Probing the cosmological variation of the fine-structure constant:
Results based on VLT-UVES sample

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Abstract. Development of fundamental physics relies on the constancy of various fundamental quantities such as the fine-
structure constant. Detecting or constraining the possible time variations of these fundamental physical quantities is an im-
portant step toward a complete understanding of basic physics. High-quality absorption lines seen in the spectra of distant
QSOs allow one to probe time variations of several of these quantities. Here we present the results from a detailed many-
multiplet analysis, to detect the possible variation of the fine-structure constant, performed using high signal-to-noise ratio,
(∼70 per pixel), high spectral resolution (R ≥ 45 000) observations of 23 Mg II systems detected toward 18 QSOs in the redshift
range 0.4 ≤ z ≤ 2.3 obtained using UVES at the VLT. We validate our procedure and define the selection criteria that will avoid
possible systematics using a detailed analysis of a simulated data set. The spectra of Mg II doublets and Fe II multiplets are
generated considering variations in α and specifications identical to that of our UVES spectra. We show that our Voigt profile
fitting code recovers the variation in α very accurately when we use single component systems and multiple-component systems
that are not heavily blended. Spurious detections are frequently seen when we use heavily blended systems or systems with very
weak lines. Thus we avoided these system while analysing the UVES data. To make the analysis transparent and accessible to
the community for critical scrutiny, all the steps involved in the analysis are presented in detail. The weighted mean value of the
variation in α obtained from our analysis over the redshift range 0.4 ≤ z ≤ 2.3 is Δα/α = (∼0.06 ± 0.06) × 10−5. The median
redshift of our sample is 1.55 and corresponds to a look-back time of 9.7 Gyr in the most favored cosmological model today.
The 3σ upper limit on the time variation of α is −2.5 × 10−16 yr−1 ≤ (Δα/αΔt) ≤ +1.2 × 10−16 yr−1. To our knowledge this is the
strongest constraint from quasar absorption line studies to till date.

Key words. galaxies: quasars: absorption lines – cosmology: observations

1. Introduction

Contemporary theories of fundamental interactions, such as SUSY GUT or Super-string theories, treating gravity and quan-
tum mechanics in a consistent way, not only predict a de-
pendence of fundamental physical constants on energy (which has been observed in high energy experiments) but also al-
low their cosmological time and space variations (Uzan 2003). Detecting or constraining the possible time variations of funda-
mental physical quantities is an important step toward a complete understanding of basic physics. In the framework of
standard Big-bang models one can probe the evolution of various physical quantities over the elapsed time using measure-
ments that are performed at different redshifts (z). As the
energy of atomic transitions depends on the electromagnetic coupling constant α (its well measured laboratory
value is e2/ℏc = 1/137.0359976(50), see Mohr & Taylor 2000), its possible time variation will be registered in
the form of small shifts in the absorption line spectra seen toward high-z QSOs (Bahcall et al. 1967). Initial
attempts to measure the variation in α were based on alkali-
doublets (Wolfe et al. 1976; Levshakov 1994; Potekhin & Varshalovich 1994; Cowie & Songaila 1995; Murphy
et al. 2001). This method (AD method) uses the difference in the wavelengths of the doublets originating from the same ground state (i.e., 2S1/2 →2P3/2 and 2S1/2 →2P1/2 transitions). The constraints on the variation in α, Δα/α, are obtained by assuming that the measured difference in the wave-
length centroid of the doublets is proportional to α2 to the lowest order. The best constraint obtained using this method is
Δα/α = (−0.5 ± 1.3) × 10−5 (Murphy et al. 2001). Like absorption doublets one can also use the central wavelength of
multiple emission lines originating from the same initial excited state. For example, Bahcall et al. (2004) use the nebular

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Observatory (ESO), under the Large Programme “QSO Absorption
Line Systems” ID No. 166.A-0106 with UVES on the 8.2 m KUEYEN
 telescope operated at the Paranal Observatory, Chile.
O III emission lines at λ5007 and λ4959 originating from 1D2 excited level and derive \( \Delta \alpha/\alpha = (0.7 \pm 1.4) \times 10^{-5} \) based on 73 quasar spectra over the redshift range 0.16 \( \leq z \leq 0.80 \). Studies based on molecular absorption lines seen in the radio/mm wavelength range are more sensitive than those based on optical/UV absorption lines. At present good constraints are available for two systems, \( \Delta \alpha/\alpha = (-0.10 \pm 0.22) \times 10^{-5} \) at \( z = 0.2467 \) and \( \Delta \alpha/\alpha = (-0.08 \pm 0.27) \times 10^{-5} \) at \( z = 0.6847 \) if one assumes constant proton \( g \)-factor (Murphy et al. 2001a). It is also possible to have joint constraints on the fundamental constants using molecular lines (see Chengalur & Kanekar 2003). However, such studies at high \( z \) are not available due to the lack of molecular absorption systems at high \( z \). The most suitable and accurate method for measuring \( \Delta \alpha/\alpha \) at high redshift is called the many-multiplet method (MM method). This is a generalization of the AD method and was introduced by Dzuba et al. (1999). Unlike the AD method, this method uses absolute wavelength measurements of numerous absorption lines from different species. This method has been shown to provide an order of magnitude improvement in the measurement of \( \Delta \alpha/\alpha \) compared to AD method (Murphy et al. 2003, and references therein). However, this method is also extremely vulnerable to a number of systematical errors. Accurate laboratory wavelengths (up to an accuracy of few mÅ) are available for most of the important transitions that are regularly detected in QSO spectra (see Table 1). The sensitivity coefficients of the different line transitions from different multiplets to the variation in \( \alpha \) were computed using many-body calculations taking into account the dominant relativistic effects (Dzuba et al. 1999, 2002). If one uses the parameterization of these authors then the accuracy in probing the possible variations in \( \alpha \) depends very much on how well one measures the wavelength of the absorption lines at high \( z \). In simple terms, the MM method exploits the fact that the energy of different line transitions varies differently for a given change in \( \alpha \). For example, rest wavelengths of Mg II doublets and Mg I are fairly insensitive to small changes in \( \alpha \), thereby providing good anchors for measuring the systemic redshift. On the other hand, the rest wavelengths of Fe II multiplets are very sensitive to small variations in \( \alpha \). Thus constraining relative shifts between an anchor and different Fe II lines make it possible to study the variations in \( \alpha \). However the accuracy depends on how well the absorption line profiles are modeled. The absorption profiles are usually modeled using multiple Voigt profiles that are defined by column density (\( N \)), velocity dispersion (\( b \)) and redshift in addition to the rest-wavelength of the species. In general the Voigt profile decomposition is robust. However, the profile decomposition is not unique when the absorption lines are heavily blended and/or if the signal-to-noise ratio of the spectrum is not high enough. In addition, in real data, small relative shifts between absorption lines from different species can be introduced due to various systematic effects such as chemical and ionization inhomogeneities in the absorbing region, incorrect wavelength calibration, isotopic abundances, and atmospheric dispersion effects etc. It is usually argued that these random systematic effects can be canceled by using a large number of measurements. Indeed, applying the method to 128 systems over a large redshift range 0.2 \( < z < 3.7 \), Murphy et al. (2003) measured \( \Delta \alpha/\alpha = (-0.54 \pm 0.12) \times 10^{-5} \). The detailed work based on the MM method has been performed by only one group using simultaneous fitting of absorption line profiles of species covering a wide range of ionization states. Their important results need to be confirmed independently using a data set that is optimized for the purpose. This forms the main motivation of this work. One of the main criticism made of the MM method is the apparent lack of transparency in the complex analysis (Bahcall et al. 2004). Here, we perform a similar analysis to the one used by Webb and collaborators on a homogeneous and very high quality data set. To make the whole process accessible for critical scrutiny, we give a detail description of the procedure and the individual fits. The main idea is that it is better to derive strong constraints from small, though statistically significant, sample of well chosen systems rather than loose constraints from a larger sample of strongly blended systems. The details of the observations and the quality of the data used in the analysis are discussed in Sect. 2. In Sect. 3 we discuss simulations that are performed to validate our Voigt profile fitting code in extracting the variations in \( \alpha \). We also use the simulation results to construct an optimum selection criterion that will ensure the best possible detection limit in \( \Delta \alpha/\alpha \). In Sect. 4 we list all the Mg II, Fe II systems that are selected in our sample. Comments on individual Voigt profile fitting are given in Sect. 5. The discussion and conclusions are presented in Sect. 6.

2. Observations and data sample

The data used in this study have been obtained using the Ultra-violet and Visible Echelle Spectrograph (UVES) mounted on the ESO KUEYEN 8.2 m telescope at the Paranal observatory for the ESO-VLT Large Programme “QSO absorption lines”. This corresponds to a homogeneous sample of 18 QSO lines of sight suitable for studying various properties of the intergalactic medium over a redshift range 1.7–3.2. All the objects were observed in good seeing conditions with 1 arcsec slit width. The data were reduced using the UVES pipeline, a set of procedures implemented in a dedicated context of MIDAS, the ESO data reduction package. The main function of the pipeline is to perform a precise inter-order background subtraction for science frames and master flat-fields, and to allow an optimal extraction of the object signal, rejecting cosmic ray impacts and performing sky-subtraction at the same time. Usually a given wavelength range is covered by more than 6 exposures, and removal of cosmic rays and odd flux fluctuations in a given exposure is robust. The reduction is checked step by step. Wavelengths are corrected to vacuum-heliocentric values and individual 1D spectra are combined. Air-vacuum conversions and heliocentric corrections were done using standard conversion equations (Elden 1966; Stumpff 1980). The slit was always oriented along the parallactic angle, and calibration exposures were taken before or after the scientific exposures. Thus most of the important concerns raised by Murphy et al. (2003), are taken care of in our observations. As we are using the \( \chi^2 \) minimization method to detect sub-pixel-scale shifts, the error spectrum is crucial for our analysis. Great care was taken in computing the error spectrum while combining the individual exposures. Our final error is the quadratic sum of appropriately
Table 1. Summary of atomic parameters that are used in our analysis.

<table>
<thead>
<tr>
<th>Species</th>
<th>( \lambda_0 ) (Å)</th>
<th>Ref. (( \lambda_0 ))</th>
<th>( \omega_0 ) (cm(^{-1} ))</th>
<th>( q )</th>
<th>id</th>
<th>( f )</th>
<th>Ref. (( f ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg I</td>
<td>2852.96310(8)</td>
<td>L</td>
<td>35051.277(1)</td>
<td>+86(10)</td>
<td>i</td>
<td>1.830</td>
<td>A</td>
</tr>
<tr>
<td>Mg II</td>
<td>2796.3543(2)</td>
<td>L</td>
<td>35760.848(2)</td>
<td>+211(10)</td>
<td>a</td>
<td>0.6123</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>2803.5315(2)</td>
<td>L</td>
<td>35669.298(2)</td>
<td>+120(10)</td>
<td>b</td>
<td>0.3054</td>
<td>A</td>
</tr>
<tr>
<td>Al II</td>
<td>1670.7887(1)</td>
<td>M</td>
<td>59851.972(4)</td>
<td>+270(30)</td>
<td>j</td>
<td>1.88</td>
<td>B</td>
</tr>
<tr>
<td>Si II</td>
<td>1526.70709(2)</td>
<td>M</td>
<td>65500.4492(7)</td>
<td>+68(30)</td>
<td>k</td>
<td>0.133</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>1808.01301(1)</td>
<td>M</td>
<td>55309.3365(4)</td>
<td>+531(30)</td>
<td>l</td>
<td>0.00208</td>
<td>A</td>
</tr>
<tr>
<td>Fe II</td>
<td>1608.45085(8)</td>
<td>P</td>
<td>62171.625(3)</td>
<td>−1200(300)</td>
<td>c</td>
<td>0.0577</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>2344.2130(1)</td>
<td>Q</td>
<td>42658.2404(2)</td>
<td>+1254(150)</td>
<td>d</td>
<td>0.114</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>2374.4603(1)</td>
<td>Q</td>
<td>42114.8329(2)</td>
<td>+1640(150)</td>
<td>e</td>
<td>0.0313</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>2382.7642(1)</td>
<td>Q</td>
<td>41968.0642(2)</td>
<td>+1498(150)</td>
<td>f</td>
<td>0.320</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>2586.6496(1)</td>
<td>Q</td>
<td>38660.0494(2)</td>
<td>+1520(150)</td>
<td>g</td>
<td>0.0691</td>
<td>D</td>
</tr>
<tr>
<td></td>
<td>2600.1725(1)</td>
<td>Q</td>
<td>38458.9871(2)</td>
<td>+1356(150)</td>
<td>h</td>
<td>0.239</td>
<td>D</td>
</tr>
</tbody>
</table>

Reference:
- oscillator strength:– (A) Morton (1991); (B) Prochaska et al. (2001); (C) Morton (2003); (D) Bergeson et al. (1996);
- wavelength:– (L) Pickering et al. (1998); (M) Griesmann & Kling (2002); (P) Pickering et al. (2002); (Q) Nave et al. (1991).

interpolated weighted mean errors and possible errors due to scatter in the individual flux measurements. In all cases, the spectra cover the wavelength range of 3000–10 000 Å. A first-guess continuum is fitted with an automatic continuum fitting code. However, for the systems that are eventually used in the analysis, a local continuum normalization is performed using lower order polynomials. In all cases we have used the flux in the core of the saturated lines to estimate the background subtraction uncertainties. This corrections to the background subtraction uncertainties are applied whenever needed. We notice that continuum fitting is not unique in the Lyα forest region and in the IR region where atmospheric lines are plenty. In our analysis we have therefore avoided the absorption lines that lie on the lower wavelength side of Lyα emission from the quasar and lines that could be blended with atmospheric lines.

The properties of the spectra in our sample are summarized in Fig. 1. A typical \( S/N \sim 60–80 \) per pixel (typically 0.035 Å wide) is achieved in the whole wavelength range of interest (see left-hand top panel in Fig. 1). This is a factor 2 improvement compared to the one used earlier for the analysis of \( \Delta \alpha/\alpha \) (see Fig. 2 of Murphy et al. 2003). The spectral resolution measured from the lamp spectrum is typically \( \pm 44,000 \).

We have investigated the accuracy of the wavelength calibration using lamp spectra that are extracted in the same way as the object. We notice that individual emission lines are well modeled by a Gaussian. This confirms that the instrumental profile can be modeled with a single Gaussian while fitting the Voigt profiles. The median difference between the extracted and the actual wavelengths at most of the settings are close to zero. The root mean square of this deviation is always smaller than 3 mÅ at all the settings. We confirm the robustness of the wavelength calibration using other consistency checks. In particular, the wavelengths of Fe II transitions strongly depend on \( \alpha \) but the relative position of the lines (apart from Fe II\( \lambda 1608 \)) does not depend much on \( \alpha \). This means that, whatever the variation of \( \alpha \) is, the relative velocity shifts between Fe II transitions should be consistent with zero in the case of good calibration. This is precisely what is observed (Panel b in Fig. 1). Note in addition that the Fe II rest wavelengths are known to very high accuracy (\( \sim 0.1 \) mÅ). In principle a similar exercise can be performed using Si IV and C IV absorption lines. However, unlike Fe II lines, C IV wavelengths are known only to 2 mÅ accuracy. Even in this case the velocity shifts measured are consistent with zero within the allowed laboratory wavelength uncertainties (Petitjean & Aracil 2003). When fitting the narrow doublets and multiplets from a given species in individual settings, we have allowed for some adjustments in the instrumental resolution. This seems to be an important step as we realized that the seeing was much better than the slit width during most of the observing run. The metal line systems in the reduced spectra are identified using the standard procedure.

The next step is to validate our Voigt profile fitting codes and make some estimate of various possible systematic effects. To do this we first simulate spectra with specifications similar to a typical UVES spectrum and perform the analysis on them. The details of the spectral simulations and the data analysis process is described in the following section.

3. Chasing the systematics using simulated data

As one tries to measure \( \Delta \alpha/\alpha \) through Voigt profile decomposition of absorption lines, various systematic effects can cause false-alarm detection of \( \Delta \alpha/\alpha \). The effects of random systematics can be averaged out by combining large numbers of measurements. However, the best strategy will be to have an a priori estimate of various systematic effects so that the proper selection criteria can be applied while choosing the sample to achieve the best possible results. In this section we try to understand various effects using simulated spectra. Such an exercise is also important to see how well our Voigt profile fitting code
Fig. 1. Properties of our sample: Panel a) shows the distribution of median signal-to-noise (S/N) ratio per pixel computed over 30 Å intervals. A typical S/N = 60–80 per pixel is achieved over the wavelength range 4000–8000 Å. Panel b) shows the distribution of measured relative velocity shifts between the Fe II λ2344 and Fe II λ2600 lines in our sample. As the q coefficients are similar for these transitions the velocity shift is expected to be distributed around zero in the case of good wavelength calibrations. Thus the plot provides an internal consistency check for the wavelength calibration. We notice that the accuracy in the wavelength calibration is of the order of 1–3 mÅ in the wavelength range of interest. Panel c) shows the distribution of the spectral resolution measured from the calibration spectrum. The typical spectral resolution achieved is larger than 44 000. Continuous and dotted distributions in these panels are for the blue (4000–5500 Å) and red (5500–8500 Å) spectral range respectively. Panel d) gives the histogram of the number of sight lines at a given redshift in which Mg II and Fe II absorption lines are in the observed wavelength range. The 18 sight lines in our sample cover the redshift range between 1 and 2 uniformly.

reverses the input value of $\Delta \alpha/\alpha$ that is used while generating the spectrum.

3.1. Generation of absorption lines

In this section, we simulate data for different assumed values of $\Delta \alpha/\alpha$ and apply our procedure to recover the input value of $\Delta \alpha/\alpha$. We mainly concentrate on absorption lines from the Mg II doublet and Fe II multiplets. We assume the ratio, $N$(Mg II)/$N$(Fe II), to be constant in individual components. In order to mimic real data, the Voigt profile of the absorption line is convolved with an instrumental profile and appropriate noise is added to the spectrum. We model the instrumental profile as a single Gaussian with $\sigma = 1.69$ pixels (with pixel size of 0.03 Å) appropriate for $\lambda/\Delta \lambda = 42,000$ as achieved in a typical UVES observation (note however that the actual resolution achieved is somewhat higher in the real data).

The noise in the spectrum is the combination of photon noise, readout noise, and the residuals from background subtraction. While the former dominates the error in the continuum the latter two will be the source of error in the core of saturated lines. We notice from our UVES data that the root mean square of the residual at the bottom of highly saturated lines is $\approx 0.4\%$ of the unabsorbed continuum flux. The readout noise added to a typical normalized spectrum mimics a Gaussian distribution with zero mean and a standard deviation of $4 \times 10^{-3}$. The photon noise spectrum is obtained using Poisson statistics. To obtain a spectrum with a given signal-to-noise ratio, ($S/N$), we have generated Poisson random numbers ($P[(S/N)^2]$), with a mean equal to ($S/N$)$^2$, and then assigned photon noise to the continuum of the normalized spectrum to be ($P[(S/N)^2] - (S/N)^2)/(S/N)^2$, having a Poisson distribution with $\sigma = (S/N)^{-1}$. In the absorbed portion of the spectrum this value is scaled by the square root of the intensity. In all our simulations we assume $S/N$ in the
Fig. 2. Illustration of the methodology and reliability of our Voigt profile-fitting code: the panels on the left-hand side show the velocity plot of the simulated Mg II and Fe II absorption lines (points with error bars) and the best-fit Voigt profiles (solid curve). The panel at the top right-hand side shows the $\chi^2$ as a function of $\Delta\alpha/\alpha$. The best-fit value and 1σ error are shown in the panel. This is consistent with the input value of $\Delta\alpha/\alpha = 0$. The lower right-hand panel shows the relationship between the input and recovered values of $\Delta\alpha/\alpha$. Each point is a result obtained from a single realization (there are 30 of them for a given input $\Delta\alpha/\alpha$). Typical errors in these measurements are shown in the bottom right corner of the panel. The dark circles with error bars represent the weighted mean value of $\Delta\alpha/\alpha$ obtained from 30 realizations.

To simulate a spectrum incorporating a varying fine-structure constant ($\alpha_z$), we use the analytic fitting function given by Dzuba et al. (2002),

$$\omega = \omega_0 + qx,$$

(1)

Here $\omega_0$ and $\omega$ are, respectively, the vacuum wave number (in unit of cm$^{-1}$) measured in the laboratory and in the absorption system at redshift $z$. $x$ is a dimensionless number defined by $x = (\alpha_z/\alpha_0)^2 - 1$ where $\alpha_0$ refers to the present value of the fine-structure constant and $\alpha_z$ to its value at redshift $z$. The numerical values of the parameter $q$ that are obtained using many-body relativistic calculations (see Dzuba et al. 1999, 2002) for different species are listed in the fifth column of Table 1. Originally $\omega$ is defined as $\omega = \omega_0 + q_1 x + q_2 y$ with $y = (\alpha_z/\alpha_0)^4 - 1$ and $x$ as defined above (Dzuba et al. 1999). When $\Delta\alpha/\alpha$ is small $q$ is basically $q_1 + 2q_2$. The table also gives the laboratory values of wavelengths (with standard isotopic composition for different elements) ($\lambda_0$), wavenumbers ($\omega_0$) and oscillator strengths ($f$) used in this study. References to various parameters used are also provided in this table.

3.2. Voigt profile fitting code

To measure the relative shifts between an anchor and Fe II lines, we fit all the lines simultaneously. In addition, we have modified our Voigt profile fitting code (Khare et al. 1997) to take into account the dependence of rest wavelengths of different species on $\alpha$. We improved the convolution algorithm using a 41-point Gaussian quadrature integration scheme. The latter is crucial for fitting narrow lines in high signal-to-noise data. The FWHM of the Gaussian kernel is considered to be $\lambda/\Delta\omega/\alpha$. Here $\lambda$ is the median wavelength over the region of the absorption line being fitted and $\Delta\omega/\alpha$ is the spectral resolution. We have taken a ±3σ ($\sigma = FWHM/2.3548$) wavelength range around each pixel of absorption line for convolution. The Gaussian quadrature integration method is adopted because it provides the most accurate estimate of the area under the Gaussian kernel that is used for normalizing the truncated Gaussian probability density function.
function. Our code also takes into account the variation of spectral resolution across the spectrum.

3.3. Measurement of $\Delta \alpha/\alpha$ in a single component systems

In order to find the best $\Delta \alpha/\alpha$ consistent with our data, we fit the absorption lines using the modified rest wavelengths (see Eq. (1)), varying $\Delta \alpha/\alpha$ in the range from $-5.0 \times 10^{-3}$ to $5.0 \times 10^{-3}$ in steps of $0.1 \times 10^{-3}$. For each value we obtain $\chi^2_{\text{min}}$ by varying $N$, $b$ and $z$. The value of $\Delta \alpha/\alpha$ at which $\chi^2_{\text{min}}$ is minimum is accepted as the best possible $\Delta \alpha/\alpha$ value, provided the reduced $\chi^2$ of fit is $\approx 1$. Following the standard statistical procedure (Press et al. 2000, see pages 690–691) we assign $1 \sigma$ errors to the above defined best value of $\Delta \alpha/\alpha$ by computing the required change in $\Delta \alpha/\alpha$ so that $\Delta \chi^2 = \chi^2 - \chi^2_{\text{min}} = 1$. To be on the conservative side we have taken the maximum change in $\Delta \alpha/\alpha$ around the best value of $\Delta \alpha/\alpha$ that is required to produce $\Delta \chi^2 = 1$ as our error on $\Delta \alpha/\alpha$.

We test the method discussed above using a simulated single component system (see Fig. 2). The profiles of $\text{Mg II}$ and $\text{Fe II}$ absorption lines that are generated for $\Delta \alpha/\alpha = 0$ are shown on the left-hand panels of the figure. In the top right-hand panel we show the best fitted $\chi^2$ as a function of $\Delta \alpha/\alpha$. The location of the minimum and $1 \sigma$ error in $\Delta \alpha/\alpha$ are shown using a point with an error bar. To investigate the systematics in detail, we have simulated single component systems with column densities in the range $10^{13}$ to $10^{16}$ cm$^{-2}$ (we consider that the number of systems with a given $N(\text{Mg II})$ is proportional to $N(\text{Mg II})^{-1.4}$, Srianand & Khare 1994), the velocity dispersion uniformly distributed in the range 3 to 10 km s$^{-1}$ and the input $\Delta \alpha/\alpha$ varying between $-10^{-5}$ and $+10^{-5}$. In addition we have varied the seed for the random number generator that produces the error spectrum. We use our Voigt profile fitting code to recover $\Delta \alpha/\alpha$ in each case. The recovered value of $\Delta \alpha/\alpha$ in each realization is plotted (dots) as a function of the input $\Delta \alpha/\alpha$ in the lower right-hand panel in Fig. 2. The typical error in an individual measurement is also shown in the plot. There are about 30 realizations for a given value of input $\Delta \alpha/\alpha$ with a varied $N$, $b$ and noise spectrum. The dark circles with error bars represent the weighted mean value of the output $\Delta \alpha/\alpha$ using all the 30 measurements. The weighted mean recovered values of $\Delta \alpha/\alpha$ are consistent with the input values.

It is clear from Fig. 2 that our procedure works well for simple single-component systems. We notice that the distribution of differences between $\Delta \alpha/\alpha$ output and input values is well fitted by a Gaussian (see Fig. 5) with $\sigma = 0.23 \times 10^{-3}$. Based on the central limit theorem and a Gaussian distribution of the recovered values around the mean we expect the accuracy of the estimated mean value to be $\sim \sigma/\sqrt{N}$ for $N$ measurements. Strong blending and internal structure in the cloud will imply larger uncertainty.

The improvement we expect from the enhanced $S/N$ ratio in our sample compared to other studies is visualized in Fig. 3. Here, we show the results obtained for individual realizations of single-component systems with input $\Delta \alpha/\alpha = 0$ for $S/N = 30$ and 70 in the top and bottom panels respectively. The shaded region in the figure shows the mean and $\sigma$ in the distribution of the recovered values. The distribution of individual values has $\sigma$ of $0.409 \times 10^{-5}$ and $0.239 \times 10^{-5}$ for signal-to-noise ratios 30 and 70 respectively.

It is clear that the improvement in the $S/N$ ratio gives a factor 2 improved accuracy in the recovered value compared to what has been done previously. In all the simulations discussed till now we have used only strong but unsaturated $\text{Fe II}$ lines (i.e. $N(\text{Fe II}) \gtrsim 10^{13}$ cm$^{-2}$). This ensures that the absorption profiles are well defined. However, when the lines are weak the flux in the line is appreciably affected by Poisson noise. One can therefore infer that the scatter in the measurements will be larger if weak lines are used in the analysis. We demonstrate this using simulations of weak lines. For simplicity we have assumed input $\Delta \alpha/\alpha = 0$, $N(\text{Mg II}) = 1.5 \times 10^{12}$ cm$^{-2}$ and $N(\text{Fe II}) = 5 \times 10^{11}$ cm$^{-2}$. We have simulated 30 systems by just changing the error spectrum. The results are summarized in Fig. 4. In the left-hand panel we show the profiles for one realization. From this figure one can appreciate the distortion in the absorption profile of $\text{Fe II}$ lines caused by Poisson fluctuations. The top panel on the right-hand side shows the recovered values of $\Delta \alpha/\alpha$. As expected, the error bars are larger (one $\sigma$ of the distribution around the mean is $2.92 \times 10^{-5}$). Typical errors are then a factor 10 higher than what we derive with strong single-component systems. Thus if one is going to use $1/\sigma^2$ weighting of different measurements, only one single strong and narrow component can be as important as about
In what follows we use these results as guidelines for choosing the systems in our sample for the final analysis.

### 3.5. $\Delta \alpha/\alpha$ measurements from well separated blends

The next step is to investigate the case of multicomponent systems where subcomponents are marginally blended. That is the cases where the separation between individual components is larger than the individual $b$ values. Results are summarized in Fig. 6. The left-hand side panel shows velocity profiles for one of the realizations. In the right-hand-side panels we compare results obtained for the case considered here (middle panel) with that obtained for single component systems (lower panel) and strongly blended systems (top panel). The $\sigma$ of the distribution around the mean are $0.221 \times 10^{-5}$, $0.261 \times 10^{-5}$ and $0.519 \times 10^{-5}$ for the single, weakly blended and strongly blended systems respectively. It is apparent that the mildly blended case gives very similar results as that obtained with single components.

### 3.6. Conclusions from simulations

In short, we can conclude from these simulations that:

1. Best constraints on $\Delta \alpha/\alpha$ are obtained either from single component systems or well resolved multiple component systems.
2. Increasing the signal-to-noise ratio from $S/N = 30$ to 70 increases the accuracy of $\Delta \alpha/\alpha$ measurements by about a factor of two.
3. It is better to avoid weak lines while extracting $\Delta \alpha/\alpha$, as their profiles can be distorted by Poisson noise. Thus, weak lines in the low-signal-to-noise data can result in false-alarm detections of non-zero $\Delta \alpha/\alpha$ values.
4. There is a non-negligible probability for deriving a statistically significant deviation from the actual value when one considers highly blended systems (i.e., systems where the component separations are smaller than the individual $b$ values). Thus it is better to avoid complex blends in the analysis.

In what follows we use these results as guidelines for choosing the systems in our sample for the final analysis.
Fig. 5. Effect of blending in the determination of $\Delta \alpha/\alpha$: absorption spectra of Mg II and Fe II were simulated for given $N$, $b$, and spectral resolution similar to that of our data, introducing spectral shifts corresponding to a given value of $\Delta \alpha/\alpha$. The top panels show the relationship between the input and derived value of $\Delta \alpha/\alpha$ in the case of a single clean component (left-hand side) and a blend of two components (right-hand side). A typical absorption profile is also shown in these panels. Dots are the values from individual realizations; the points with the error bars represent the weighted mean over 30 realizations. The lower panels give the distribution of the recovered $\Delta \alpha/\alpha$ around the true one. Single (left) and blended (right) cases are considered respectively. Fits by Gaussian distributions are over-plotted.

4. Description of the UVES sample

The Mg II systems with detectable Fe II absorption lines present in our data are listed in Table 2. The table provides the QSO name, the emission redshift ($z_{\text{em}}$), the redshift of the absorption line systems ($z_{\text{abs}}$) and the species that are detected (notations are as in Table 1). No such Mg II systems are detected along 2 sight lines (toward HE 1158–1843 at $z_{\text{em}} = 2.449$ and Q 0420–388 at $z_{\text{em}} = 3.177$). The last column of the table is meant for comments on the systems.

In total, there are 50 systems with detected Mg II and Fe II lines at a detection limit of $\sim 1.5 \times 10^{11}$ cm$^{-2}$ for the strongest Fe II $\lambda2374$ line. The corresponding limit for Fe II $\lambda2374$ is $1.5 \times 10^{12}$ cm$^{-2}$. In the following we discard systems with $N(\text{Fe II}) < 2 \times 10^{12}$ cm$^{-2}$, as simulations have shown that spurious effects can affect the results when the lines are weak. Therefore systems with $N(\text{Fe II})$ less than this limit (15 systems) are marked as “weak” in Table 2 and are not considered in our main analysis. However, for completeness, we present the results obtained from these systems also in the concluding section.

Two systems at $z_{\text{abs}} = 1.0598$ and $z_{\text{abs}} = 1.9183$ toward HE 0940–1050 could not be considered in our analysis because there are no good anchor lines in these systems. In the former system only Mg II $\lambda2796$ is present in the spectrum. In the case of the $z_{\text{abs}} = 1.9183$ system, Si II and Al II are blended with Ly$\alpha$ lines and Mg II is heavily affected by atmospheric lines. In addition, at $z_{\text{abs}} = 0.7261$ toward Q 0453–423, $z_{\text{abs}} = 0.8367$ toward Q 0002–422 and $z_{\text{abs}} = 0.6631$ toward PKS 2126–158, the Fe II lines are heavily blended with Ly$\alpha$ systems and are therefore marked as “Ly$\alpha$”. Thus we are left with 29 systems that have strong Fe II lines, good anchor lines and are not contaminated by Ly$\alpha$ or atmospheric absorption. These systems form the basic dataset that we use for measurement of $\Delta \alpha/\alpha$.

In summary, based on the results from the simulations, we apply the following selection criteria to derive reliable $\Delta \alpha/\alpha$:

1. We consider only lines with similar ionization potentials (Mg II, Fe II, Si II and Al II) as they are most likely to originate from similar regions in the cloud.
2. We avoid absorption lines that are contaminated by atmospheric lines.
Table 2. List of Mg II and Fe II systems.

<table>
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<tr>
<th>QSO</th>
<th>$z_{em}$</th>
<th>$z_{abs}$</th>
<th>Transitions</th>
<th>Comments</th>
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</table>

Transitions as identified in Table 1.

“weak” refers to systems with $N(Fe II) < 2 \times 10^{12}$ cm$^{-2}$.

“+” indicates systems that are not used in the analysis (see discussion).

“Lyα” refers to systems which are blended in the Lyα forest.

“no anchor” refers to systems without good anchor lines.

Fig. 6. Results for well-separated blends: left-hand-side panels show the velocity plot and the best-fit Voigt profiles to Mg II and Fe II absorption lines from one of the realizations. In all cases considered here the velocity separation between the components is always more than the individual values of the $b$ parameters. In the right-hand-side panels we compare the results obtained for heavily blended systems (top panel), well separated blends (middle panel) and single-component systems (lower panel). The standard deviation measured around the mean in different cases is also given in the corresponding panels. It is clear that systems with well separated sub-components have similar $\sigma$ as that of single-component systems for $\Delta \alpha$ measurement.

3. We consider only systems that have $N(Fe II) \geq 2 \times 10^{12}$ cm$^{-2}$ which ensures that all the standard Fe II multiplets are detected at more than 5$\sigma$ level.

4. We demand that at least one of the anchor lines is not saturated so that the redshift measurement is robust.

5. We also avoided sub-damped Lyα (i.e. $N(H I) \geq 10^{19}$ cm$^{-2}$). Here after we denote them as sub-DLAs as these systems may have ionization and chemical inhomogeneities.

6. We do not consider strongly saturated systems with large velocity spread (complex blends); however in such systems whenever we find a well detached satellite components we include these components in the analysis.

7. Finally, based on the component structure resulting from the Voigt profile fits of systems that are not complex blends, we retain only systems for which the majority of components are separated from its neighboring components by more than the $b$ parameters.

Application of the above conditions resulted in 23 systems on which to perform the measurement (six single component systems, six well separated two components (doubles), six systems with three components having at least one well detached from the rest, and five systems with more than 3 components).
5. Notes on individual systems in our sample

5.1. Systems along the line of sight toward HE 1341–1020

There are 3 Mg II systems with Fe II along this sight line. All these three systems are strong enough to be included in our sample.

5.1.1. \( z_{\text{abs}} = 0.8728 \) system toward HE 1341–1020

The Voigt profile fits to some of the lines from this system are shown in Fig. 7. This system shows Al II, Mg I \( \lambda 2852 \), Ca II \( \lambda \lambda 3934, 3969 \) lines in addition to Mg II and Fe II lines. The Mg II profile is fitted with 8 components, the three strongest of which show absorption in other species. The Fe II \( \lambda 2600 \) line is blended with absorption lines from other intervening systems and the Fe II \( \lambda 2374 \) profile is affected by a bad pixel. In order to avoid unsystematic errors we do not consider these lines for the \( \Delta \alpha / \alpha \) measurement analysis. However we use these profiles to check the consistency in the number of components and the level of saturation in each component. The Al II line is affected by noise and the Ca II lines are too weak to give any meaningful constraints. We therefore use Fe II \( \lambda \lambda \lambda 2344, 2382, 2586, \) Mg II \( \lambda \lambda 2796, 2803 \) and Mg I \( \lambda 2852 \).

5.1.2. \( z_{\text{abs}} = 1.2778 \) system toward HE 1341–1020

The absorption profile of this system is spread over 200 km s\(^{-1}\). Absorption lines of Mg I, Mg II, Al II, Al III, Fe II, Ca II, Si II, Mn II, Ni II, and Ca II are detected. The core is strongly blended over 100 km s\(^{-1}\) but a well detached narrow satellite is seen ~100 km s\(^{-1}\) away. We have used Fe II and Mg II absorptions from this sharp component for the measurement of \( \Delta \alpha / \alpha \). We do not use Mn II, Ni II and Ca II because they are weak. Si II and Al II lines fall in the Ly\( \alpha \) forest and the profile of Fe II \( \lambda 2586 \) is affected by the wing of a contaminating neighboring line. The satellite component is well fitted with two components (see middle panel in Fig. 7). Even though the two components are merged, we notice that the components are separated by more than the velocity dispersion of the individual component.

5.1.3. \( z_{\text{abs}} = 1.9154 \) system toward HE 1341–1020

This system shows absorption lines from Mg II, Mg I, Al II, Al III, C IV, Si II, and Si IV. The high-ionization lines are weak and their profiles are very different from that of the low-ionization lines. The Mg II lines are saturated and blended with atmospheric absorption lines. Thus, we use Si II \( \lambda \lambda 1526, \) Si II \( \lambda 1808 \) and Fe II lines for \( \Delta \alpha / \alpha \) measurement. We do not use Fe II \( \lambda 2586 \) however as its profile is affected by noise. The system is fitted with 4 distinct components that are clearly visible in the profiles of the weak lines (see right panel in Fig. 7). Even though the overall fit is good, one can notice that the Si II \( \lambda 1808 \) line profile is apparently not reproduced well. This is due to the effect of noise in a single pixel in the red wing of the strongest absorption component. This shows how delicate the measurement can be.

5.2. Systems along the line of sight toward Q 0122–380

There are 4 Mg II systems along this line of sight. The system at \( z_{\text{abs}} = 1.9102 \) has weak Fe II absorption lines and does not fulfill our criteria to enter our main sample.

5.2.1. \( z_{\text{abs}} = 0.8221 \) system toward Q 0122–380

This system is defined by narrow Mg II, Mg I and Fe II absorption lines. All the lines are well fitted by a single component. An extra component in the red wing is required for the Mg II doublets. The Voigt profile fits to the system are shown in Fig. 8.

5.2.2. \( z_{\text{abs}} = 0.8593 \) system toward Q 0122–380

This system is defined by Mg II, Mg I, Al II, Al III, Ca II, and Fe II absorption lines. As the spectral range in which Al II absorption is redshifted is of lower signal-to-noise and Ca II doublets and Fe II \( \lambda 2374 \) are weak we have not used these lines for the \( \Delta \alpha / \alpha \) analysis. The resulting Voigt profile fits are shown in Fig. 8. We notice that the Fe II \( \lambda 2586 \) line is slightly under-predicted in our best-fit model. However the other three Fe II lines are very well fitted. We also notice that the value of \( \chi^2 \) per degree of freedom is very good when using different \( b \) parameters for Fe II and Mg II. The result for \( \Delta \alpha / \alpha \) presented for this system is therefore obtained with different \( b \) parameters for Mg II and Fe II lines.
5.2.3. $z_{\text{abs}} = 1.2433$ system toward Q 0122–380

The Mg II absorption profiles in this system are spread over $\sim 70$ km s$^{-1}$. The components are heavily blended. The strongest Mg II component shows Mg I, Al III, Si II, Si IV, C IV, Ca II, Ni II and Fe II. Fe II $\lambda 2586$ and Fe II $\lambda 2600$ lines are not covered by our spectrum. Three Fe II lines and the Mg II doublet can be fitted with 4 components. The Ni II $\lambda 1709$ line is blended and Ni II $\lambda 1741$ is weak and falls in the noisy region of the spectrum. We notice small profile inconsistencies between the Ca II doublets. We therefore avoid using Ni II and Ca II lines in the analysis. The results are summarized in Fig. 8.

5.3. Systems along the line of sight toward PKS 0237–23

This QSO is known for the presence of a supercluster of C IV absorption lines at $z_{\text{abs}} \sim 1.6$. Our spectrum reveals 5 Mg II systems one of which at $z_{\text{abs}} = 1.1846$ has a very weak Fe II line and is not considered for our main analysis.

5.3.1. $z_{\text{abs}} = 1.3650$ toward PKS 0237–23

This system shows absorption lines of Fe II, Al II, Al III, C I, C I', C II, C II', Si II, Si IV, Ca II, Mg I and Mg II lines. The Ly$\alpha$ line is just below our wavelength coverage. The absorption profiles of strong resonance lines spread over 230 km s$^{-1}$. The main component has a large number of sub-components heavily blended. There is a well detached satellite component that could be used for the $\Delta \alpha/\alpha$ measurements. The presence of C I suggests that this could possibly be a sub-DLA. In such systems inhomogeneities are generic. Thus it is better to avoid them in the $\Delta \alpha/\alpha$ measurements. This system happens to be both in our data and in those of Murphy et al. (2003). Thus we perform the $\Delta \alpha/\alpha$ measurement in the satellite components. We use Al II, all Fe II lines (apart from Fe II $\lambda 2383$), and Mg II lines for our analysis. Murphy et al. (2003) have found $\Delta \alpha/\alpha = -0.19 \pm 0.5$ for this system. Unlike us Murphy et al. have used the whole profile of Al II, Si II, Al III, and two Fe II lines. Our analysis gives a best-fit value of $0.0 \pm 0.1$. This is consistent with that measured by Murphy et al. and our final result. The increased accuracy in our measurement is contributed to by the enhanced S/N and our choice of the well detached and unblended satellite component for the analysis. For the reason mentioned above we have excluded this system from our analysis.

5.3.2. $z_{\text{abs}} = 1.6371$ toward PKS 0237–23

The absorption profiles of this system are spread over $\sim 280$ km s$^{-1}$. Absorption lines from Mg I, Mg II, Al II, Al III, Si II and Fe II have been detected. Even though C IV and Si IV absorption lines are detected the system is dominated by low-ionization species. There are two distinct velocity components at $z_{\text{abs}} = 1.63588$ and 1.63715 that we use for $\Delta \alpha/\alpha$ measurements. The Voigt profile fits to these systems are shown in Fig. 9.

5.3.3. $z_{\text{abs}} = 1.6574$ toward PKS 0237–23

This system shows absorption lines from Mg II, Mg I, Al II, Al III, Si II, C IV, Si IV in addition to Fe II. The low- and high-ionization lines show distinct profiles. C IV and Si IV absorption lines are spread over 400 km s$^{-1}$. We use the uncontaminated Fe II lines with Si II and Al II lines as anchors. The Mg II line is heavily blended and has a large number of extra components compared to Fe II, and Mg I is weak. The results of the Voigt profile fits are shown in Fig. 9.
5.3.4. $z_{\text{abs}} = 1.6724$ toward PKS 0237–23

This is one of the few sub-DLAs that are detected in the large programme sample. Mg I, Mg II, C I, C I*, C II, C II*, Si II, Si IV, C IV, Ni II, Zn II, Cr II, Mn II and Fe II lines are present. Most of the standard resonance lines are saturated. Zn II, Mn II, Cr II and Ni II absorption lines are weak and therefore will not give good constraints. The presence of the whole range of ionization states and of fine-structure absorption lines strongly suggests strong inhomogeneities in the gas. To keep our analysis unbiased we did not use this sub-DLA system in our analysis.

5.4. Systems along the line of sight toward HE 0001–2340

There are 5 Mg II systems detected along this sight line. Two of them at $z_{\text{abs}} = 0.9489$ and 1.6517 are weak systems and are not considered in our main sample.

5.4.1. $z_{\text{abs}} = 0.4524$ system toward HE 0001–2340

This system has two very narrow components separated by $\sim 70$ km s$^{-1}$. The low-redshift component shows Mg II, Mg I, Fe II and Ca II $\lambda 3934$ absorption lines. The high-redshift component has very weak Fe II lines. Thus we use only the low-redshift component for $\Delta \alpha/\alpha$ measurement. Voigt profile fits to the absorption lines from this component are shown in Fig. 10. Fe II $\lambda 2344$ and Fe II $\lambda 2586$ are blended with other absorption lines and were not used in the analysis.

5.4.2. $z_{\text{abs}} = 1.5855$ system toward HE 0001–2340

Like the $z_{\text{abs}} = 0.4524$ system this system has two well detached components separated by $\sim 110$ km s$^{-1}$. Mg II, Mg I, Fe II, Al II, Al III, C IV, Si II and Si IV absorption lines are seen in both components. However Fe II lines are strong only in the high-$z$ component. We notice that Mg II $\lambda 2803$ is contaminated by an atmospheric line. As Si II $\lambda 1526$ is also blended, we use Al II and Mg I absorption lines as anchors. Even though the overall fit to the data is good, our profiles consistently over-predict the blue wing of Fe II $\lambda 2374$. This is probably due to one single pixel in the wing being affected by noise. As the two components that are required to fit the profiles are very close to each other this system is not considered for the final analysis.

5.4.3. $z_{\text{abs}} = 2.1839$ system toward HE 0001–2340

Absorption profiles produced by this system are spread over 400 km s$^{-1}$. Mg I, Mg II, Al II, Al III, Si II, Si IV and C IV absorption lines are detected. The Fe II lines are strong in only two of these components (at $z_{\text{abs}} = 2.1854$ and $z_{\text{abs}} = 2.1872$). As Fe II $\lambda 2383$, Fe II $\lambda 2586$ and Fe II $\lambda 2600$ absorption lines fall in the wavelength range affected by atmospheric absorption, we consider only the clean lines in each case. The results of profile fitting are shown in Fig. 10.

5.5. Systems along the line of sight toward Q 0109–3518

There are two Mg II system seen along the line of sight toward this QSO. We used both of them for $\Delta \alpha/\alpha$ measurement.

5.5.1. $z_{\text{abs}} = 1.1827$ system toward Q 0109–3518

This system shows absorption lines from Mg II, Mg I and Fe II. The Fe II and Mg I lines are fitted with a single component but Mg II requires additional components in the red wing. The results of Voigt profile fitting are shown in Fig. 11.
5.5.2. \( z_{\text{abs}} = 1.3483 \) system toward Q 0109–3518

This is a strong Mg \( \text{II} \) system with heavily blended absorption profiles spread over 300 km s\(^{-1}\). Absorption lines from Mg I, Mg II, Fe II, Al II, Al III, Si II, Si IV and C IV are detected. There are two well detached satellite components. We used the \( z_{\text{abs}} = 1.3483 \) satellite for \( \Delta \alpha/\alpha \) measurement. As Mg II is blended, Si II, Al II and Mg I lines are used as anchors. For the other satellite the anchor lines are all blended. The results of Voigt profile fitting are presented in Fig. 11.

5.6. Systems along the line of sight toward
\( z_{\text{em}} = 2.414 \) HE 2217–2818

There are five Mg II systems along the line of sight toward this QSO. Three of these systems (at \( z_{\text{abs}} = 0.7862, 1.6277 \) and 1.5556) are weak and are not considered in the main sample. The other two systems were used for \( \Delta \alpha/\alpha \) measurement.

5.6.1. \( z_{\text{abs}} = 0.9425 \) system toward HE 2217–2818

This system shows absorption lines from Mg II and Fe II. There are two distinct subsystems that are separated by \( \sim 200 \) km s\(^{-1}\). Only the high-redshift subsystem shows Fe II absorption. The Fe II lines are fitted with four well separated components and Mg II requires an extra component in the red wing. We notice that the goodness of the fit in this system is decided by how well we model the Mg II profiles. The results of the profile fitting are summarized in Fig. 11.

5.6.2. \( z_{\text{abs}} = 1.5556 \) system toward HE 2217–2818

This system shows absorption due to Mg II, Al II, Si II, Si IV, C IV and Fe II. The Mg II absorption profiles are spread over \( \sim 100 \) km s\(^{-1}\). However only the central velocity component shows other singly ionized species (see Fig. 11). Surprisingly this component also shows absorption due to O I. High-ionization lines such as C IV, Si IV are weak and Al III is absent.

5.7. Systems along the line of sight toward
\( z_{\text{em}} = 2.435 \) QSO Q 0329–385

There are two absorption systems detected along this line of sight. The \( z_{\text{abs}} = 1.43799 \) system is weak and is not considered in our sample.

5.7.1. \( z_{\text{abs}} = 0.7627 \) system toward Q 0329–385

This system shows absorption lines due to Mg I, Mg II, and Fe II. The Mg II absorption profile is spread over 200 km s\(^{-1}\). However the other lines are detected only in the central component. The Voigt profile fits to this component are shown in Fig. 12. The profiles are fitted with two components, and as the velocity separation is smaller than the largest \( b \) value of the components we do not consider this system in our study.

5.7.2. \( z_{\text{abs}} = 1.8583 \) system along the line of sight toward Q 0453–423

This is a narrow single-component absorption system detected in Mg II, Al II, Al III, Si II, Si IV and Fe II. The results of the single-component Voigt profile fit to three prominent Fe II lines and the Mg II doublet are shown in Fig. 12.
5.8.3. \( z_{\text{abs}} = 2.3004 \) system along the line of sight toward Q 0453–423

This system shows absorption lines of Mg II, Fe II and Al II. Fe II \( \lambda 2586 \) and Fe II \( \lambda 2600 \) are not covered by our spectrum. Fe II \( \lambda 2374 \) is very weak. The signal-to-noise ratio in the spectral range where other Fe II lines are seen is \( \sim 30 \) only. In addition, both Mg II lines are contaminated by atmospheric absorption. There are also profile inconsistencies between the weak Al II lines (a potential anchor) and Fe II lines. Thus we do not consider this system in our study.

5.9. Absorption systems along the line of sight toward \( z_{\text{em}} = 2.767 \) Q 0002–422

There are 5 Mg II systems with detected Fe II absorption lines along this line of sight. The \( z_{\text{abs}} = 0.8367 \) system is a very strong Mg II system whose absorption profile is spread over \( \sim 600 \) km s\(^{-1}\). As the corresponding absorption profiles are heavily blended we do not consider this system in our analysis, as is the case for the \( z_{\text{abs}} = 1.9885 \) system in which Fe II lines are too weak. The remaining 3 systems are considered in our main analysis.

5.9.1. \( z_{\text{abs}} = 1.5418 \) system toward Q 0002–422

This system is defined by Mg II, Mg I, Al II, Al III, and Fe II absorption lines. The Mg II absorption lines are spread over \( \sim 180 \) km s\(^{-1}\). The other low-ionization lines are seen only in the strongest central component. Voigt profile fits to the absorption lines are shown in Fig. 13.

5.9.2. \( z_{\text{abs}} = 2.1678 \) system toward Q 0002–422

This system is defined by Mg II, Mg I, Al II, Al III, Fe II, Si II, Si IV and C IV absorption lines. Like most systems in our sample the high- and low-ionization profiles are different. The Mg II absorption profile is spread over \( 80 \) km s\(^{-1}\). However, only the strongest central component shows other low-ionization absorption lines. As the central Mg II component is blended in the red wing with other components we use Si II and Al II as anchors. The presence of Fe II \( \lambda 1608 \) with negative \( q \) coefficient makes this system very sensitive to \( \Delta \alpha/\alpha \). The system is fitted with two components and the velocity separation between them is larger than the individual \( b \) values.

5.9.3. \( z_{\text{abs}} = 2.3008 \) system toward Q 0002–422

This system is defined by Mg II, Mg I, Al II, Al III, Fe II, Si II, Si IV and C IV absorption lines. Here also high- and low-ionization profiles are different, like most of other systems. The Mg II absorption profile is spread over \( 200 \) km s\(^{-1}\). The main component is complex and blended. There is a well detached satellite in the blue wing of the profile. We use this for our \( \Delta \alpha/\alpha \) measurement. We also use the unblended Fe II lines and Si II \( \lambda 1526 \) and Al II lines for our analysis. The three components that fit the profile are well resolved.

6. Results

6.1. Fitting procedure

As discussed before, the accuracy of the method depends on how well absorption line profiles are modeled using Voigt profiles. As can be seen from Table 1 the relative shifts between different Fe II lines (apart from Fe II \( \lambda 1608 \)) are small compared to the shifts between the Fe II lines and the anchors. We use this fact to refine our fitting procedure.

We first fit all the Fe II lines simultaneously using laboratory wavelengths (\( \Delta \alpha/\alpha = 0 \)). This allows us to find out about (i) bad pixels, (ii) unknown contaminations and (iii) the velocity component structure in Fe II. We therefore can remove inadequate profiles (because of contamination or bad pixels) from the \( \Delta \alpha/\alpha \) analysis. A similar exercise was carried out for...
Mg II doublets and other anchors. Based on these preliminary fits, a first set of parameters is generated to start the Voigt profile fitting procedure that includes Δα/α variations as described in Sect. 3.3. In all our fits, we have constrained the redshifts parameter of all the species under consideration to be same. We do not know a priori whether the b parameters of different species are identical or not. To take this uncertainty into account we perform the fit of the systems for two cases: (i) assuming that b parameters of individual components are same for all species and (ii) assuming different b parameters for different species. We determine the best-fit Δα/α values for both cases. It is found that in all cases the two values are consistent with one another within 1σ uncertainty. For our final result we consider the fit that gives the smaller reduced χ².

### 6.2. Determination of Δα/α

The χ² analysis for individual systems is shown in Fig. 14. Here, we plot χ² as a function of Δα/α. The dot with error-bar in each panel gives our best-fit estimate of Δα/α. The vertical line in each panel shows the value Δα/α = 0. It is clear from the figure that apart from few cases the curve is well behaved and the best-fit values are consistent with zero in most cases. The numerical values of the fits are given in Table 3. In this table we list in the first four columns the name of the QSO, the emission redshift (z_em), absorption redshifts (z_abs) and velocity dispersions b of individual components. When the best-fit solutions are obtained with different b parameters the b values quoted are for the main anchor line (Mg II, Si II or Al II lines). Column 5 lists the transitions used in the analysis with the same notations as in Table 1. Nc given in Col. 6 is the number of subcomponents used to fit the Fe II absorption profiles. The last two columns give the determination of Δα/α obtained using laboratory wavelengths of Mg II, Mg I, and Si II absorption lines with terrestrial isotopic abundances (case 1) and with the wavelengths of the dominant isotopes (case 2, see next section). Numbers in brackets are the reduced χ² for the fit. The numbers marked with an asterisk (*) are values obtained using different b values for different species.

In Fig. 15 we plot individual determinations of Δα/α as a function of z_abs. We also plot the existing results from the literature. The horizontal dotted line gives the weighted mean...
<table>
<thead>
<tr>
<th>Name</th>
<th>$z_{em}$</th>
<th>$z_{abs}$</th>
<th>$\Delta \alpha/\alpha$</th>
<th>Transitions</th>
<th>$\Delta \alpha/\alpha$ (in units of $10^{-5}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PKS 0237–23</td>
<td>2.222</td>
<td>0.18395</td>
<td>0.1263</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
<tr>
<td>PKS 1239–20</td>
<td>2.190</td>
<td>0.92056</td>
<td>0.9200</td>
<td>$E1431$</td>
<td>1.59</td>
</tr>
<tr>
<td>PKS 1240–20</td>
<td>2.263</td>
<td>0.23616</td>
<td>0.2360</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
<tr>
<td>PKS 1254–23</td>
<td>2.414</td>
<td>0.62504</td>
<td>0.6250</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
<tr>
<td>PKS 1257–23</td>
<td>2.470</td>
<td>0.23616</td>
<td>0.2360</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
<tr>
<td>PKS 1258–23</td>
<td>2.470</td>
<td>0.23616</td>
<td>0.2360</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
<tr>
<td>PKS 1259–23</td>
<td>2.470</td>
<td>0.23616</td>
<td>0.2360</td>
<td>$E14370$</td>
<td>1.54</td>
</tr>
</tbody>
</table>
Fig. 15. Results from the UVES sample: the measured values of $\Delta \alpha/\alpha$ from our sample (filled circles) are plotted against the absorption redshifts of Mg II systems. Each point is the best-fit value obtained for individual systems using $\chi^2$ minimization as shown in Fig. 14. The open circle and stars are the measurements from the Oklo phenomenon and from molecular lines respectively. The weighted mean and 1σ range measured by Murphy et al. (2003) are shown by the horizontal long dashed lines. Clearly most of our measurements are inconsistent with this range.

The shaded region marks the weighted mean and its 3σ error obtained from our study \[
\langle \Delta \alpha/\alpha \rangle_w = (-0.06 \pm 0.06) \times 10^{-5}.
\]

Our data give a 3σ constraint on the variation of $\Delta \alpha/\alpha$ to be $-2.5 \times 10^{-16} \text{ yr}^{-1} \leq (\Delta \alpha/\alpha) \Delta t \leq 1.2 \times 10^{-16} \text{ yr}^{-1}$ in the case of a flat universe with $\Omega_\Lambda = 0.7$, $\Omega_m = 0.3$ and $H_0 = 68 \text{ km s}^{-1} \text{ Mpc}^{-1}$ for a median redshift of 1.55.

Just for completeness we also fitted the weak systems that show at least two Fe II lines. The recovered values of $\Delta \alpha/\alpha$ based on these are given in Table 4. As expected the individual measurements have large errors. The weighted mean value of $\Delta \alpha/\alpha$ measured from these systems is $-0.40 \pm 0.36$. Most of the individual measurements and the weighted mean are consistent with the value obtained from our main sample.

A summary of the results for different sub-samples is given in Table 5. The sample identification is given in the first column. The number of systems used and the median redshift of the samples are given in Cols. 2 and 3. The mean, weighted mean and σ of the measured $\Delta \alpha/\alpha$ values are given in Cols. 4, 5, and 6 respectively. Last column gives the reduced $\chi^2_{\text{W}}$ obtained from all our measurements for the measured weighted mean. It can be seen that there is no sub-sample in which we find a significant change in $\alpha$.

6.3. The magnesium isotopic abundance

One major uncertainty in the many-multiplet analysis comes from the determination of the effective rest wavelengths. Even though laboratory wavelengths are measured with a precision of a few 0.1 mÅ, values given in Table 1 assume terrestrial
Table 4. Results for the weak systems that show at least two detectable Fe II lines.

<table>
<thead>
<tr>
<th>QSO</th>
<th>(z_{\text{abs}})</th>
<th>(\Delta \alpha/\alpha) (in (10^{-5}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q 0122--380</td>
<td>1.9102</td>
<td>−0.9 ± 1.6</td>
</tr>
<tr>
<td>PKS 1448--232</td>
<td>1.5847</td>
<td>+0.4 ± 2.0</td>
</tr>
<tr>
<td>PKS 0237--23</td>
<td>1.1846</td>
<td>−0.9 ± 2.7</td>
</tr>
<tr>
<td>HE 0001--234</td>
<td>0.9489</td>
<td>−1.1 ± 2.3</td>
</tr>
<tr>
<td>HE 2217--2818</td>
<td>1.6277</td>
<td>+0.3 ± 0.6</td>
</tr>
<tr>
<td>Q 0329--385</td>
<td>1.4379</td>
<td>−0.4 ± 0.6</td>
</tr>
<tr>
<td>HE 1347--257</td>
<td>1.5082</td>
<td>−3.9 ± 1.4</td>
</tr>
<tr>
<td>PKS 0329--255</td>
<td>0.9926</td>
<td>−0.3 ± 1.3</td>
</tr>
<tr>
<td>HE 2347--4342</td>
<td>1.7962</td>
<td>−0.4 ± 2.0</td>
</tr>
</tbody>
</table>

isotopic abundances. This assumption may not be valid at high redshift. In astrophysical settings the effect of isotopic shifts could be important for Si and Mg (see Murphy et al. 2003; Ashenfelter et al. 2003). Indeed, the relative abundances of different isotopes may depend on the overall metallicity of the gas. Gay & Lambert (2000) have shown that the abundance of \(^{25}\text{Mg}\) and \(^{26}\text{Mg}\) relative to \(^{24}\text{Mg}\) decreases with decreasing metallicity. In low-metallicity gas (\(Z \lesssim 0.01 Z_\odot\)) most of the metals will be in their dominant isotopic state. Thus in the extreme case of very low metallicity, the effective rest wavelengths could take values in the range from terrestrial to production to wavelengths corresponding to the dominant isotope. This range is less than 0.5 mÅ for Mg I and Si II absorption lines but is of the order of 1 mÅ for Mg II lines. To accommodate this uncertainty we fit the systems using the wavelengths of the species from the dominant isotope. The measurements are given in the last column of Table 3. As expected, using these abundances leads to a lower \(\alpha\) determination (see Table 5). Note however that even in this extreme case, the variation stays smaller than what has been claimed from previous studies. Note that the assumption of very low metallicity is extreme as: (i) the systems in the redshift range 0.4--2 are more likely to have metallicity larger than 0.1 \(Z_\odot\) (e.g. Ledoux et al. 2002); (ii) the measured mean ratios of \(^{24}\text{Mg}^{25}\text{Mg}^{26}\) in a cool dwarf with metallicity \(Z = −1.5\) to \(−1.0 Z_\odot\) is 80:10:10 (from Table 1 of Yong et al. 2003). This gives weighted mean wavelengths close to the terrestrial wavelengths; and (iii) We also notice that the minimum \(\chi^2\) for the fit in most cases (apart from 4 cases) is lower when we use the laboratory wavelengths in our analysis. Therefore, although some additional uncertainty and scatter could come from the isotopic abundances being different from that of terrestrial ones, our result using the laboratory wavelengths is most probably robust.

7. Conclusion

We have applied the MM method to a homogeneous sample of 50 Mg II systems observed along 18 QSO lines of sight observed with UVES at the VLT. Using extensive simulations we show that our Voigt profile fitting procedure works well for the recovery of the input value of \(\Delta \alpha/\alpha\) for simple single component systems. We can recover the input \(\Delta \alpha/\alpha\) values with an error of \(\sigma = 0.23 \times 10^{-5}\) in the case of individual components for data of quality comparable to the UVES data (\(S/N \sim 70\)). We show that the uncertainty is about twice as large for data of \(S/N \sim 30\) as used by previous surveys. In addition at this \(S/N\) ratio, the use of strongly blended systems leads to larger uncertainties. We show that weak lines should be excluded from the determination as their use can lead to a false determination of \(\Delta \alpha/\alpha\).

We devise the selection criteria for our sample using extensive simulations. This leads to 23 systems excluding strongly blended systems and systems with weak absorption lines. Individual systems and fits to all absorption lines are discussed in detail. We also avoided the two sub-DLAs from the analysis. In one of the systems that is in common with Murphy et al. (2003) our results are consistent with the earlier measurement albeit with smaller error (see Sect. 5.3.1).

The weighted mean value of the variation in the fine-structure constant obtained from our analysis over the redshift range \(0.4 \leq z \leq 2.3\) is \(\Delta \alpha/\alpha = (−0.06 \pm 0.06) \times 10^{-5}\). For the median redshift of our sample (\(z = 1.54\)) we obtain a 3\(\sigma\) constraint on the variation of \(\Delta \alpha/\alpha\) to be \(< 2.5 \times 10^{-16} \text{ yr}^{-1} < (\Delta \alpha/\alpha \Delta t) < 1.2 \times 10^{-16} \text{ yr}^{-1}\) in the case of a flat universe with \(\Omega_m = 0.7\), \(\Omega_b = 0.3\) and \(H_0 = 68 \text{ km s}^{-1} \text{ Mpc}^{-1}\). To our knowledge this is the strongest constraint from QSO absorption line studies to date.

We show the effect of varying the isotopic abundances. Only if we assume isotopic abundances for very low metallicity would \(\alpha\) show some variation. This is not acceptable for (i) assuming that all systems in the redshift range 0.4--2 have extremely low metallicity is incorrect as metals are seen in these systems and metallicity is measured to be larger than 0.1 \(Z_\odot\) (e.g. Ledoux et al. 2002); (ii) at \(Z < 0.1 Z_\odot\), which would be a more reasonable assumption, the composite wavelength we get will be close to the laboratory value we have used in our study (we use Yong et al. 2003 for isotopic abundances). Therefore, although some additional uncertainty and scatter could come from the isotopic abundances being different from terrestrial ones, our result using laboratory wavelengths is most probably robust.

In conclusion, our study does not support claims by previous authors of a statistically significant change in \(\Delta \alpha/\alpha\) with cosmic time at \(z > 0.5\). Our result still does allow smaller variations in excess of what is found based on the Oklo phenomenon. Future very high resolution \((R \sim 100,000)\) spectroscopic studies are needed to probe the variations in \(\alpha\) with much better accuracy.

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Table 5. Summary of results for various sub-samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Number of systems</th>
<th>$z$</th>
<th>$\Delta\alpha/\alpha$ $(10^{-5})$</th>
<th>$\chi^2$</th>
<th>$\sigma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single + double (case 1)</td>
<td>12</td>
<td>1.54</td>
<td>$+0.01 \pm 0.15$</td>
<td>$-0.08 \pm 0.07$</td>
<td>0.27</td>
</tr>
<tr>
<td>Full sample (case 1)</td>
<td>23</td>
<td>1.54</td>
<td>$-0.02 \pm 0.10$</td>
<td>$-0.06 \pm 0.06$</td>
<td>0.41</td>
</tr>
<tr>
<td>Weak (case 1)</td>
<td>9</td>
<td>1.51</td>
<td>$-0.80 \pm 0.58$</td>
<td>$-0.40 \pm 0.36$</td>
<td>1.27</td>
</tr>
<tr>
<td>Weak + All (case 1)</td>
<td>32</td>
<td>1.54</td>
<td>$-0.24 \pm 0.18$</td>
<td>$-0.07 \pm 0.06$</td>
<td>0.81</td>
</tr>
<tr>
<td>Single + double (case 2)</td>
<td>12</td>
<td>1.54</td>
<td>$-0.31 \pm 0.14$</td>
<td>$-0.33 \pm 0.09$</td>
<td>0.33</td>
</tr>
<tr>
<td>Full Sample (case 2)</td>
<td>23</td>
<td>1.54</td>
<td>$-0.33 \pm 0.11$</td>
<td>$-0.36 \pm 0.06$</td>
<td>0.44</td>
</tr>
</tbody>
</table>

Case 1: laboratory wavelengths given in Table 1 are used.
Case 2: rest wavelengths of dominant isotopes for Mg and Si are used.

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