The solar oxygen abundance from an empirical three-dimensional model

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ABSTRACT

The oxygen abundance in the solar photosphere, and consequently the solar metallicity itself, is still a controversial question with far-reaching implications in many areas of astrophysics. This paper presents a new determination obtained by fitting the forbidden O I line at 6300 Å with an observational 3D model. The approach presented here is novel because previous determinations were based either on 1D empirical stratifications or on 3D theoretical models. The resulting best-fit abundances are log $\epsilon$(O)=8.90 and log $\epsilon$(Ni)=6.15. Nevertheless, by introducing minor tweaks in the model and the procedure, it is possible to retrieve very different values, even down to log $\epsilon$(O)=8.70. This extreme sensitivity of the abundance to possible systematic effects is not specific to this particular work, but probably reflects the real uncertainty inherent to all abundance determinations based on a prescribed model atmosphere.


1. Introduction

Oxygen is, after hydrogen and helium, the third most abundant element in the Universe. A precise knowledge of its abundance in stellar interiors is of critical importance in many areas of astrophysics because of its contribution to opacity and free electrons, and also because it serves as a reference for other relevant elements that cannot be directly measured in the photospheric spectrum. Unfortunately, there are reasons to believe that our knowledge of even the solar oxygen abundance is still far from being precise. Several factors conspire to make it so elusive. First of all, oxygen is a very volatile element and therefore its meteoritic abundance is not a valid proxy. Second, there are very few atomic indicators in the photospheric solar spectrum, and they are all either extremely weak forbidden lines or complex transitions suffering from non-LTE effects and affected by additional uncertainties in the relevant atomic processes.

In spite of the uncertainties, the prevailing paradigm that emerged since the 1970s was that the solar oxygen abundance was log $\epsilon$(O) in the range between 8.8 and 8.9, in the logarithmic scale commonly used in astrophysics, where H has a reference value of 12 (e.g., Cameron 1973; Anders & Grevesse 1989; Grevesse & Sauval 1998). In all cases, the procedure of determining chemical abundances was always the same: From a prescribed model atmosphere, synthetic spectral lines are computed, and the abundances are adjusted until a satisfactory agreement is found with some atlas observations. The models employed were one-dimensional (1D), usually derived empirically by fitting very many lines and continua, and the photospheric observations were unresolved in space and time.

The paradigm was called into question by the controversial paper of Asplund et al. (2004) (hereafter AGSAK), who claimed that the solar O abundance needed to be revised downward by almost a factor of two (log $\epsilon$(O)=8.66). The procedure was conceptually similar to what had been done before, except that in addition to updated atomic parameters, the authors employed a three-dimensional (3D) model resulting from a hydrodynamical numerical simulation. Obviously, such a large revision would have a considerable impact on many areas of astrophysics. In particular, it would ruin the existing excellent agreement between solar interior models and measurements from helioseismology. Attempts to reconcile the models with the new abundances have been unsuccessful thus far (e.g., Basu & Antia 2008 and references therein; Basu & Antia 2013).

The debate on whether the proposed revision should be adopted has been very intense, and several papers have been published with new abundance determinations, including those of Ayres et al. (2006); Socas-Navarro & Norton (2007); Ayres (2008); Centeno & Socas-Navarro (2008); Caffau et al. (2008); Scott et al. (2009); Pereira et al. (2009); Caffau et al. (2011).

The frequency of publications on the so-called solar oxygen crisis seems to have declined in recent years, not because the problem has been satisfactorily resolved, but probably because there are no new arguments or data in favor of one view or the other. Regardless of the final outcome, the work of AGSAK has been of great importance because at the very least, it has blown the whistle on a grossly overlooked problem, namely how much we can trust our abundance determinations. The choice of a suitable atmospheric model is critical, and the model uncertainties are not yet clear, nor is it established how these uncertainties propagate into the final result. Ayres (2008) correctly remarked that while a 3D atmosphere is preferable over a 1D atmosphere, it is not clear that a model resulting from a numerical simulation is better than an empirical one when it is employed to fit observations. In addition, the work of Caffau et al. (2008, 2011) demonstrated that working in 3D does not necessarily result in lower abundances.

The present paper is an attempt to resolve the dilemma of theoretical 3D versus empirical 1D by taking the best of both approaches. A 3D model obtained from observations is used to derive the O and Ni abundances in the conventional way (i.e., fitting unresolved atlas observations). As in my previous works on this subject (Socas-Navarro & Norton 2007; Centeno & Socas-Navarro 2008), the focus is not so much the final result, but in-
introducing a novel methodology in the hope that it might open a new path for better, more robust determinations in the future.

2. Observations and the model

The 3D model was derived from a reanalysis of the data used in Socas-Navarro (2011, hereafter SN11) with a slightly different approach and a new version of the code NICOLE (Socas-Navarro et al. 2014). The new version incorporates several improvements in accuracy and stability that make the model smoother and with a lower inversion noise (the pixel-to-pixel fluctuation exhibited by the retrieved parameters, such as the photospheric temperature). Some other improvements include the following:

- The new NICOLE version supports two-component inversions. Pixels that exhibit a polarization signal above a certain threshold (32% of the total) are inverted here with a full two-component treatment, which is more consistent than the approach taken in SN11.
- A regularization term is added to the χ² merit function to favor (when possible) smoother solutions over those with spurious high-frequency vertical fluctuations.
- A Bezier spline interpolation scheme is now used to reconstruct the model atmosphere from the node values. Previously, the model was constructed from linear segments between the nodes. The new scheme results in smoother height runs of the retrieved physical parameters, thus removing unsightly sharp corners and at the same time minimizing the possibility of overshooting between the nodes. For stability reasons, it is important to keep the model within reasonable ranges of the various physical parameters at every step of the iterative procedure. The algorithm sets constrains on the node values, but the interpolation between nodes might overshoot beyond the safe range. This risk is minimized by the use of Bezier interpolation, which behaves similarly to splines with tension.
- The new version includes hyperfine structure arising from the angular momentum coupling between electron and nuclear spin.

The dataset from which the model has been derived is exactly the same as in SN11. A brief description is provided here for the sake of completeness, but the reader is referred to that paper for more details. The data come from the Spectro-Polarimeter (SP) of the Hinode satellite’s Solar Optical Telescope (SOT; Kosugi et al. 2007; Ichimoto et al. 2008; Shimizu et al. 2008; Suematsu et al. 2008; Tsuneta et al. 2008), in particular from a quiet-Sun observation acquired at UT 19:32:10 on 2007 September 24.

The SP scan spans the wavelength range between 6300.89 and 6303.29 Å with a sampling of 21.4 mÅ. Unfortunately, the O/Ni blend at 6300.27 Å that is the target of this study is just outside the observed range. It is therefore impossible to perform a pixel-by-pixel abundance analysis here in the way of Socas-Navarro & Norton (2007), which would be of great interest to gauge with high confidence and precision the total error of the entire procedure, for instance. Instead, element abundances are derived here by fitting the disk-center average intensity from the Kitt Peak Fourier Transform Spectrometer (FTS) atlas of Neckel & Labs (1984), following the traditional procedure of synthesizing line profiles from the model and taking the spatial average of the synthetic spectra to compare with the atlas. Absolute wavelengths were obtained by comparing the average observed spectrum to those of the FTS atlas. The spectral point spread function of the Hinode SP is known and has been considered in the analysis by applying it on the emerging synthetic spectra. This means that at each iterative step, the synthetic profiles produced by the proposed model are convolved with the Hinode PSF before they comparing them to the observed profiles.

The field of view observed by the Hinode SOT is a very quiet region at disk center. Spatial sampling is about 0.15″, which is approximately half of the actual spatial resolution given by the SOT point spread function. Standard flatfielding and polariometric calibration procedures were applied to the spectra. Some other corrections were applied a posteriori, as detailed in SN11.

The inversion fits both Fe i lines at 6301.5 Å and 6302.5 Å simultaneously. There are some small departures from LTE (Shchukina & Trujillo Bueno 2001) that are taken into account in the procedure using the approximation described in SN11. Three different inversions were carried out by varying the assumed Fe abundance (7.40, 7.45, and 7.50), resulting in three slightly different 3D models. This is helpful to assess the influence of our uncertainties in the model on the abundances obtained.

The atomic line data employed are listed in Table 1. They are mostly those resulting from a query to the VALD database (Piskunov et al. 1995; Kupka et al. 1999; Kupka et al. 2000), following the traditional procedure of synthesizing unsightly sharp corners and at the same time minimizing the possibility of overshooting between the nodes. For stability reasons, it is important to keep the model within reasonable ranges of the various physical parameters at every step of the iterative procedure. The algorithm sets constrains on the node values, but the interpolation between nodes might overshoot beyond the safe range. This risk is minimized by the use of Bezier interpolation, which behaves similarly to splines with tension.

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As noted above, the inversion is made differently depending on whether the spatial pixel to invert is magnetic or not. Magnetic pixels are defined as those that exhibit polarization signals above the 3 − σ noise level in any one of the Stokes profiles Q, U or V. In both cases the model is initialized with the Harvard-Smithsonian Reference Atmosphere (HSRA) as a starting guess (Gingerich et al. 1971) and then iterated in two successive cycles. The number of nodes used in the inversions for the various physical parameters is listed in Table 2. Note that the inversion corresponding to the second cycle of magnetic pixels uses a two-component model, which has a different number of nodes for each component. In these cases (magnetic pixels), the atmosphere was parameterized as consisting of a magnetic component (Comp 2 in the table) coexisting with some filling factor inside the pixel with a non-magnetic surrounding (Comp 1).

Some representative fits are shown in Figure 1 to give an idea of how accurately the synthetic lines match the observations. Perhaps the most relevant plot for the discussion here is the top left panel, which shows the spatially averaged profiles. This is not a fit but, the result of combining all the individual synthetic profiles and comparing them to the average of the ob-

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Fig. 1. Some representative fits obtained with the empirical 3D model. In all panels the solid line represents the Hinode observations, the dashed line the synthetic profile computed with NICOLE. Upper left: Spatial average over the entire field of view. Lower left: Fit to a non-magnetic profile whose $\chi^2$ is similar to the average $\chi^2$ in all the non-magnetic pixels. The quality of this fit may then be considered as typical of the entire region. Right panels: Fits to Stokes $I$ (upper) and $V$ (lower) profiles from a pixel with a magnetic signal whose $\chi^2$ is similar to the average $\chi^2$ in all the magnetic pixels. The quality of this fit may then be considered as typical of the entire region.

Figure 2 illustrates the spatial distribution of some sample quantities. No effort has been made to place the model on a common geometrical scale since that is irrelevant for the purpose of computing the emerging profile from each model column. Therefore, the reference height scale employed in this paper is the monochromatic continuum optical depth at 500 nm, $\log(\tau_{500})$.

As with the earlier version, the new 3D model is publicly available and may be downloaded from the CDS via anonymous ftp to cdsarc.u-strasbg.fr (130.79.128.5) or via http://cdsweb.u-strasbg.fr/cgi-bin/qcat?J/A+A/.

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3. Analysis

3.1. The Sc\(\text{ii}\) line and the missing dynamics

Very close to the O/Ni blend that constitutes the main target of this study lies another interesting spectral feature of similarly weak nature, emerging from a Sc\(\text{ii}\) transition. This line is of great interest for reasons explained below, and it has been included in the synthetic spectrum computed from the 3D model. To calculate the Sc\(\text{ii}\) profile, its hyperfine structure needs to be considered. Fortunately, accurate atomic parameters exist in the literature (e.g., in the VALD database), and these values can easily be fed into NICOLE to produce realistic line shapes.

The inclusion of hyperfine structure for the Sc\(\text{ii}\) line slows down the calculation by nearly a factor 100. As a result, computing the 200x200 columns of the 3D model takes a significant (but still acceptable) amount of time. However, it quickly becomes prohibitive to repeat the full model synthesis for each point in the three-parameter grid of abundances of Fe, Ni, and O required for the abundance determination in Sect. 4 below. It is then convenient, if not entirely necessary, to find computationally cheaper alternatives.

The overall effect of the hyperfine structure on the spatially averaged Sc\(\text{ii}\) intensity profile is simply a broadening of the
Fig. 2. Spatial distribution of various magnitudes derived from the empirical 3D model used in this work. Top panels: Horizontal cuts of the temperature at the base of the photosphere (left) and at the height where $\log(\tau_{500}) = 10^{-2}$ (right). The rest of the panels show a comparison between various observed (left) and synthetic (right) quantities.
Fig. 5. Response functions (in absolute value) of the spectral features considered in this work. The O\textsubscript{Ni} blend and the Sc\textsubscript{ii} response that overlaps almost entirely with that of the continuum. To show has been subtracted from each wavelength in this figure. Each response function has been normalized to its highest value to facilitate visualization.

The discrepancy of the Sc\textsubscript{ii} observed profile. ble to deepen the line core by increasing its abundance or oscillator strength, but it would still be too narrow compared to the line. With some experimentation, it is straightforward to see that the average Sc\textsubscript{ii} synthetic profile emerging from the 3D model is nearly identical to one computed without hyperfine structure, but adding a microturbulent velocity of 0.8 km s\textsuperscript{-1}, as seen in Fig (3). The extra parameters needed for the hyperfine structure calculation in addition to those from Table 1 (Young et al. 1988; Arnesen et al. 1982) are \( I^{up}_{1}=3.5, A_{low}=-27.9, B_{low}=19, A_{up}=125.4, B_{up}=7, \) with all A and B constants in MHz. In view of this result, all the calculations shown in the remainder of this paper have been carried out simulating the effects of hyperfine structure in the Sc\textsubscript{ii} line by applying an “artificial” microturbulence of 0.8 km s\textsuperscript{-1} (only to this line!), which is a very good approximation for the spatially averaged intensity.

A first comparison of the synthetic spectrum to the atlas observation, plotted in Fig (4), shows that it is possible to reach a reasonable agreement in the O/Ni blend, but only a very poor fit to the Sc\textsubscript{ii} line. The abundance values for the calculation in the figure are \( \log e(\text{Fe})=7.50, \log e(O)=8.80, \) and \( \log e(Ni)=6.20. \) The discrepancy of the Sc\textsubscript{ii} feature is too large to be reconciled by tweaking its abundance or the atomic parameters. It is possible to deepen the line core by increasing its abundance or oscillator strength, but it would still be too narrow compared to the observed profile.

To bring the synthetic Sc\textsubscript{ii} line into agreement with the observation, it is necessary to add some additional broadening. The natural explanation for the missing broadening is to assume that even with Hinode’s high spatial resolution, there are still some small-scale plasma flows at the base of the photosphere that are not being sufficiently resolved in the observations. It is important to note that the Fe\textsubscript{i} lines are very well reproduced without requiring any microturbulence, which indicates that most of the small-scale dynamics seen by these lines has been properly captured in the 3D model. Therefore, the missing dynamics is probably located in the deeper layers, where the O/Ni and the Sc\textsubscript{ii} lines form but the Fe\textsubscript{i} lines have very little sensitivity.

Figure 5 shows the response functions of the lines considered here in a typical quiet-Sun model (HSRA) to illustrate their respective formation heights. Response functions describe how the emerging profiles are altered when the atmosphere is perturbed at a given point and constitute a good indicator of the line formation height. In all cases, the continuum response was subtracted. The response for each line was normalized to its maximum because otherwise the Fe\textsubscript{i} line would dominate, making the others barely visible. The figure clearly shows that the Sc\textsubscript{ii} feature samples the same heights as the O/Ni blend, whereas the Fe\textsubscript{i} line spans a much broader atmospheric range.

A comparison of the 3D model employed here with the hydrodynamical simulation of AGSAK further supports the notion of the missing dynamics scenario. The \( rms \) spread of vertical velocities at the \( \log(\tau_{500})=1 \) level is much broader in the simulation than in the 3D empirical model. With the information provided by the Sc\textsubscript{ii} line, it is straightforward to correct this problem by applying an ad hoc enhancement factor \( v_{enh} \) to the velocities in the model and tweaking this factor until the line is properly fitted. Since the Sc\textsubscript{ii} line has a strength and formation region very similar to the O/Ni blend, this enhancement will also correct any possible missing broadening in the O and Ni fea-
tures. After some experimentation, the optimal rms spread for the Sc ii line was found to be \( v_{rms} = 1.36 \text{ km s}^{-1} \), about a factor of 2 greater than in the empirical model. This figure is considerably lower than the 2.47 km s\(^{-1}\) of the AGSAK model. In fact, adopting their value of the rms velocity spread would result in excessive broadening of the observed Sc ii line, incompatible with the observations of these weak lines. Reproducing the behavior of small-scale dynamics in stellar atmospheres is not a trivial affair and important challenges still exist (Steffen et al. 2013). For our purposes here, we simply applied an ad-hoc enhancement factor to our model that was derived empirically and independently from the Sc ii line. After this correction, the agreement between observed and average synthetic profiles improves enormously, as shown below.

4. Abundance determinations

With a 3D model that accurately reproduces the Sc ii line, we can now proceed with confidence to derive the O and Ni abundance by the usual procedure. One computes the synthetic profiles of the O/Ni blend for an array of \([\log \epsilon(O), \log \epsilon(Ni)]\) values at each model column, takes the spatial average over the entire field of view and compares the result to the observation in order to determine which pair of abundances produces the best match. The comparison was performed by defining for each set of abundances a \( \chi^2 \) function as the sum of the quadratic difference between synthetic and observed profiles for all wavelengths in the range.

Since there are three slightly different 3D models, resulting from using different Fe abundances (log \( \epsilon(Fe) = 7.40 \), 7.45 and 7.50) in the inversion of the Hinode Fe i profiles, we performed the comparison for all three cases, which resulted in an effective three-dimensional grid of abundances (log \( \epsilon(Fe) \), log \( \epsilon(O) \), and log \( \epsilon(Ni) \)). Even though the best value for the solar Fe abundance is probably log \( \epsilon(Fe) = 7.50 \), as concluded by previous works (Shchukina & Trujillo Bueno 2001), it is nevertheless interesting to use the three inversion results because this will hopefully provide some insight into how slight changes in the model can influence the inferred abundances.

The profiles plotted in Fig. 6 represent the best fit. The corresponding triplet of abundance values is log \( \epsilon(Fe) = 7.50 \), log \( \epsilon(O) = 8.90 \), and log \( \epsilon(Ni) = 6.15 \). As a side note, the Sc abundance required to fit this line is the meteoritic value of 3.05.

To illustrate how the fit degrades away from the optimal set of abundances, Fig. 7 shows the second-best fit in the grid. This new fit was obtained with log \( \epsilon(Fe) = 8.95 \) and log \( \epsilon(Ni) = 6.10 \), and the resulting \( \chi^2 \) is only 20% higher than before. The figure shows that the fit is remarkably tight, even though the difference in O abundance is rather significant. A comparison of the results obtained with all three models (corresponding to log \( \epsilon(Fe) = 7.40 \), 7.45 and 7.50) shows that the best fit is always achieved with log \( \epsilon(O) = 8.90 \) and log \( \epsilon(Ni) \) varying between 6.10 and 6.15, in all cases with similarly good fits to those shown in Fig. 6. If we also include the second-best match in the comparison, we have three more values to consider. As before, the \( \chi^2 \) is only slightly worse than for the best (30% higher, at the most) and the fits look visually very similar to those in Fig. 6. In this case, the spread of good fit abundances increases to the range from 8.85 to 8.95 for log \( \epsilon(O) \) and remains between 6.10 and 6.15 for log \( \epsilon(Ni) \).

To further explore the sensitivity of the abundance results to the model employed for the calculation, the same procedure was applied to a slightly perturbed version of the log \( \epsilon(Fe) = 7.50 \) model. The perturbation consists of a linear addition to the temperature in the range of optical depths log(\( \tau \)) = (-1,1) and has a total amplitude of 50 K, going from +50 K at log(\( \tau \)) = 1 down to -50 K at log(\( \tau \)) = -1. With such a small perturbation, the inferred abundances of O and Ni change to 8.78 and 6.02, respectively (Fig. 8).

Other factors that affect the result are the continuum reference and the wavelength calibration. If the continuum is chosen slightly lower or higher, both O and Ni abundances will change accordingly. This is a common problem in all abundance determinations. As an example, if we take the analysis shown in Fig. 6 and lower the continuum reference by only 0.1%, the best-fit profile in the grid is now the one corresponding to log \( \epsilon(O) = 8.83 \) and log \( \epsilon(Ni) = 6.20 \).

Similarly, a slight wavelength shift of a few mÅ might be compensated for by altering the ratio of O and Ni in the blend to recover a good fit. Finally, the line broadening produced by the dynamics in the lower photosphere has a significant impact as well. The AGSAK dynamics is incompatible with the Sc ii line, but still, even slight changes in the adopted rms velocity will affect the result.

If these factors are varied, it is even possible to achieve an O abundance compatible with that of AGSAK. The actual fit is shown in Fig 9. The fit is still remarkably good, given how
discrepant these new abundances are compared to the previous figures, which again illustrates how extremely sensitive these results are to small details of the model or of the procedure in general. To produce this low abundance, the velocity enhancement at the base of the photosphere was reduced by 15% (bringing the $\text{rms}$ velocity at that layer down from 1.36 to 1.15 km s$^{-1}$), the wavelength scale was shifted by 2 mÅ, and the continuum reference was decreased by 10%.

Fig. 8. Best fit abundances using a slightly perturbed model with a perturbation in temperature ranging from 50 K at $\log(\tau_{500})=1$ to -50 K at $\log(\tau_{500})=-1$.

Fig. 9. Fits with a low O abundance obtained after minor tweaks to the model atmosphere and the continuum reference selection. The goodness of this fit illustrates the high sensitivity of abundance determinations to the particular details of the model or the procedure.

5. Discussion

This paper presents a novel approach to the O abundance determination using a 3D model that has been obtained from observations, as opposed to most recent works, which used a hydrodynamical simulation. Since the model is obtained by inverting the much stronger Fe I lines at 6301.5 and 6302.5 Å observed by the Hinode SP, it seems to be missing some of the dynamics that occur in the deeper formation region of the O/Ni blend. This could be due to a lack of spatial resolution or to the weak sensitivity of the stronger Fe I lines to plasma motions in a layer very close to the continuum formation region. Ayres (2008) indicated that this might be a problem and noted, with remarkable perspicacity, that “a possible disadvantage is that the derived motions will depend on the instrumental resolution and might not be as vigorous as in fully resolved 3D theoretical models”. Fortunately, this problem can be overcome by using the nearby Sc II line, which is of very similar strength to the O/Ni blend and probes the same atmospheric region.

In principle, the results obtained are in the high range ($\log (\epsilon(O))=8.9\pm0.1$), compatible with the “old” solar composition. However, in view of how sensitive the results are to the details of the model atmosphere (it is even possible to produce values compatible with the “new” composition with very minor tweaks), they should probably be taken with some reservation. In fact, the same caution should probably be adopted with all previous results on abundance determinations based on the traditional technique of fitting atlas observations with a prescribed model atmosphere. Ultimately, the reliability of our abundances is limited by the uncertainty in the models employed, of which we still lack such detailed understanding. It is then almost impossible to ascribe meaningful error bars to those measurements. A thorough study of the systematic errors arising from the use of a prescribed model and other parameters employed in the abundance analysis is urgently needed to gain a better understanding of our limitations.

An important conclusion to draw from this study is that the “low” O abundance of AGSAK ($\log (\epsilon(O))=8.66$) is not necessarily obtained from a 3D model with recent atomic data. Intermediate (Caffau et al. 2008) and high (this work) abundances may be obtained from 3D models as well. Furthermore, given the sensitivity of the results to the details of the analysis, we should also question to what degree an analysis based on equivalent widths (which is what has been employed in most cases thus far) is enough to capture all the subtleties that are observed in a detailed fit. It seems that in difficult situations, such as this one, it is necessary to work with a full line analysis quantified by some merit function (e.g., the $\chi^2$ used here).

In the solar case there is an obvious way to improve the overall procedure and increase our confidence in the measurement of element abundances. Using spatially resolved observations, it is possible to conduct a pixel-by-pixel analysis, similarly to what was reported in Socas-Navarro & Norton (2007). If the systematic errors are known, there is no reason to expect that all of the analyzed spectra will yield consistent results. Therefore, the pixel-to-pixel distribution is a perfect sanity check to diagnose possible systematic errors. In that paper we found a fluctuating pattern of abundances, correlated with the granulation distribution. This inconsistency was probably a reflection of the uncertainties in the non-LTE line formation physics. Conducting a similar study with simpler lines such as the forbidden transition analyzed here would undoubtedly cast some more light into this important question.

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The VALD database (referenced in the text) was used to obtain atomic parameters for the Sc ii line.

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Table 1. Spectral line data, where \( r_0 \) is the Böhr radius, \( \gamma_{\text{rad}} \), \( \gamma_{\text{Stark}} \), and \( \gamma_{\text{vdW}} \) are the radiative, Stark and van der Waals damping enhancement constants.

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<th>Element</th>
<th>( \lambda (\text{Å}) )</th>
<th>Excitation(eV)</th>
<th>( \log(g) )</th>
<th>( \sigma(r_0^2) )</th>
<th>( \alpha )</th>
<th>( \gamma_{\text{rad}} )</th>
<th>( \gamma_{\text{Stark}} )</th>
<th>( \gamma_{\text{vdW}} )</th>
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Table 2. Inversion nodes

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