matlab/Python codes for specific workflows and instruments (e.g., [Allison et al. 2021; Wieser et al. 2021]).

DiadFit is a high-level python package which aims to bridge the gap between generic Raman packages which allow experienced coders to build highly customizable workflows for the specific application of interest, and easier-to-use GUI-based methods which are hard to automate. DiadFit is aimed at volcanologists with a wide range of coding experience. As well as providing functions for peak fitting, this package also addresses the fact that there are no widely-available python tools for common calculations associated with fluid inclusion data, such as calculating CO2 densities from microthermometry data, and converting CO2 densities from fluid inclusions into pressure and depths using the CO2 or CO2-H2O equation of state. At present, EOS calculations are typically performed using computer programs with a graphical or text-based user interface (e.g., FLINCOR, [Brown 1989], FLUIDS, [Bakker 2003]), which can be difficult to install on modern operating systems. Many available tools require calculations to be performed manually for each fluid inclusion.

By integrating functions for peak fitting and subsequent EOS calculations into a single package, entire workflows can be performed in DiadFit without having to swap between different software tools (Fig. 1). In addition to reducing time spent data processing, DiadFit will also result in more consistent fitting between different groups, increasing reproducibility. For example, to align with FAIR principles (Findable, Accessible, interoperable, reproducible), users can publish a
supporting folder containing their spectra, relevant metadata, and Jupyter notebooks showing exactly how spectra were fitted and subsequent calculations performed (e.g. [Dayton et al. 2023; DeVitre and Wieser 2024]).

The shapes and locations of the Fermi diad and Ne lines are well defined, and spectra collected on any given Raman instrument have similar features; DiadFit capitalizes on this similarity to automate many workflows. Each user should tweak generic peak fit parameters for their specific analytical set up until they get good visual fits to representative spectra and acceptable residuals/peak fitting errors. Then, hundreds of spectra can be fitted automatically within minutes using the tweaked parameters, eliminating tedious (and subjective) workflows requiring lots of clicking. DiadFit also outputs the uncertainty associated with each peak fit, which allows for a deeper appreciation of the errors associated with different instrument hardware and acquisition conditions.

To demonstrate the time-saving nature of DiadFit, we compared the time taken to fit Neon lines and CO2 acquisitions collected over a 24 hour period during instrument calibration by DeVitre et al. [2021] in Fityk vs. DiadFit. Fitting these spectra took ~8 hrs using Fityk, and ~15 minutes using DiadFit on a typical laptop with 16 GB of RAM and an i7 processor. Given the potential for fluid inclusion analyses to provide rapid estimates of magma storage depths during volcanic crises [Dayton et al. 2023; DeVitre and Wieser 2024; Zanon et al. 2024], it is vital to speed up data processing as much as possible to reap the full benefits of this speedy technique. We anticipate that users who are not familiar with Python will simply use the provided Jupyter Notebooks and narrated YouTube videos, changing simple parameters like the path to their files and peak fit parameters to adjust for the different appearance of spectral peaks on different Raman instruments. More experienced coders can adapt the base functions in DiadFit to create their own workflows to address whatever science question they desire.

2 Code Installation, Underlying Dependencies and Architecture

For detailed instructions on code installation and dependencies (i.e., underlying packages required by DiadFit) and supported file types we refer readers to the ReadTheDocs page (https://diadfit.readthedocs.io/en/latest/). Briefly, DiadFit is available on PyPI (https://pypi.org/project/DiadFit/), so can be installed using pip within a Jupyter Notebook or through your command prompt. DiadFit uses a number of functions from different python libraries. The SciPy [Virtanen et al. 2020] find_peaks function is used to identify approximate peaks positions and attributes such as height, prominence, and presence/absence of certain peaks (based on pixel values, not fitted spectra). These estimated peak parameters can be used to subdivide spectra into groups with similar characteristics prior to fitting. Grouping spectra by characteristics (e.g., Fig. 2) and having good estimates of peak position and prominence greatly increases computational efficiency during iterative fitting of multiple overlapping peaks. Peak-fitting functions utilize the Python package lmfit, which supports iterative fitting of different peak types (e.g., Gaussian, Voigt, PseudoVoigt) and provides errors on peak fitting parameters (e.g., peak center, area, amplitude) from the covariance matrix. Numpy [Harris et al. 2020] is used for all basic math and regression operations, including fitting splines to spectra, and extracting subregions from spectra. Pandas [pandas development team 2020] is used for importing data from spreadsheet-type datafiles (.xlsx, .csv), and to output fit parameters in a tabular format. Matplotlib [Hunter 2007] is used for all plots produced by functions. EOS calculations using the Span and Wagner [1996] model are implemented through CoolProp [Bell et al. 2014]. This package is not a dependency for all of DiadFit, but if users want to perform calculations using this model, they need to have CoolProp installed (else they will receive an error containing installation instructions). Calculations using this EOS should cite both DiadFit and CoolProp.

3 Python Jargon

A user interacts with DiadFit by calling various functions. Each of these functions requires specific input parameters (called arguments). Five main datatypes are used in DiadFit as inputs/arguments and outputs:

1. A *string* is a piece of text. These are used to tell a function something about your data input, or specify a certain thing you want the function to do. For example, users must specify their file-type when using functions that load in spectra (e.g., spectra_filetype='headless.txt').

2. A numpy.array is a column or array of data without headings. DiadFit uses these to store spectral data after it has been extracted from proprietary instrument files. The wavenumber is stored in the 1st column (index 0 in Python), and the intensity in the 2nd column (index 1 in Python). Numpy allows for very fast computation.

3. A pandas.Series is a column of data with a heading.

4. A pandas.DataFrame is a collection of pandas.Series, and can be visualized as a single sheet in an excel workbook with labelled columns. In DiadFit, these are used to store peak fit parameters in columns with column headings (Fig. 5). Information in a given column of the dataframe (called df) can be accessed using the column heading: df['column_heading'].

5. dataclasses are used to define default peak finding and fitting configurations. These defaults can be tweaked as much or as little as required for each specific Raman spectrometer. For example, the default parameters for fitting Diad1 are stored in the dataclass diad1_fit_config:

```python
Diad1_fit_config(model_name='PseudoVoigtModel',
                 fit_peaks=2, fit_gauss=False,
                 gauss_amp=1000, diad sigma=0.2,
                 diad sigma_min_allowance=0.2,
                 diad sigma_max_allowance=5,
                 N_poly_bck_diad=1,
                 lower_bck_diad=(1180, 1220),
```
Figure 1: Schematic diagram showing the different workflows that can be performed in DiadFit.
Once these datACLasses are tweaked for a given instrument/set of samples, they can be used to automatically loop through large numbers of spectral files.

## 4 WORKED EXAMPLES

We have produced a number of example Jupyter Notebooks demonstrating specific workflows. These notebooks are available on GitHub ([https://github.com/PennyWieser/DiadFit](https://github.com/PennyWieser/DiadFit)) and the ReadTheDocs page ([https://diadfit.readthedocs.io/en/latest/](https://diadfit.readthedocs.io/en/latest/)). We include notebooks showing how to:

1. Fit peaks from CO$_2$-rich fluid inclusions and melt inclusion vapour bubbles and correct data for instrument drift using Ne line acquisitions. Some of these examples also show how to quantify the area of carbonate and SO$_2$ peaks, and calculate molar ratios of different gas species.

2. Perform calculations using the CO$_2$ and CO$_2$-H$_2$O equation of state, including converting homogenization temperatures from microthermometry into CO$_2$ densities, converting CO$_2$ densities from Raman or microthermometry into pressures, and then converting these pressures to depths using different crustal density profiles.

3. Propagate uncertainty in microthermometry and Raman-based fluid inclusion barometry into pressure and depth distributions.

4. Model fluid inclusion re-equilibration for different magma ascent, stalling and quenching scenarios following DeVitre and Wieser [2024].

5. Quantify peak asymmetry using the approach of DeVitre et al. [2023a] to help identify fluid inclusions which contain both liquid and vapour phases at the time of analysis.

6. Quantify H$_2$O contents in silicate glasses using an approach adapted from Di Genova et al. [2017] and Schiavi et al. [2018].

7. Quantify the ratio of H$_2$O to silicate glass peaks on Raman acquisitions taken on unexposed olivine-hosted melt inclusions. This can also be used to quantify H$_2$O contents in exposed silicate glasses (following [Di Genova et al. 2017]).

8. Propagating Raman and volume uncertainty when determining the contribution from vapour bubble CO$_2$ to the total CO$_2$ budget of melt inclusions.

## 5 FITTING THE CO$_2$ FERMI DIAD

The Raman spectrum of CO$_2$ consists of two relatively strong, well-defined peaks collectively termed the Fermi diad. These peaks result from the interaction of a symmetrical stretching mode and an active bending mode in the CO$_2$ molecule by the process of Fermi resonance ([Fermi 1931; Rosso and Bodnar 1995; Lamadrid et al. 2017]). One of the peaks is centered at ~1280–1290 cm$^{-1}$ (referred to as Diad1), and the second peak at ~1386–1390 cm$^{-1}$ (Diad2). It is well accepted that the distance between Diad1 and Diad2 (commonly referred to as splitting, diad splitting, Fermi diad separation, or Δ) correlates with the density of the CO$_2$ fluid. This is because with increasing CO$_2$ density, Diad1 moves to lower wavenumbers (shift of 6 cm$^{-1}$ from 0 to 0.8 g/cm$^3$), while the position of Diad2 stays more constant (only varying by ~1–2 cm$^{-1}$). The exact relationship between diad splitting and CO$_2$ density, commonly called a ‘densimeter’, has been shown to vary as a function of instrument hardware and acquisition conditions ([Lamadrid et al. 2017; DeVitre et al. 2021]).

The Fermi diad is flanked by low intensity hot bands (HB1 to the left of Diad1 and HB2 to the right of Diad2, Fig. 2), which arise from molecules populating the first excited vibron due to their thermal energy at room temperature, which are also perturbed by a Fermi resonance effect. The peak height of the HBs relative to the diad peaks increases with increasing temperature [Rosso and Bodnar 1995], and in general, hotbands become more pronounced in spectra collected on denser CO$_2$ fluids. Diad2 also has a flanking $^{13}$C peak to its left which becomes more visible as the CO$_2$ density increases and with increasing proportions of $^{13}$C relative to $^{12}$C ([Wang and Lu 2022]). Fitting these peaks can be used to quantify the relative abundance of $^{12}$C/$^{13}$C (see [Lu et al. 2023]).

In the literature, density estimates from CO$_2$-rich fluids such as melt inclusion vapour bubbles or fluid inclusions have been obtained by fitting the position of the two diad peaks using various background models combined with Voigt, Pseudovoigt, Gaussian or Lorentzian peaks. These peak fitting routines greatly increase the precision at which the diad splitting can be determined relative to the spectral resolution of the instrument (distance between datapoints). While Raman instruments typically acquire an intensity reading every 0.1–1.5 cm$^{-1}$, the error of the peak position on a fitted peak can be 10–30x smaller if an appropriate peak profile is fitted (e.g., Pseudovoigt/Voigt for diads, [Fukura et al. 2006; Lin et al. 2007; Yuan and Mayanovic 2017], see also Section 9.1). After the peaks are fit, splitting is calculated and corrected for instrument drift, an instrument-specific densimeter is used to convert corrected splitting into CO$_2$ density [Lamadrid et al. 2017; DeVitre et al. 2021].

When using DiadFit to peak fit CO$_2$ spectra, the first step is to find all the spectral files you wish to fit (by filtering based on file names, Fig. 3). Then, a single file is selected, and the DiadFit function `identify_diad_peaks` is used to tweak the `SciPy find_peaks parameters, displaying the identified peaks with yellow stars. This step is necessary because different Raman instruments have vastly different intensities and spectral resolutions. We find that the prominence, defined as the vertical distance above the neighbouring background, is the most important parameter to tweak between different instruments (some instruments have peaks 100s-1000s of counts above background while others are 100,000-1 million counts above background). Then, the function `loop_approx_diad_fits` is used to loop through all files to obtain approximate peak positions and prominences for the diad, hotbands and $^{13}$C peaks for each spectra, which is stored in a dataframe. This
function also calculates various additional parameters that are very helpful when grouping spectra in later steps (e.g. signal to noise ratios, elevation of the diad-HB region above background).

These approximate peak parameters are then used alongside a filter to remove any cosmic rays present in the spectra. The filtering process calculates the intensity factor $F_i$ based on the intensity ($I$) of any given spectral datapoint relative to the intensities of the datapoints to the left ($i-1$) and right ($i+1$):

$$F_{i-1} = \frac{I_i - I_{i-1}}{I_i}$$

$$F_{i+1} = \frac{I_i - I_{i+1}}{I_i}$$

These two factors are then multiplied together:

$$M_i = F_{i-1} \times F_{i+1}$$

If the spectra has a lower intensity to the left of datapoint $i$, $F_{i-1}$ will be $>1$. If the spectra also has lower intensity on the right hand side of point $i$, $F_{i+1}$ will be $>1$, so $M_i$ will be $>>1$. High values of $M_i$ for individual pixels are indicative of short, sharp peaks (i.e. most cosmic rays). A datapoint is considered a cosmic ray if $M_i$ exceeds a user determined value (e.g., $\text{dynfact}=0.001$). The highest pixel of Diads, HB and $^{13}$C peaks can also have high $M_i$ values, so any points with an $x$ coordinate equal to an identified peak position ($\pm$ the spectral resolution) are ignored. Users can also enter a custom range, within which no cosmic rays are removed (e.g., the area around the narrow SO$_2$ peak). If a cosmic ray is identified, the filter is run again, in case the cosmic ray was wider than a single pixel. The pixels identified as cosmic rays are then removed from the spectrum and a new file is saved with the same filename with the addition of the suffix _CRR_DiadFit.

After removal of cosmic rays, the identify_diad_peaks function is used again to obtain approximate peak parameters without inference from cosmic rays. These parameters are then used to divide spectra into groups. Dividing spectra into groups is advantageous because spectra from CO$_2$ fluids vary greatly in their spectral characteristics as a function of the sample size, depth, CO$_2$ density, laser power, and acquisition time (Fig. 2). Spectra collected on CO$_2$ fluids with low densities, at relatively deep depths, or using low laser power tend to have a weak to non existent hot bands (Fig. 2a-c). Stronger spectra have prominent hot bands and $^{13}$C peaks which overlap with the tails of the diad peaks (Fig. 2d-f). In the strongest spectra, the entire region around the diad peaks and hotbands is elevated above the near-linear background seen at greater distance from the diad peaks (Fig. 2g-i). Different fit parameters would be needed to obtain satisfactory fits on these different spectral types.

A full description of the different parameters used for classification is provided on the ReadTheDocs page, and the most useful parameters will depend greatly on your Raman instrument and samples. In general, spectra should be classified as ‘Weak’ when none of the peaks are strong enough to interfere with each other (e.g., Fig. 2c, where the hot bands and diad peaks do not have overlapping tails). Spectra should be classified in the ‘Medium’ group once the hotbands and diads begin to overlap substantially. This will likely result in the user wanting to specify background positions which are further from the peaks than for the ‘Weak’ group. Often, a Gaussian background is required in addition to the polynomial background for Diad2, because it becomes elevated. Spectra with very strong hot bands, clear $^{13}$C peaks, and greatly elevated troughs between diad peaks and HBs should be placed into the ‘Strong’ category. These spectra will need to be fitted with the addition of a Gaussian background to recreate the elevation of the spectra for both Diad1 and Diad2 (fit_gauss=True, Fig. 2c). The choice of groups is somewhat arbitrary, and is best viewed as an easy way to be able to set three different peak fit routines. In many studies, all spectra will fit in a single group. Users should not stress to much about whether to add a Gaussian background or not. If a Gaussian background isn’t really required (e.g., Fig. 2f) but it is requested, the code will converge on a Gaussian with a very small amplitude. Thus, the only real advantage of putting those not really needing Gaussian backgrounds into a ‘Medium’ Category is that the iterative fitting routine will be a bit faster (favorable when fitting large numbers of spectra).

One file from each group is then used to tweak the fit parameters (Fig. 3), which are applied to all spectra in the group (taking 1-5 s per spectra on a 16 GB intel i7 laptop). A figure of each fit is saved in the same folder as the Spectra and displayed in the notebook - fits should be carefully inspected. In particular, attention should be paid to the subplot showing the residual around the peak, and the selected background positions and background fit. The returned dataframe of fitted parameters also contains the uncertainties in peak positions, which can also be very helpful for identifying poor fits (Figure 10). In addition to plots for visual inspection of residuals, this function also calculates a measure of the residual for each peak:

$$\text{Residual} = \frac{\sum \sqrt{(y_{\text{meas}} - y_{\text{fitted}})^2}}{N_{\text{points}}}$$

where $y_{\text{meas}}$ is the $y$-coordinate of the background-subtracted data, $y_{\text{fitted}}$ is the $y$ coordinate of the best composite model fit, and $N_{\text{points}}$ is the number of discrete points along the $x$ axis used in the fit.

The next step is to combine the dataframe of peak fit parameters from each group, and combine this with information extracted from the metadata. In particular, it is necessary to obtain the time stamp of the analysis to apply the Ne correction model. Finally, splittings corrected for instrument drift are converted into CO$_2$ densities using a instrument-specific densimeter. Uncertainties from each step of this process are propagated (see section 9.1).

5.1 Fitting peaks to secondary phases

Raman spectra collected from fluid inclusions and melt inclusion vapour bubbles commonly contain peaks arising from other gaseous species (e.g., SO$_2$, N$_2$) or solid phases on the inclusion wall (e.g., carbonates, sulfates, [Schiavi et al. 2020], [Frezzotti et al. 2012]). It can be helpful to identify which spec-
Figure 2: Spectra from DeVitre et al. [2021] showing different characteristics to demonstrate how to group spectra for ease of fitting. Spectra classified as ‘Weak’ (a-c) have no $^{13}$C peak, and a small HB, which does not overlap with the diad (c). Spectra classified as ‘Medium’ (d-f) have a $^{13}$C peak, and the tails of the $^{13}$C, diad and HB peak all overlap (f). In this case, a Gaussian background was not added, but for any stronger spectra it would be required as the red dots are beginning to diverge from the best fit line (green curve) inbetween Diad2 and HB2. In spectra classified as ‘Strong’ (g-i), the troughs between $^{13}$C, Diad2 and HB2 are greatly elevated above the background. This requires a Gaussian background (purple curve) in addition to the 3 other peaks (red, blue, cyan) to minimise residuals.

tra contain such phases, and quantify characteristics of these peaks (e.g., peak heights, positions, relative peak areas, [Burke 2001]). The first step is to obtain all the spectra you wish to fit (Fig. 4). Then, the function plot_secondary_peaks plots the spectra in a specific wavenumber range from all selected files. Intensities are normalized based on the range of intensities in the selected wavenumber window, allowing multiple spectra to be stacked and inspected. It is clear from this visualization strategy that files #42 and #33 in Fig. 4 have prominent carbonate peaks. For any given set of spectra, you can inspect as many regions as you want in this way. For example, the second panel in Fig. 4 is centered around the SO$_2$ peak at $\sim$1151 cm$^{-1}$.

As well as plotting spectra to visually identify secondary phases, the function plot_secondary_peaks can identify peak positions within a user-defined wavenumber range using scipy find_peaks or a prominence filter. Identified peaks will be marked on the returned figure with a yellow star (Fig. 4). The function also returns a dataframe of approximate peak parameters based on the maximum pixel (position, height, prominence) - where no peaks are found, these columns are filled with NaNs. After identifying spectra containing secondary peaks, peak fitting parameters should be tweaked based on one selected spectra. Using the model_name parameter, secondary peaks can be fit with Gaussian, PseudoVoigt or Voigt curves, or a cubic spline. Tweaked fit parameters are then applied to all files. More than one type of secondary peak can be quantified in a given notebook. In Step 5, all identified secondary peaks are merged together with fitting parameters for diad peaks (see Fig. 3).

6 Fitting Ne lines

During any given analytical session, there may be some stretching or contraction of the Raman spectra; these changes are often referred to as ‘non-linearity of the Raman shift axis’. For brevity and to draw parallels with various mass spectrometry techniques, we term this ‘instrument drift’. Commonly, instrument drift within the spectral region containing the Fermi diad is assessed by repeatedly measuring the emission spectrum of Neon (Ne) produced by a lamp every $\sim$ 5-10
Fitting Fermi Diads

Get filenames to fit 
get_files()

Fit 1 spectra. Inspect +tweak if needed. 
fit_diad_1_w_bck() 
fit_diad_2_w_bck()

Overall fit for Diad 1

For each group, set fit parameters 
diad1_fit_config() 
diad2_fit_config()

Individual fit components

Inspect residuals

• For each group, fit all files 
  using these parameters
  • Save fit parameters
  • Repeat for each group

• Merge fits for different groups together
• Extract Ne correction model, apply to splittings 
  stitch_metadata_in_loop(),
  extracting_filenames_generic()

• Convert splitting to density using instrument 
  specific densimeter
• Inspect uncertainties

Check for cosmic rays, remove if present 
(cosmicray_filter(), filter_singleray(), filter_raysinloop())

Sort files into Weak, Medium and Strong 
(based on approx fit parameters)

plot_peak_params(), filter_splitting_prominence(), 
identify_diad_groups(), plot_diad_groups()

diad1_fit_config() 
diad2_fig_config()

stitch_metadata_in_loop(), 
extracting_filenames_generic()

Fit secondary peaks if relevant (see Fig. 4)

Loop over all files to find approx 
peak positions, heights, etc.

loop_approx_diad_fits()

Get filenames to fit 
get_files()

Fitting Fermi Diads
Get filenames to fit 
get_files()

Identify approx peak positions by 
tweaking SciPy 'find_peaks'
diad_id_config(), identify_diad_peaks()

Loop over all files to find approx 
peak positions, heights, etc.

loop_approx_diad_fits()

Check for cosmic rays, remove if present 
(cosmicray_filter(), filter_singleray(), filter_raysinloop())

Sort files into Weak, Medium and Strong 
(based on approx fit parameters)

plot_peak_params(), filter_splitting_prominence(), 
identify_diad_groups(), plot_diad_groups()

Figure 3: Schematic of the workflow used to efficiently fit diad peaks with very different spectral characteristics.

minutes. As well as correcting drift during a given session, Ne lines are also useful for variations between sessions, allowing continued use of a Raman densimeter months to years after it was calibrated [DeVitre et al. 2021]. An alternative approach to correct for instrument drift uses repeated analyses of a sample/capillary with known CO\textsubscript{2} density [Le et al. 2021; Wieser et al. 2021]. This approach is particularly useful on instruments without a Ne lamp, or with insufficient spectral resolution to quantify the narrow Ne peaks.

The Ne emission spectrum has a number of distinctive "lines", several of which lie close to the position of the CO\textsubscript{2} Fermi diad. The wavelength of the different Ne lines in air (\(\lambda_1\)) are converted into Raman shifts (\(\Delta\nu\)) in cm\textsuperscript{-1} using the excitation wavelength of the laser (e.g., \(\lambda_0=532.05\text{ nm}\); [Lin et al. 2007]):

\[
\Delta\nu = \frac{10^7}{\lambda_0(\text{nm})} - \frac{10^7}{\lambda_1(\text{nm})}
\]

Traditionally, a measured diad splitting has been corrected for instrument drift and non-linearity by comparing the measured distance between two selected Ne lines (\(\Delta_{\text{Measured}}\)) and the theoretical distance between those lines using Equation 5 (\(\Delta_{\text{Theoretical}}\)):

\[
\text{Correction Factor} = \frac{\Delta_{\text{Theoretical}}}{\Delta_{\text{Measured}}} \quad (6)
\]

This correction factor is then multiplied by the measured diad splitting:

\[
\Delta_{\text{Corrected}} = \text{Correction Factor} \times \Delta_{\text{Measured}} \quad (7)
\]

This approach has been termed the 'line segment' technique by Bakker [2021], and a typical workflow of how this method is performed in DiadFit is summarized in Fig. 5. First, identify all the files which contain Ne lines based on a unique string in their name (e.g. 'Ne'). Next, the function calculate_Ne_line_positions is used to calculate the wavenumber of each Ne line using the specific wavelength of the excitation laser, and a threshold intensity to filter out weaker lines:
Identifying and fitting peaks from other phases

Get filenames

\texttt{get_diad_files()}

Inspect all files in a specific spectral region

\texttt{generic_peak_config(), plot_secondary_peaks()}

OR 2 options to ID peak positions

Plot spectra to visually assess peak presence

Peaks

see step 3

Identifying and fitting peaks from other phases

Get filenames

\texttt{get_diad_files()}

config peak params for each phase/distinct peak

\texttt{generic_peak_config(), fit_generic_peak()}

Carbonate peak - PseudoVoigt

SO$_2$ peak - spline

Loop through all files using these tweaked parameters

\texttt{get df of peak parameters}

Merge with diad peak fitting parameters. Calculate molar proportions using instrument factors.

Figure 4: Schematic of workflow used to fit secondary peaks (e.g., carbonate, SO$_2$). Secondary peak fitting results are then merged with diad parameters (allowing calculation of molar proportions of gas species).

\begin{verbatim}
df_Ne=pf.calculate_Ne_line_positions(
    wavelength=531.885, cut_off_intensity=2000)
\end{verbatim}

This returns a dataframe, with the Raman shift (wavenumber) expected for each Ne line position for your specific wavelength (See Fig. 5 Step 2).

Two Ne lines are selected (1117 cm$^{-1}$ and 1447 cm$^{-1}$ in this example), and the theoretical splitting is calculated for the closest lines to those specified wavenumbers. Next, one representative Ne acquisition file is selected, and used to tweak SciPy \texttt{find_peaks} parameters stored in the class \texttt{Neon_id_config} to identify the approximate position of each Ne line (and its intensity) by identifying the largest peak within ±10 datapoints of the Ne line of interest (e.g. for a spectra resolution of 0.4 cm$^{-1}$ and for line 1117, the spectral region 1113-1121 cm$^{-1}$ would be searched). Obtaining approximate peak positions and intensities greatly improves the computational efficiency of the fitting process.

After identifying approximate peak positions for each file, peak fitting parameters are tweaked in the class \texttt{Ne_peak_config}. The peak fitting function fits a polynomial background to two regions either side of the Ne line, discarding points within this region which are outside of a certain number of standard deviations of the median background position (default 3σ, can be tweaked in \texttt{Ne_peak_config}). This background is then subtracted. For most Ne lines, a single PseudoVoigt/Voigt curve can be fitted to background-subtracted data (See ~1453 cm$^{-1}$ peak, Fig. 5). For the line at ~1122 cm$^{-1}$, two peaks need to be fitted iteratively because of the prominent shoulder to the left of this line (red and cyan curves, combined into the green curve, Fig. 5).

In general, the default peak fitting values stored in a dataclass \texttt{Ne_peak_config} generally only need tweaking once for each Ne line on each instrument, because repeated Ne acquisitions are so similar. The most important parameters to change when swapping between lines are the positions of the background. These are expressed in terms of distance from the peak center identified above. For example, specifying \texttt{lower_bck_pk1=(-40, -25)} positions a background window 40-25 wavenumbers to the left of the pixel with maximum intensity. Other important parameters include the degree of polynomial to fit between background points \texttt{X_poly_pk1_baseline=2}, and the approximate sigma of the PseudoVoigt/Voigt curve \texttt{(pk1_sigma=0.6, pk2_sigma=0.7)}. These tweaked peak fit parameters are then used to fit one spectra file with the function \texttt{fit_Ne_lines}. This function returns the peak positions and other fit parameters, as well as...
Correcting for instrument drift with Ne

Input laser λ, calculate positions of Ne lines above a threshold intensity

calculate_Ne_line_positions()

Get filenames to fit

get_files()

Enter 2 approx line positions, calculated theoretical splitting

calculate_Ne_splitting()

Tweak peak fitting parameters for 2 selected lines:

Ne_peak_config()

• Background (bck) positions
• Degree of bck poly
• # of peaks to fit (e.g., 2 for 1117 line)
• Approx σ of Ne lines

Figure 5: Schematic showing how DiadFit can be used to fit Ne lines, and build a model of correction factor vs. time with full propagation of uncertainty.

graphs showing the overall best fit, the residual of the fit, and the background positions.

After inspecting these graphs for one Neon acquisition and tweaking any fit parameters as necessary (e.g., background positions, plotting parameters), the tweaked find peaks and peak fit parameters can then be used to loop through all Ne spectra. On a $1000 laptop (e.g., 16 GB RAM, Intel i7), it takes approximately 0.2 s to fit each Ne file. There is also an option for the function to save a figure showing the fit for each file in a subfolder it creates within the spectra path. This increases the run time by a factor of 5 for each file, but allows users to check the fits, or publish fits as part of a data repository accompanying a paper. After all files are looped through, DiadFit returns a pandas dataframe with the filename, and all the peak fit parameters (including the all important Ne correction factor).

In most published studies, Fermi diad acquisitions have been corrected using the average correction factor of the Ne line acquired before and after a specific sample (although occasionally concurrent acquisitions are used). However, the relatively narrow shape of the Ne peaks means that the error on the peak center of each line, and thus the Ne correction factor, is relatively large (see error bars on Ne correction factor on Fig. 5). Thus, we suggest it may be better to perform a regression of the correction factor as a function of time to average out this random peak fitting noise. On many instruments (e.g., WITec, some HORIBAs), the spectral file doesn’t include a useful time stamp (and the file write/edit date may be the time of export, not acquisition). Sometimes (e.g., WITec), the timestamp is stored in the metadata file, meaning that data and metadata files can be stitched together to obtain the time for each Ne correction factor (see documentation for examples). In this instance, DiadFit is designed to work best when spectra and metadata files have the same name - if users wish to use different names, they will have to adapt the automatic stitching functions. If a metadata file is not available, DiadFit contains functions to extract the time based on the read or edit time stamp on the spectra file. Once a DataFrame is obtained that contains both the Ne correction factor and the time, it is useful to inspect changes in Neon correction factors and peak positions with time using built-in visualization functions (e.g., plot_Ne_corrections). This allows outliers to be discarded that differ substantially from adjacent acquisitions using the function filter_Ne_lines_neighbours. Finally, a polynomial or spline model is used to parameterize the change in Ne correction factor as a function of time (expressed as seconds after midnight, Fig. 5), along with the associated confident interval. This model is saved, so that it can be loaded when...
7 EQUATION OF STATE CALCULATIONS

An equation of state (EOS) describes the relationship between physical properties of a system such as pressure, temperature and density. EOS calculations are widely used in volcanology - they allow conversion of fluid inclusion densities into pressures, conversion of homogenization temperatures from microthermometry into CO₂ densities, and calculations of densities inside optical cells widely used to calibrate Raman spectrometers. At present, DiadFit supports calculations using pure CO₂ and mixed CO₂-H₂O EOS. In future, we could implement additional EOS upon request from users (so check ReadTheDocs).

7.1 Pure CO₂ EOS

DiadFit currently supports calculations using the EOS for pure CO₂ using the model of Sterner and Pitzer [1994] (hereafter SP94) and Span and Wagner [1996] (hereafter SW96). While the SP94 EOS is hard coded directly into DiadFit, we perform calculations using the SW96 EOS through the Python package CoolProp [Bell et al. 2014], which needs to be installed to use the DiadFit functions relying on this EOS. There are three core functions used to perform EOS calculations in DiadFit:

1. calculate_rh0_for_P_T: calculates CO₂ density for a known pressure and temperature.
2. calculate_P_for_rh0_T: calculates pressure for a known CO₂ density and temperature.
3. calculate_T_for_rh0_P: calculates temperature for a known pressure and CO₂ density.

7.1.1 Calculating CO₂ density for a given P and T

The function calculate_P_for_rh0_T can be used to calculate CO₂ density for a specified Pressure (P) and Temperature (T). This calculation is very useful when processing data collected from Raman gas cell calibration apparatus, where T is measured by a thermistor and P is measured with a pressure transducer in the fluid within the optical cell [DeVitre et al. 2021]. An entire spreadsheet of pressures and temperatures can be imported and all densities calculated:

```
df=pd.read_excel('Cali_Data.xlsx')
dens_SW96=pf.calculate_rh0_for_P_T(P_kbar=df['P_kbar'], T_K=df['T_K'], EOS='SW96')
```

7.1.2 Calculating pressures from CO₂ densities and temperatures

The function calculate_P_for_rh0_T can be used to calculate pressure for a specified CO₂ density (rho) and temperature (T). This calculation is commonly used to calculate fluid inclusion pressures. This pressure may represent the entrapment pressure under the assumption that the volume and mass of the inclusion is fixed from the [Roedder 2018], or a re-equilibration pressure if the inclusion underwent prolonged stalling [Hansteen 1991].

To convert a CO₂ density of 0.5 g/cm³ into a pressure at a temperature of 1200 K using the Span and Wagner [1996] EOS (SW96):

```
P_SW96=pf.calculate_P_for_rho_T(CO2_dens_gcm3=0.5, T_K=1200, EOS='SW96')
```

The Sterner and Pitzer [1994] EOS (SP94) can be used instead simply by changing the EOS argument:

```
P_SP94=pf.calculate_P_for_rho_T(density_gcm3=0.5, T_K=1200, EOS='SP94')
```

In addition to single calculations, this function can also be applied to a pandas dataframe with as many rows as the user wants. For example, to load an entire spreadsheet with columns for CO₂ densities and an estimate of entrapment temperatures in °C (as well as an optional column with the sample name):

```
df=pf.read_excel('FI_densities.xlsx')
P_SP94=pf.calculate_P_for_rho_T(T_K=df['Temp in C']+273.15, CO2_dens_gcm3=df['Density_gcm3'], EOS='SW96', Sample_ID=df['Sample'])
```

This returns a pandas dataframe with calculations done for each input row.

7.1.3 Comparing two EOS

DiadFit makes it very simple to compare calculations using the EOS of Sterner and Pitzer [1994] and Span and Wagner [1996]. In Fig. 6, we calculate pressure for 11 discrete densities between 0.1 and 1.1 g/cm³ and temperatures between 34 and 2000 °C (see ReadTheDocs for a worked example). These calculations would have been tedious in existing tools, but can be performed in several lines of code in DiadFit (and in seconds of computational time). This figure demonstrates that the EOS are extremely similar at temperatures relevant to basaltic magmas (<2% discrepancies).

7.1.4 Converting homogenization temperatures to CO₂ densities

Microthermometry provides an alternative method to Raman spectroscopy to determine the density of CO₂-rich fluids. A fluid inclusion is cooled down, and then slowly heated up to determine the temperature at which a two-phase inclusion containing liquid and vapour transitions to a single homogeneous phase (the homogenization temperature). CO₂ densities can be calculated from these homogenization temperatures using the function calculate_CO2_density_homog_T, which
Figure 6: Comparison of the CO$_2$ EOS of Sterner and Pitzer [1994] and Span and Wagner [1996]. The y axis on b) shows the pressure calculated by SP94 minus that from SW96 divided by the average of these two pressures.

Figure 6: Comparison of the CO$_2$ EOS of Sterner and Pitzer [1994] and Span and Wagner [1996]. The y axis on b) shows the pressure calculated by SP94 minus that from SW96 divided by the average of these two pressures.

This returns a DataFrame showing the bulk density, the density of the co-existing liquid and vapour just before homogenization, and the user-inputted parameters.

An entire excel sheet of homogenization temperatures and phases can be loaded as a DataFrame. There is also an optional input of entering a sample name for each row (so it is returned in the outputted DataFrame). Uncertainties in homogenization temperature can be propagated to determine errors in density using Monte-Carlo techniques (see example on ReadTheDocs).

7.2 CO$_2$-H$_2$O EOS

In reality, many igneous systems will have some proportion of H$_2$O in the exsolved fluid phase, and thus the entrapped fluid inclusion. In this scenario, entering CO$_2$ densities measured by Raman spectroscopy or microthermometry into a pure CO$_2$ equation of state to calculate pressure is problematic, and a mixed CO$_2$-H$_2$O EOS must be used instead. However, detecting the presence of H$_2$O in fluid inclusions, let alone quantifying H$_2$O mole proportions required to perform calculations using a H$_2$O-CO$_2$ EOS, is very challenging for a number of reasons [Morgan et al. 1993]. First, H$_2$O may no longer be present in the inclusion, because of diffusive loss during stalling, ascent and syn-eruptive quenching, or because it reacted with the mineral host to form hydrous phases [Morgan et al. 1993; Mavrogenes and Bodnar 1994; Zanon et al. 2024]. If H$_2$O is still present, H$_2$O has a very low solubility in H$_2$O fluids at temperatures typical of routine Raman analyses (~ 0.5 mol%, [Spicher et al. 2003; Frezzotti and Peccerillo 2007]). The small proportions of molecules that do dissolve will be characterized by a weak and relatively narrow (15 cm$^{-1}$) peak at ~ 3640 cm$^{-1}$ that may not be detectable [Wopenka and Pasteris 1987; Frezzotti and Peccerillo 2007; Berkesi et al. 2009]. The extremely thin nature of this film makes it very challenging to detect optically or with Raman spectroscopy. Another issue is that at room temperature, peaks resulting from H$_2$O (dissolved or in films) are at 3000-3800 cm$^{-1}$ [Frezzotti et al. 2012], which is not in the typical spectral window used when measuring CO$_2$ at high spectral resolution, requiring a separate acquisition to detect them. Heating the fluid inclusion to ~ 120-150 °C, where the solubility of dissolved H$_2$O is far higher, results in a larger peak, which can be more easily identified/quantified [Berkesi et al. 2009]. However, perhaps because the H$_2$O has been lost in many cases, and even with heating, quantification is challenging, the majority of fluid inclusion studies simply pick a fixed value of X$_{H_2O}$ (e.g., 0.1 for [Sandoval-Velasquez et al. 2023], [Zanon et al. 2024]. [Forte et al. 2023]). More recently, DeVitre and Wieser 2024 determine X$_{H_2O}$ as a function of pressure for the volcanic system of interest using melt inclusion measurements (many solubility models used to calculate saturation pressures also calculate X$_{H_2O}$ at the point of vapour saturation, e.g., [Ghiorso and Gualda 2015]).
To calculate pressures for mixed fluids, an assumption also must be made about whether H$_2$O was lost or remained in the fluid inclusion at the time of measurement. By default, DiadFit calculates the pressure for both scenarios. For the calculation assuming H$_2$O was lost prior to measurement, DiadFit follows the correction method of [Hansteen and Klügel 2008], where the measured CO$_2$ density is corrected back to a bulk density of the trapped CO$_2$-H$_2$O mix (where 44 is the molar mass of CO$_2$, 18 of H$_2$O):

$$\alpha = X_{H_2O}/(1 - X_{H_2O})$$

$$\rho_{bulk} = \rho_{CO_2, meas}(1 + \frac{\alpha}{44})$$

If H$_2$O is assumed to still be present in the fluid inclusion as a thin film of liquid around the edge, the bulk density at the time of trapping can be calculated as follows (assuming no volume changes). Following the law of partial pressures:

$$P_{Tot} = P_{CO_2} + P_{H_2O}$$

$$P_{CO_2} = P_{Tot}X_{CO_2}^{mol}$$

$$P_{H_2O} = P_{Tot}X_{H_2O}^{mol}$$

The pure CO$_2$ EOS is used to calculate the pressure of the CO$_2$ fluid ($P_{CO_2}$) from the measured density and analysis temperature (Fig. 7c). Re-arranging equation 11 allows calculation of $P_{Tot}$. Then equation 12 is used to calculate $P_{H_2O}$ for a fixed value of $X_{mol}^{CO_2}$. Unlike the scenario where H$_2$O is lost, this calculation requires knowledge of the analysis temperature, which is used alongside $P_{H_2O}$ to calculate $\rho_{H_2O}$ using the pure H$_2$O EOS of Wagner and Prüß [2002] (implemented in [Bell et al. 2014]). By converting $X_{mol}^{H_2O}$ to $X_{H_2O}^{max}$, the volume of CO$_2$ and H$_2$O phases can be calculated from their densities. By re-arranging, $\rho_{bulk}$ can be calculated, which can then be entered in a CO$_2$-H$_2$O EOS (Fig. 7c).

DiadFit performs calculations using the CO$_2$-H$_2$O of [Duan and Zhang 2006]. This EOS was implemented in DiadFit by translating and adapting the C code of Yoshimura [2023] for the interaction coefficients. In DiadFit, pressure or molar volume are solved iteratively from Eq 9 of Duan and Zhang [2006] using a Newton-Raphson method. At certain bulk densities and $X_{H_2O}$ values, a negative pressure is converged upon during iteration. To resolve this issue, and other convergence issues, we use a generate a look up table of molar volumes for 40 temperatures (T = 300-2000 K), 40 $X_{H2O}$ values (0-1), and 40 pressures (P=1-100 kbar), yielding 62,000 look up values for all possibly combinations of these (40x40x40). When a user enters a bulk density, this is first converted to molar volume, and then the closest match to the entered conditions in the look up table is identified, and the corresponding pressure is used as the start point for the algorithm. This resolves convergence issues, and speeds up computation time.

Overall, to convert fluid inclusion CO$_2$ densities into pressures using the CO$_2$-H$_2$O EOS, users should use the function calculate_entrainment_pressure, specifying measured CO$_2$ densities, entrapment temperature, analysis temperature, and $X_{H2O}^{mol}$. The function returns the pressure for the mixed EOS for the H$_2$O loss and no H$_2$O loss scenario, and the calculated bulk densities. It also returns pressures for the pure CO$_2$ EOS of SW96, SP94 and Duan and Zhang [2006] for comparison. Unlike existing tools, this allows pressures to be calculated for hundreds of fluid inclusions very quickly, using different $X_{H2O}$ for each inclusion if so desired. This function can also be used to visualize the magnitude of the correction factor for different CO$_2$ densities and molar proportions of H$_2$O in the fluid (e.g., Fig. 8).

We also include the function calc_prop_knownP_EOS_DZ2006 for calculating molar volumes, compressibility factors, fugacity and activities of each species using the Duan and Zhang [2006] EOS where pressure, temperature and $X_{H2O}$ are known. We anticipate this will be useful for experimentalists and/or those building solubility models or other parameterizations requiring this thermodynamic data.

8 ADDITIONAL CALCULATIONS RELATING TO FI BAROMETRY

8.1 Converting Pressures into depths

Building on functionality in Thermobar [Wieser et al. 2022], DiadFit can convert pressures to depths using various options (Fig. 1h) implemented in the function convert_pressure_to_depth:

- A fixed crustal density (e.g., $\rho$=2700 kg/m$^3$)
- A 2-step or 3-step crustal density profile (e.g., $\rho$=2700 kg/m$^3$ at <10 km, $\rho$=3300 kg/m$^3$ at >10 km)
- Pressure-depth models of:
  1. A crustal profile for arc magmas from Rasmussen et al. [2022] (‘rasmussen’).

We will add more density profiles as they are published, so we encourage users to check the documentation for more details.

8.2 Modeling fluid inclusion re-equilibration

DeVitre and Wieser [2024] implement a Python3 version of the mechanical re-equilibration model for olivine-hosted fluid inclusions of Wanamaker and Evans [1989] in their package RelaxIF. We incorporate this methodology into DiadFit for ease of installation and maintenance, because it uses the underlying EOS and density-depth conversions from DiadFit to track changes in fluid inclusion volume, density, and pressure for different stalling, ascent and quenching paths. Worked examples are available on ReadTheDocs. If this model is
Using a pure CO\textsubscript{2} fluid

\[ P_{\text{entrap}} = 155.38 \text{ MPa (Duan + Zhang 2006)} \]

\[ P_{\text{entrap}} = 159.84 \text{ MPa (Sterner+Pitzer, 1994)} \]

\[ P_{\text{entrap}} = 155.97 \text{ MPa (Span+Wagner, 1996)} \]

\begin{align*}
\rho_{\text{CO}_2, \text{meas}} &= 0.4 \text{ g/cm}^3 \\
\rho_{\text{CO}_2, \text{bulk}} &= ?
\end{align*}

Using a pure CO\textsubscript{2} EOS at 37°C:

\[ P_{\text{CO}_2} = 8.26 \text{ MPa} \]

Law of partial pressures:

\[ P_{\text{tot}} = X_{\text{CO}_2, \text{mol}} \cdot P_{\text{CO}_2} \]

\[ P_{\text{H}_2\text{O}} = 74.35 \text{ MPa} \]

Using a pure H\textsubscript{2}O EOS at 37°C:

\[ \rho_{\text{H}_2\text{O}} = 0.9936 \text{ g/cm}^3 \]

Given mass (m) / Volume (V) = density (ρ)

\[ V_{\text{CO}_2} = \frac{m \cdot (1 - X_{\text{H}_2\text{O}, \text{mol}})}{\rho_{\text{CO}_2}} \]

\[ V_{\text{H}_2\text{O}} = \frac{m \cdot X_{\text{H}_2\text{O}, \text{mol}}}{\rho_{\text{H}_2\text{O}}} \]

\[ \rho_{\text{bulk}} = \frac{\rho_{\text{H}_2\text{O}} \cdot V_{\text{H}_2\text{O}} + \rho_{\text{CO}_2} \cdot V_{\text{CO}_2}}{0.4107 \text{ g/cm}^3} \]

Using Duan and Zhang (2006) EOS

\[ P_{\text{entrap}} = 173.7 \text{ MPa} \]

\begin{align*}
\alpha &= X_{\text{H}_2\text{O}, \text{mol}} / (1 - X_{\text{H}_2\text{O}, \text{mol}}) = \text{H}_2\text{O/CO}_2 \text{ molar ratio} \\
\rho_{\text{H}_2\text{O, \text{ass}} + \text{CO}_2} &= 0.4 \text{ g/cm}^3 \\
\rho_{\text{H}_2\text{O}} &= (1 + 0.0003 \text{ molar ratio}) \rho_{\text{H}_2\text{O, \text{bulk}}}
\end{align*}

DiadFit provides a framework for propagating the three main sources of analytical uncertainty when quantifying CO\textsubscript{2} density by Raman spectroscopy:

1. Peak fitting error when determining the peak center of each diad peak (Fig. 10a-b).
2. Error associated with correcting for instrument drift (Fig. 10a-c).
Figure 9: Screenshot of the output from `pf.stretch_in_ascent` for isothermal ascent (1200 °C) towards the surface for a FI with an initial radius of 1 μm for an ascent rate of 1 m/s. The returned figure shows the fractional change in the fluid inclusion radius as a function of depth (e.g., it gets bigger as it ascents), and the accompanying drop in CO₂ density caused by the larger FI volume. The returned `pandas.DataFrame` shows changes in internal and external pressure, fluid inclusion radius, along with other parameters.

3. Error associated with the densimeter used to convert drift-corrected splitting into CO₂ density (Fig. 10d).

DiadFit obtains the 1σ error associated with each peak center from `lmfit` (which uses the covariance matrix). As each peak is fitted independently because the Diad1 and Diad2 peaks do not overlap, the error associated with the position of each peak center is combined in quadrature to get the error on the peak splitting. The Ne line correction factor vs time regression model (Fig. 5) is used to determine the error on the Ne correction factor for each acquisition (Fig. 10c). This error is combined with the splitting error in quadrature to get the uncertainty in the corrected splitting (black crosses, Fig. 10a). To propagate the uncertainty associated with the densimeter, DiadFit uses the regression model through the calibration data. This error is combined with the error on the corrected splitting to yield the overall error in density (σ Density g/cm³, Fig. 10d). Assessing the relative contribution from each source of uncertainty helps optimization of analytical routines (Fig. 10). If the majority of the error is being introduced at the peak fitting stage, it may be that longer acquisitions are required to get stronger signals, or a higher resolution grating is required. Fig. 10a,f shows that a few acquisitions at ~0.18–0.2 g/m³ have far larger peak fitting errors than other acquisitions from the same analytical session. These anomalously large errors would justify inspection of the spectra and fits in more detail, and perhaps re-analysis.

9.2 Uncertainty in fluid inclusion pressures

DiadFit can be used to propagate uncertainty when calculating pressures and depths from fluid inclusions based on Raman and microthermometry measurements. The function `propagate_FI_uncertainty` allows propagation of uncertainties in input parameters using Monte Carlo methods. The function input arguments are the magnitude and distribution of uncertainty for CO₂ density, temperature, the mol fraction of H₂O (X_{H₂O}), and the crustal density. Input errors for each variable can be absolute (e.g., ±50 K for temperature) or percentage errors (e.g., ±5%), and can follow a normal or uniform distribution. For example, a user may specify that the uncertainty in their temperature is best approximated with a normal distribution with a standard deviation of 50 K. For each fluid inclusion, the function generates N duplicates of each input parameter randomly drawn from these specified prior distributions. It is assumed that all input parameters are independent of each other. Pressure and depths are calculated for each of these N duplicates. The function outputs a figure showing the simulation for a single specified fluid inclusion (here, FI1, Fig. 11), in addition to two dataframes. The first dataframe shows the mean, median, standard deviation of calculated pressures and depths for each fluid inclusion (Fig. 11, Output1). For skewed distributions, the standard deviation may be best approximated by half the difference between the 84th - 16th percentile (also provided). The second dataframe showing the results for all simulations - if N=1000 duplicates
Figure 10: Assessing different sources of analytical error. a) The black cross shows the total error on the corrected splitting from propagating the error in fitting CO₂ peaks and Ne correction model for an instrument with a spectral resolution of 0.56 cm⁻¹. Individual contributions from each peak and the Ne correction model are shown. Note, the total error is smaller than the sum of each error shown, as a result of propagating these errors in quadrature. b) To allow easier comparison of the proportional contribution from each source of error, we also take each individual error and divide by the total error on the corrected splitting. This show that the fitting error on Diad1 is the largest source of uncertainty. c) Ne regression model with uncertainty represented by red-dashed lines, along with the uncertainty for a single diad acquisition. d) Error associated with the densimeter shown with dashed lines, with the underlying acquisitions shown as dots. The densimeter is regressed in two parts for low and medium densities (see DeVitre et al. 2021). e) Percentage % error on calculated density, showing the contribution from errors on the corrected splitting (see a) and from the densimeter. f) 1σ errors on density and corrected splitting.
and N=10 fluid inclusions, the dataframe has 10,000 rows (Fig. 11, Output2).

9.3 Uncertainty in equivalent CO2 contents of vapour bubbles
The Raman method of reconstructing vapour bubbles present within melt inclusions relies on measuring the CO2 density, and then converting this into an equivalent amount of CO2 that would have been dissolved in the glass at the time of melt inclusion entrapment by mass balance:

\[ CO_2^{\text{equivalent glass ppm}} = 10^3 \frac{V\text{Rvol}}{\rho_{\text{melt}}} \cdot \rho_{\text{CO}_2} \]  \hspace{1cm} (13)

Where \( \rho_{\text{CO}_2} \) is the density of CO2, \( \rho_{\text{melt}} \) is the density of silicate melt, and \( V_{\text{Rvol}} \% \) is the Volume % occupied by the vapour bubble.

Similar to the fluid inclusion method described above, DiadFit contains functionality to propagate the uncertainties in each of these variables using Monte-Carlo methods. The function `propagate_CO2_in_bubble` allows users to specify the magnitude of the error for each parameter, whether the error is a % error or an absolute error, and whether the error is normally or uniformly distributed. This function returns two dataframes - the first with the average and standard deviation for each melt inclusion, and the second showing all simulation outputs. The uncertainty in CO2 density can be obtained from DiadFit. The uncertainty in melt density can be obtained from Iacovino and Till [2019]. Uncertainty in estimating bubble volumes depend greatly on the method used and the geometry of inclusions (e.g., [DeVitre et al. 2023b], [Tucker et al. 2019]).

10 Other useful functions for volcanological workflows
10.1 Quantifying peak asymmetry/skewness to identify coexisting liquid and gaseous CO2
At room temperature (18-22 °C), a CO2 fluid with a density between ~0.18–0.21 and 0.75–0.79 g/cm³ will consist of an inner shell of CO2 vapour, and an outer shell of CO2 liquid [Span and Wagner 1996]. Using the `calculate_CO2_homog_T` function, we can calculate that at 20 °C the gaseous phase will have a density of 0.194 g/cm³ and the liquid will have a density of 0.773 g/cm³. DeVitre et al. [2023a] show that Raman measurements performed in inclusions with two coexisting phases at room temperature don’t always show two distinct peaks for the gaseous and liquid CO2 phase, but often show a single skewed peak, with contributions from both phases. Quantifying peak asymmetry can help to identify such spectra, which cannot be reliably quantified for CO2 density.

To assess asymmetry in each spectra, an \( N^{th} \) degree polynomial is fitted between specified baseline positions, and this background is subtracted. A cubic-spline is fitted between the baseline positions. The x and y coordinates of the highest point on this spline fit are identified. The position on each shoulder of the peak with an intensity equal to a certain fraction of the peak height is identified. In Fig. 12, this cut off parameter \( \text{intcutoff} \) is set at 0.3. The ratio of the x-distance between the peak center and these shoulder points (green and grey lines in Fig. 12) defines the peak asymmetry.

After tweaking these parameters for one spectra, the function `loop_diad_skewness` can be used to loop through all files in a specific folder, and stitch the results into a dataframe. Typical skewness parameters have to be identified for each instrument, after which spectra with high skewness can be identified, and reanalysed about the critical temperature where a single peak will be present (see [DeVitre et al. 2023a]).

10.2 H2O fitting
It is well established that Raman spectroscopy can be used to quantify H2O within silicate glasses. The glass alumino-silicate region consists of multiple bands; a low frequency/low wavenumber (LF or LW) band centered on ~550 cm⁻¹, a medium frequency/wavenumber (MF, MW) band centered at ~550 cm⁻¹, and a high frequency/wavenumber (HF, HW) band at ~1000 cm⁻¹ (Fig. 13, see [van Gerve and Namur 2023] for detailed description of the relevant vibrations). The H2O region is characterized by a broad, asymmetrical peak centered at \( \sim 3500 \) cm⁻¹. There are many different methods to use these different peaks to quantify H2O, some use the height or intensity of the water Raman band (~3000-3800 cm⁻¹), and others using the ratio of the area under the water region divided by the area under the alumino-silicate band at ~200–1250 cm⁻¹ [Thomas 2000; Zajacz et al. 2005; Behrens et al. 2006; Thomas et al. 2006; Mercier et al. 2010; Le Losq et al. 2012; Di Genova et al. 2017; Schiavi et al. 2018].

The exact relationship between peak areas/ratios/heights and the H2O content depends on instrument hardware and the glass major element composition, so standards with known H2O contents should be used for calibration (see [Schiavi et al. 2018] and refs within).

In terms of quantification strategies, Behrens et al. [2006] provide two empirical equations relating silicate:water areas to absolute water contents. One equation is for albite, haplogranite and dacite samples, and the other is for intermediate and depolymerized compositions. Le Losq et al. [2012] build on this to produce a global empirical calibration. Importantly, they note that different silicate glass compositions require different background positions prior to quantifying silicate peak areas. Schiavi et al. [2018] refine this method further, noting that as well as being sensitive to glass composition, the baseline is also sensitive to reduct state. They suggest fitting a default cubic baseline through predefined baseline positions for 4 melt compositions (basalt, basanite, andesite, rhyolite) results in the area ratio (LW+HW+MW to H2O area) being reasonably independent of melt composition. However, Gonzalez-Garcia et al. [2021] note that the presence of nanolites with a peak at 670-690 cm⁻¹ complicate total silicate area methods, because this nanolite region overlaps with the LW silicate region used in the ‘total area method’ of Schiavi et al. [2018]. They propose a new protocol using only the HW area to characterize the silicate portion, rather than the overall silicate region as in Schiavi et al. [2020].

To allow maximum flexibility when using these different methods, DiadFit quantifies three different silicate band areas, and the overall silicate and water area (Fig. 13), with lots of options for users to tweak background positions, under the assumption protocols will continue to change. When fitting
Outputs

Output 1: a dataframe with the average for each FI

<table>
<thead>
<tr>
<th>Sample</th>
<th>CO₂ dens g/cm³</th>
<th>Single FI D_km</th>
<th>Single FI P_kbar</th>
<th>Mean MC P_kbar</th>
<th>Med MC P_kbar</th>
<th>std dev MC P_kbar</th>
<th>std dev MC P_kbar from percentile</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>0.436888</td>
<td>6.765613</td>
<td>1.792008</td>
<td>1.797897</td>
<td>1.788961</td>
<td>0.183864</td>
<td>0.185257</td>
</tr>
<tr>
<td>F2</td>
<td>0.417796</td>
<td>6.119572</td>
<td>1.60891</td>
<td>1.635096</td>
<td>1.620293</td>
<td>0.169176</td>
<td>0.164091</td>
</tr>
<tr>
<td>F3</td>
<td>0.459045</td>
<td>10.899166</td>
<td>1.924561</td>
<td>1.927098</td>
<td>1.926690</td>
<td>0.194317</td>
<td>0.188625</td>
</tr>
</tbody>
</table>

Output 2: A dataframe showing the results for all N simulations for each FI

<table>
<thead>
<tr>
<th>Sample</th>
<th>Pressure (kbar)</th>
<th>Pressure (MPa)</th>
<th>Depth (km)</th>
<th>input crust dens g/cm³</th>
<th>MC_T_K</th>
<th>MC CO₂ dens g/cm³</th>
<th>MC_XH2O</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>1.937356</td>
<td>193.735604</td>
<td>6.999241</td>
<td>282.151.6108</td>
<td>None</td>
<td>128.371368</td>
<td>0.461293</td>
</tr>
<tr>
<td>F2</td>
<td>2.046047</td>
<td>204.604678</td>
<td>7.462096</td>
<td>2795.025232</td>
<td>None</td>
<td>1251.033969</td>
<td>0.466559</td>
</tr>
<tr>
<td>F3</td>
<td>1.765151</td>
<td>176.515067</td>
<td>6.203195</td>
<td>2900.663361</td>
<td>None</td>
<td>1338.388635</td>
<td>0.431808</td>
</tr>
<tr>
<td>F4</td>
<td>1.847866</td>
<td>184.786569</td>
<td>7.650040</td>
<td>2462.281330</td>
<td>None</td>
<td>1340.861811</td>
<td>0.419261</td>
</tr>
</tbody>
</table>

Output 3: A figure for one FI

Simulations for sample = F1

- Input distribution Temp: Normally-distributed, 1σ = 30 K
- Input distribution CO₂ density: Normally-distributed, 1σ = 0.02 g/cm³
- Input distribution crustal density: Normally-distributed, 1σ = 5%
- Input Distribution XH2O: Normally-distributed, 1σ = 0.05 kg/m³

Figure 11: Schematic showing the workflow used to propagate uncertainty in CO₂ density, Temperature, and crustal density into error distributions in pressure and depth for each fluid inclusion.
Peak Asymmetry

Get filenames to fit, select 1 test file
get_files()

Tweak peak fit parameters+intensity cut off
diad1_fit_config(), diad2_fit_config()

Inspect fit to test file, tweak if necessary
assess_diad1_skewness(), assess_diad2_skewness()

Loop through all files using these parameters
loop_diad_skewness()

spectra acquired on hydrous glasses in DiadFit, the first step is to select the files of interest. As for diad peaks and Ne lines, default peak fit parameters are stored in dataclasses. There are dataclasses for the four suggested background positions of Schiavi et al. [2018], e.g., sil_bck_pos_Schiavi_basalt, sil_bck_pos_Schiavi_andesite. Users can tweak these background positions for their specific samples if they wish (e.g., for a basaltic andesite). For example, the lower background position can be easily changed for basalt from the default of 300–340 to 320-350 cm<sup>-1</sup>:

```
pf.sil_bck_pos_Schiavi_basalt(
    lower_range_sil=[320, 350])
```

These dataclasses also store options for baseline fitting. After removing baseline points outside of a specified sigma, either a polynomial of degree N or a cubic spline can be fit to all the datapoints within the baseline region. After subtracting away this polynomial, the area under the background-corrected spectra is calculated using the Simpson and the trapezoid method [Tallarida and Murray 1987] implemented in SciPy and NumPy respectively. Overall, the function function returns these areas, along with the ratio of the silicate to water areas for each area method (as well as other useful parameters such as the positions of chosen baselines, degrees of polynomials etc. for future reference).

Figure 12: Schematic showing how DiadFit can be used to assess diad assymetry/skewness following the method of De-Vitre et al. [2023a].
H$_2$O fitting in silicate glasses

- Get filenames of glass spectra
  
  ```python
  get_files()
  ```

- Select peak fit parameters for silicate
  
  ```python
  e.g., sil_bck_pos_Schiavi_basalt(), sil_bck_pos_Schiavi_andesite() etc.
  ```

- Select peak fit parameters for H$_2$O
  
  ```python
  water_bck_pos()
  ```

- Fit 1 spectra and inspect fit to H$_2$O region
  
  ![H$_2$O fit](image)

- Fit 1 spectra and inspect fit to silicate region
  
  ![Silicate fit](image)

- Loop over all files if require similar parameters, or step through and fit 1 at a time, and stitch fits saved in .csvs together
  
  ```python
  stitch_dataframes_together()
  ```

Figure 13: Schematic showing how DiadFit can be used quantify the relative areas of silicate and H$_2$O peaks in Raman spectra collected from silicate glasses.

10.2.1 H$_2$O fitting within unexposed melt inclusions

A common criticism of the Raman method when quantifying the CO$_2$ content of vapour bubbles is that it does not ac-
count for the sequestration of carbon as solid carbonate phases on the inclusion walls (e.g., [Moore et al. 2015]). DeVitre et al. [2023b] developed a technique where melt inclusions with carbonate-bearing vapour bubbles are heated to their liquidus temperature using a Linkam TS1400XY stage, causing carbonate to redissolve as CO$_2$ which can then be measured by Raman spectroscopy. A concern with any heating method when working with melt inclusions is the possibility of diffusive loss of H$_2$O (Chen et al. 2011; Gaetani et al. 2012). DeVitre et al. [2023b] develop a method to assess H$_2$O loss by acquiring Raman spectra on unexposed melt inclusion glasses before and after homogenization.

Quantifying H$_2$O in spectra acquired from unexposed melt inclusions is more complicated than in exposed glasses, because the Raman signal contains a contribution from the glass and the overlying olivine host. The contribution from the host mineral must be subtracted to obtain an ‘unmixed spectra’, allowing reliable estimation of the silicate area. Fortunately, most magmatic olivine crystals have very low H$_2$O contents; typically a few 10s of ppm, [Neucombre et al. 2020; Toubin et al. 2023], and occasionally up to 100s of ppm, [Wang et al. 2022]. Thus, peaks in the H$_2$O region of the spectra are extremely weak/completely absent, even at laser powers far higher than is typical for glass analyses (70-140 mW, [Martinek and Bolfan-Casanova 2021], [Martinek and Bolfan-Casanova 2021]), which simplifies demixing of the signals. Here, we describe the demixing method used for olivine-hosted melt inclusions, although it could be adapted for other phases which do not produce significant peaks in the H$_2$O region. We note that demixing and deconvolution of glass from other signals is also available in the program SilicH2O [van Gerve and Namur 2023].

To reliably unmix the spectra, it is best to acquire a spectra in the melt inclusion at the depth where the H$_2$O peak is the strongest, and a spectra of the olivine next to the melt inclusion. DiadFit extracts the region of these two spectra between 800-900 cm$^{-1}$ containing the strong olivine doublet. A cubic spline is fitted to both spectra to smooth out noise. SciPy is used to find the positions of the two strong peaks in the olivine spectra, and the trough position between these peaks. DiadFit then creates N spectra, where the olivine spectra is subtracted from the mixed spectra. For each of these mixed spectra, a linear regression is fit between the two points with the wavenumbers of the identified olivine peaks. The vertical (y) distance is then calculated between this linear regression and the y value at the x-coordinate of the trough (Fig. 14). When too little signal from the olivine has been subtracted from the mixed spectra, the trough position will be lower than the linear regression (e.g., Dist=-400 in sketch 1 on Fig. 14, and when too much olivine has been subtracted, the peaks will invert, and the distance will be a positive number. When exactly the right amount of olivine has been subtracted, there will be no clear peaks or troughs, so the distance will be zero. The function fits a curve to this calculated distance vs. the mixing proportion, to determine the best-fit mixing proportion where the distance is zero. The resulting, ‘unmixed’ spectra is taken as the spectra from the melt inclusion itself. Then, the work-

11 FUTURE WORK

The open-source nature of DiadFit means that users can customize functions and build their own (either on a local fork, or using a pull request on GitHub). Additionally, we anticipate that we will continue to add workflow as new applications of Raman spectroscopy appear in volcanology. To reflect the evolving nature of this tool, when citing DiadFit, users should make sure they specify the version they used, obtained using:

```python
pf.__version__
```

Care should also be taken to cite the root packages used by DiadFit. E.g., if converting CO$_2$ densities to pressures using the functions here, we recommend wording similar to this: ‘CO$_2$ densities were converted into pressures using DiadFit (Wieser and DeVitre, 2024, v.1.0.5), using the Equation of State of Span and Wagner [1996] implemented in CoolProp [Bell et al. 2014].’

12 CONCLUSIONS

DiadFit is a high-level python package that provides easy-to-use functions for common workflows involving Raman spectroscopy and microthermometry (e.g. peak fitting, EOS calculations etc.), with particular applicability to the analysis of melt inclusions and fluid inclusions. DiadFit not only greatly reduces the time spent on data processing/reduction, it also includes numerous functions to propagate uncertainty. In turn, this will be vital to identify the biggest sources of uncertainty, and identify ways to minimise these errors in many difficult volcanological applications.

AUTHOR CONTRIBUTIONS

PW conceived the project and lead the code writting (apart from the functions discussed below). CD performed extensive testing and tweaking of existing functions. She also wrote the cosmic ray removal code, the fluid inclusion relaxation code, and the code to calculate molar proportions from gas peak areas. She conceptualized the peak asymmetry method, which PW implemented in python.

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Get filenames of spectra from MIs and neighbouring Ol crystals

extract_xstal_MI_name()

get_spectra(),
smooth_and_trim_around_olivine()

check_if_spectra_negative()

Extract region around ~800-900 cm\(^{-1}\) for both spectra.
Fit a smoothed cubic spline + ID peaks and troughs

Best match when y distance between 2 peaks and trough is as small as possible in unmixed spectra
MI-0.1* Ol
MI-0.9* Ol

check_if_spectra_negative()

Subtract varying proportions of Ol from the MI spectra.

Figure 14: Schematic showing how DiaDiFit can be used to unmix the contribution of olivine and glass from spectra taken on unexposed melt inclusions. The code could be easily adapted for other silicate phases. There is an option to loop if glass compositions are sufficiently similar. Alternatively, each file can be stepped through manually to tweak positions, and the code saves the output for each file to a .csv, and these are then stitched together after all files are fitted.
DATA AVAILABILITY

All files are available on GitHub https://github.com/PennyWieser/DiadFit. Documentation is available at ReadTheDocs (https://diadfit.readthedocs.io/en/latest/), and YouTube videos explaining various aspects of the tool are available on the DiadFit YouTube channel bit.ly/DiadFitYouTube
REFERENCES


