



Searching for space-time variation of the fine structure constant using QSO spectra: overview and future prospects

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Abstract. Theories unifying gravity with other interactions suggest the possibility of spatial and temporal variation of fundamental “constants” in the Universe. Using quasar absorption systems we can probe the value of the fine-structure constant, $\alpha = e^2/\hbar c$, over the history of the universe. Previous studies of three independent samples of data, containing 143 absorption systems spread from 2 to 10 billion years after big bang, hint that α was smaller 7 – 11 billion years ago. However competing studies show no such α -variation. The studies can be improved by utilising more atomic transitions that are seen in quasar spectra, however in many cases this is hampered by a lack of accurate laboratory frequency measurements. The aim of this paper is to provide a compilation of transitions of importance to the search for α variation. They are E1 transitions from the ground state in several different atoms and ions, with wavelengths ranging from around 900 – 6000 Å, and require an accuracy of better than 10^{-4} Å. We also discuss isotope shift measurements that are needed in order to resolve systematic effects in the study. Researchers who are interested in performing these measurements should contact the authors directly.

1. Introduction

Current theories that seek to unify gravity with the other fundamental interactions suggest that spatial and temporal variation of fundamental constants is a possibility, or even a necessity, in an expanding Universe (see, for example the

review of Uzan (2003)). Several studies have tried to probe the values of constants at earlier stages in the evolution of the Universe, using tools such as big-bang nucleosynthesis, the Oklo natural nuclear reactor, quasar absorption spectra, and atomic clocks (see, e.g. Flambaum & Berengut 2009).

Table 1: High-priority transitions observed in QSO spectra. ‘Status’ refers to the experimental status as explained in the text.

Atom/ Ion	Wavelength λ (Å)	Frequency ω_0 (cm ⁻¹)	Oscillator Strength	q value (cm ⁻¹)	Status	Refs.
C I	945.188	105799.1	0.272600	130 (60)	M	[1]
	1139.793	87735.30	0.013960	0 (100)	B	
	1155.809	86519.47	0.017250	0 (100)	B	
	1157.186	86416.55	0.549500	0 (100)	B	
	1157.910	86362.52	0.021780	0 (100)	B	
	1188.833	84116.09	0.016760	0 (100)	B	
	1193.031	83820.13	0.044470	0 (100)	B	
	1193.996	83752.41	0.009407	0 (100)	B	
	1260.736	79318.78	0.039370	30 (10)	A	
	1276.483	78340.28	0.004502	17 (10)	A	
	1277.245	78293.49	0.096650	-13 (10)	A	
	1280.135	78116.74	0.024320	-21 (10)	A	
	1328.833	75253.97	0.058040	117 (10)	A	
	1560.309	64089.85	0.080410	137 (10)	A	
	1656.928	60352.63	0.140500	-24 (10)	A	
	C II	1036.337	96493.74	0.123000	168 (10)	
1037.018*		96430.32	0.123000	105 (10)	A	
1334.532		74932.62	0.127800	178 (10)	A	
1335.662*		74869.20	0.012770	115 (10)	A	
1335.707*		74866.68	0.114900	118 (10)	A	
C III	977.020	102352.0	0.762000	165 (10)	B	
C IV	1548.204	64590.99	0.190800	222 (2)	A	[2]
	1550.781	64483.65	0.095220	115 (2)	A	[2]
O I	1025.762	97488.54	0.020300	0 (20)	B	
	1026.476	97420.72	0.002460	0 (20)	B	
	1039.230	96225.05	0.009197	0 (20)	B	
	1302.168	76794.98	0.048870	0 (20)	A	
Na I	3303.320	30272.58	0.013400	57 (2)	B	
	3303.930	30266.99	0.006700	51 (2)	B	
	5891.583	16973.37	0.655000	62 (2)	M	[3,4,5]
Mg I	5897.558	16956.17	0.327000	45 (2)	M	[3,6]
	2026.477	49346.73	0.112000	87 (7)	M	[7,8]
	2852.963	35051.27	0.181000	90 (10)	M	[7,9,10,11,12]
Mg II	1239.925	80650.04	0.000267	192 (2)	B	
	2796.354	35760.85	0.612300	212 (2)	M	[7,9,13,14]
	2803.532	35669.30	0.305400	121 (2)	M	[7,9,14]
Al II	1670.789	59851.97	1.880000	270 (30)	M	[2]
Al III	1854.718	53916.54	0.539000	458 (6)	M	[2]
	1862.791	53682.88	0.268000	224 (8)	M	[2]
Si II	1190.416	84004.26	0.250200		B	
	1193.290	83801.95	0.499100		B	
	1260.422	79338.50	1.007000		A	
	1304.370	76665.35	0.094000		A	
	1526.707	65500.45	0.117094	50 (30)	I	[2]
	1808.013	55309.34	0.002010	520 (30)	I	[2]

Table 1: (continued)

Atom/ Ion	Wavelength λ (Å)	Frequency ω_0 (cm ⁻¹)	Oscillator Strength	q value (cm ⁻¹)	Status	Refs.
Si IV	1393.760	71748.64	0.528000	823 (40)	I	[2]
	1402.773	71287.54	0.262000	361 (15)	I	[2]
S II	1250.583	79962.61	0.005350		A	
	1253.808	79756.83	0.010700		A	
	1259.518	79395.39	0.015900		A	
Ca II	3934.777	25414.40	0.688000	446 (6)	M	[15]
	3969.591	25191.51	0.341000	222 (2)	M	[15]
Ti II	1910.600	52339.58	0.202000	-1564 (150)	A	
	1910.938	52330.32	0.098000	-1783 (300)	A	
	3067.245	32602.55	0.041500	791 (50)	I	[7]
	3073.877	32532.21	0.104000	677 (50)	I	[7]
	3230.131	30958.50	0.057300	673 (50)	I	[7]
	3242.929	30836.32	0.183000	541 (50)	I	[7]
	3384.740	29544.37	0.282000	396 (50)	I	[7]
Cr II	2056.256	48632.06	0.105000	-1110 (150)	I	[7,16]
	2062.236	48491.05	0.078000	-1280 (150)	I	[7,16]
	2066.164	48398.87	0.051500	-1360 (150)	I	[7,16]
Mn II	1197.184	83529.35	0.156600	-2556 (450)	B	
	1199.391	83375.65	0.105900	-2825 (450)	B	
	1201.118	83255.77	0.088090	-3033 (450)	B	
	2576.877	38806.66	0.288000	1276 (150)	I	[7]
	2594.499	38543.08	0.223000	1030 (150)	I	[7]
Fe II	2606.462	38366.18	0.158000	869 (150)	I	[7]
	1063.176	94057.80	0.060000		B	
	1063.971	93987.52	0.003718		B	
	1096.877	91167.92	0.032400		B	
	1121.975	89128.55	0.020200		B	
	1125.448	88853.51	0.016000		B	
	1143.226	87471.77	0.017700		B	
	1144.939	87340.98	0.106000		B	
	1260.533	79331.52	0.025000		A	
	1608.450	62171.63	0.058000	-1030 (300)	A ⁰	[17]
	1611.200	62065.53	0.001360	1560 (500)	A ⁰	[17]
	2249.877	44446.88	0.001821		A	
	2260.780	44232.51	0.002440		I	[7]

Table 1: (continued)

Atom/ Ion	Wavelength λ (Å)	Frequency ω_0 (cm ⁻¹)	Oscillator Strength	q value (cm ⁻¹)	Status	Refs.
Fe II	2344.212	42658.24	0.114000	1540 (400)	I	[7,18]
	2367.589	42237.06	0.000212	1904	A	
	2374.460	42114.83	0.031300	1660 (60)	I	[7,18]
	2382.764	41968.06	0.320000	1550 (60)	I	[7,18]
	2586.649	38660.05	0.069180	1540 (40)	I	[7,18]
	2600.172	38458.99	0.238780	1410 (60)	I	[7,18]
Ni II	1317.217	75917.64	0.146000		A	
	1370.132	72985.67	0.076900		A	
	1393.324	71770.82	0.022220		A	
	1454.842	68735.99	0.032300		A	
	1467.259	68154.29	0.009900		B	
	1467.756	68131.22	0.006300		B	
	1502.148	66571.34	0.006000		B	
	1703.412	58705.71	0.012240		A	[16]
	1709.604	58493.07	0.032400	-20 (250)	I	[16]
	1741.553	57420.01	0.042700	-1400 (250)	I	[16]
	1751.915	57080.37	0.027700	-700 (250)	I	[16]
Zn II	2026.137	49355.00	0.489000	2470 (25)	M	[7,16,19]
	2062.660	48481.08	0.256000	1560 (25)	I	[7,16]

⁰ The wavelength measurements of the Fe II 1608 and 1611 lines have been questioned (G. Nave, IAU General Assembly, 2009) and urgently need reanalysis.

¹ Labazan et al. (2005)

² Griesmann & Kling (2000)

³ Juncar et al. (1981)

⁴ Pescht et al. (1977)

⁵ Huber et al. (1978)

⁶ Gangrsky et al. (1998)

⁷ Aldenius et al. (2006)

⁸ Hannemann et al. (2006)

⁹ Pickering et al. (1998)

¹⁰ Hallstadius (1979)

¹¹ Boiteux et al. (1988)

¹² Salumbides et al. (2006)

¹³ Drullinger et al. (1980)

¹⁴ Batteiger et al. (2009)

¹⁵ Wolf et al. (2008)

¹⁶ Pickering et al. (2000)

¹⁷ Pickering et al. (2002)

¹⁸ Nave et al. (1991)

¹⁹ Matsubara et al. (2003)

Comparison of atomic transition frequencies on Earth and in quasar (QSO) absorption spectra can be used to measure variation of the fine-structure constant $\alpha = e^2/\hbar c$ over the last 10 billion years or so. Early studies used the “alkali-doublet” method (Savedoff, 1956), taking advantage of the simple α -dependence of the separation of a fine-structure multiplet.

More recently we developed the “many-multiplet” method (Dzuba et al., 1999a,b) which improves sensitivity to variation in α by more than an order of magnitude compared to the alkali-doublet method. Enhancement comes from the use of transitions which are more sensitive to α than the fine-structure splitting is, for example the s -wave orbital has maximum relativistic corrections to energy but no spin-orbit splitting. In addition the α -dependence varies strongly between different atoms and transitions (for example s - p and s - d transitions can have different signs) and this helps to control systematics. The number of spectral lines available for study is quite large; this gives a statistical advantage.

The first analyses using the many-multiplet method revealed hints that the fine structure constant was smaller in the early universe (Murphy et al., 2003a, 2001b,c; Webb et al., 1999, 2003, 2001). The latest results of this group, which combine data from 143 absorption systems over the redshift range $0.2 < z_{\text{abs}} < 4.2$, gives a variation of $\Delta\alpha/\alpha = (-0.57 \pm 0.11) \times 10^{-5}$ (Murphy et al., 2004). A very extensive search for possible systematic errors has shown that known systematic effects cannot explain the result (Murphy et al., 2001a). Our method and calculations have been used by other groups to analyse different data sets from different telescopes (Levshakov et al., 2005; Quast et al., 2004; Srianand et al., 2004), however their results indicate no variation of α . Recently the methodology of Srianand et al. (2004) was questioned by Murphy et al. (2007). A re-analysis of the same data, which included 23 absorption systems from VLT/UVES, gave a revised fit of $\Delta\alpha/\alpha = (-0.64 \pm 0.36) \times 10^{-5}$ (Murphy et al., 2008), increasing the error previously reported by a factor of six.

2. Discussion

To continue this work and resolve the discrepancies, several new transitions are being considered. In Table 1 we present a list of lines commonly observed in high-resolution QSO spectra. All of the lines marked ‘A’ (very important) or ‘B’ (mildly important) lack the high-accuracy laboratory measurements necessary for studies of α variation. All

transitions are from the ground state of the ion, with the exception of the C II lines marked with an asterisk which are transitions from the metastable $2s^2 2p^2 P_{3/2}^o$ level. Predominantly the wavelengths and oscillator strengths are taken from the compilations of Morton (1991, 2003). The wavelengths have errors of about 0.005 Å, although it is possible that some errors are closer to 0.05 Å. Note that the oscillator strengths presented are not as accurate as the wavelengths: these measurements are much more difficult. As a general rule, the lines are more important for α variation if they lie above 1215.67 Å (the Lyman- α line of hydrogen) due to the “Lyman- α forest” seen in QSO spectra.

Isotope shift measurements for these transitions are also needed in order to resolve a source of systematic error in the variation of α studies: the isotope abundance ratios in the gas clouds sampled in the quasar absorption spectra may not match those on Earth (Murphy et al., 2001a, 2003b). Spurious observation of α -variation due to differences in isotope abundance of any one element has been ruled out, however an improbable “conspiracy” of changes in several elements could mimic the observed effect.

Accurate measurements of the isotope shift are required to quantify these systematic effects. Additionally, if the isotope shifts are known then it is possible to simultaneously determine both any possible α -variation and the isotope abundances in the early universe directly (Kozlov et al., 2004). This can be used to constrain models of chemical evolution of the Universe and test models of nuclear processes in stars (Ashenfelter et al., 2004; Fenner et al., 2005). We have performed very complicated calculations of these isotope shifts (Berengut et al., 2003, 2005, 2006, 2008), however calculations in group 3d atoms and ions are difficult, and our accuracy may be low. Therefore measurements for at least some lines are needed to benchmark calculations in this regime. In addition to the transitions previously mentioned, in Table 1 we present lines that were used in previous studies (and hence have precise wavelength measurements), but for which the isotopic structure has not been measured. These transitions are marked with an ‘I’. Where both the transition frequency and isotope shift are known they are marked with an ‘M’: these are included here for reference only, however verification would still be useful.

We previously calculated the relativistic energy shifts, or q -values, for many of the lines seen in quasar spectra (Berengut et al., 2004, 2005, 2006; Dzuba et al., 2002; Dzuba & Johnson, 2007; Porsev et al., 2007; Savukov & Dzuba, 2008). The dif-

ference between the transition frequencies in QSO spectra (ω) and in the laboratory (ω_0) depends on the relative values of α . The dependence of the frequencies on small changes in α is given by the formula $\omega = \omega_0 + qx$, where $x = (\alpha/\alpha_0)^2 - 1$. The q values are calculated using atomic physics codes. The atomic energy levels are calculated to a first approximation using relativistic Hartree-Fock (Dirac-Hartree-Fock). Higher order effects are taken into account using a combination of configuration interaction (for many-valence-electron systems) and many-body perturbation theory; this is known as the “CI+MBPT” method (Dzuba et al., 1996). The value of α is varied in the computer codes and the energy levels are recalculated, and hence the transition frequencies. The q values are extracted as

$$q = \left. \frac{d\omega}{dx} \right|_{x=0} \quad (1)$$

We also account for complications due to level pseudo-crossing as described by Dzuba et al. (2002). In Table 1 we present our current best q -values for easy reference. Uncertainties here are representative rather than statistical.

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