The AMBRE Project: Stellar parameterisation of the ESO:FEROS archived spectra

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ABSTRACT

Context. The AMBRE Project is a collaboration between the European Southern Observatory (ESO) and the Observatoire de la Côte d’Azur (OCA) that has been established in order to carry out the determination of stellar atmospheric parameters for the archived spectra of four ESO spectrographs.

Aims. The analysis of the FEROS archived spectra for their stellar parameters (effective temperatures, surface gravities, global metallicities, alpha element to iron ratios and radial velocities) has been completed in the first phase of the AMBRE Project. From the complete ESO:FEROS archive dataset that was received, a total of 21 551 scientific spectra have been identified, covering the period 2005 to 2010. These spectra correspond to 6285 stars.

Methods. The determination of the stellar parameters was carried out using the stellar parameterisation algorithm, MATISSE (MATrix Inversion for Spectral SynthEsis), which has been developed at OCA to be used in the analysis of large scale spectroscopic studies in galactic archaeology. An analysis pipeline has been constructed that integrates spectral normalisation, cleaning and radial velocity correction procedures in order that the FEROS spectra could be analysed automatically with MATISSE to obtain the stellar parameters.

The synthetic grid against which the MATISSE analysis is carried out is currently constrained to parameters of FGKM stars only.

Results. Stellar atmospheric parameters, effective temperature, surface gravity, metallicity and alpha element abundances, were determined for 6508 (30.2%) of the FEROS archived spectra (~3087 stars). Radial velocities were determined for 11 963 (56%) of the archived spectra. 2370 (11%) spectra could not be analysed within the pipeline due to very low signal-to-noise ratios or missing spectral orders. 12 673 spectra (58.8%) were analysed in the pipeline but their parameters were discarded based on quality criteria and error analysis determined within the automated process. The majority of these rejected spectra were found to have broad spectral features, as probed both by the direct measurement of the features and cross-correlation function breadths, indicating that they may be hot and/or fast rotating stars, which are not considered within the adopted reference synthetic spectra grid. The current configuration of the synthetic spectra grid is devoted to slow-rotating FGKM stars. Hence non-standard spectra (binaries, chemically peculiar stars etc.) that could not be identified may pollute the analysis.

Key words. astronomical databases: miscellaneous – stars: fundamental parameters – techniques: spectroscopic – methods: data analysis

1. Introduction

Astronomy has entered an era of large scale astronomical surveys, the scientific goals of which have the potential to considerably expand our understanding of the formation and evolution of the Universe. In particular current and future large scale spectroscopic surveys of the Milky Way will allow astronomers to trace in incredible detail the chemical and kinematic history of our Galaxy.

These surveys are being undertaken over a range of resolutions. Low-resolution surveys ($R = 2000$ to $8000$) are now widespread, for example RAVE (Steinmetz et al. 2006) and SEGUE (Yanny et al. 2009), while high-resolution surveys are a more recent endeavour such as APOGEE (Majewski et al. 2007) and the Gaia-ESO Survey (P.I.s: Gerry Gilmore & Sofia Randich). The science goals of these ground-based surveys will significantly contribute to the studies of galactic archaeology as well as provide complementary information to the upcoming astrometric survey, the European Space Agency (ESA) Gaia Mission. The ESA Gaia satellite will observe approximately a billion stars in the Milky Way for which distances will be determined to milliarcsecond accuracies. Its Radial Velocity Spectrometer (RVS) will observe spectra at medium-resolution ($R \approx 7000–11\,500$) which will be used to obtain radial velocities for all the targets, as well as to determine stellar atmospheric parameters and chemical abundances for some ten’s of millions of stars.

The multi-object instruments at high-resolution ($R \sim 20\,000$; MIKE on Magellan and FLAMES on the Very Large Telescope), medium-resolution ($R \sim 8000$; AAOmega on the Australian Astronomical Telescope (AAT)) and low-resolution ($R \leq 2200$; Fibre Multi Object Spectrograph (FMOS) on the Subaru Telescope) are also being used to address science goals specific to the field of galactic archaeology. One key instrument that is currently being built and has been designed primarily for galactic archaeology research is the High Efficiency and Resolution Multi-Element Spectrograph (HERMES Barden et al. 2010) on the AAT.

This ongoing accumulation of large spectral datasets from surveys and individual spectrographs has compelled the development of automated stellar parameterisation algorithms that can reliably and effectively analyse every spectrum. The stellar parameterisation algorithm, MATISSE, has been developed at the Observatoire de la Côte d’Azur (OCA).
(Recio-Blanco et al. 2006) to be included in the automated pipeline for the analysis and parameterisation of the Gaia-RVS stellar spectra. MATISSE has also been developed as a standalone Java application for use in a wide variety of projects (see for instance, Gazzano et al. 2010; Kordopatis et al. 2011). The AMBRE Project team at OCA, which oversees the development of MATISSE, is connected to the Gaia Data Processing Consortium (DPAC) under the Generalized Stellar Parameterizer-spectroscopy (GSP-spec) Top Level Work Package which is overseen by Coordination Unit 8 (CU8).

With such large datasets soon to be available a crucial aspect of the analysis is to be able to compare the results from the different surveys. To do this a comprehensive set of standard objects is required that can be used to calibrate the datasets. Preparation for this is in the form of the development of spectral libraries. These are datasets of spectra with homogeneously determined characteristics, such as stellar parameters, which can be used as calibration stars for these surveys. In particular standard star lists are being developed for Gaia for both the radial velocity (Crifo et al. 2010) and stellar parameter measurements (Soubiran et al. 2010).

The work carried out for the AMBRE Project in effect converts the spectra in the European Southern Observatory (ESO) archive into a comprehensive spectral library of homogeneously determined stellar parameters. There are three primary objectives for the AMBRE Project:

1. To rigorously test MATISSE on large spectral datasets over a range of wavelengths and resolutions, including those for the Gaia RVS.
2. To provide ESO with a database of stellar temperatures, gravities, metallicities, alpha to iron ratios and radial velocities for the associated archived spectra that will then be made available to the international scientific community via the ESO Archive.
3. To create a chemical map of the Galaxy from the combined ESO archived sample upon which stellar and galactic formation and evolution archaeological analysis can be carried out.

The first phase of the AMBRE Project was the analysis of the FEROS archived spectra. From the complete ESO:FEROS archive dataset that was delivered to OCA, a total of 21,551 scientific spectra have been identified, covering the period 2005 to 2010. These spectra correspond to 6285 different stars based on a coordinate matching calculation with a radius of 10″.

The structure of this paper is as follows: Sect. 2 introduces the AMBRE Project; Sect. 3 introduces the MATISSE algorithm and the synthetic spectra grid; Sect. 4 describes the analysis pipeline that has been built around MATISSE for the analysis of the archived spectra; Sect. 5 discusses the internal errors that have been calculated for the pipeline; Sect. 6 discusses the external errors analysis based on a reference sample of stars; Sect. 7 presents the application of key rejection criteria to the spectral dataset; Sect. 8 presents the stellar parameter results for FEROS and finally Sect. 9 concludes the paper.

### 2. The AMBRE Project

Under a contract with ESO the archived spectra of four ESO spectrographs are being analysed using the automated parameterisation programme MATISSE in a project overseen by the AMBRE Project team. Table 1 lists the main characteristics of the four spectrographs: the Fiberfed Extended Range Optical Spectrograph (FEROS) (Kaufer et al. 1999); the High Accuracy Radial velocity Planet Searcher (HARPS) (Mayor et al. 2003); the Ultraviolet and Visual Echelle Spectrograph (Dekker et al. 2000); and the Fibre Large Array Multi Element Spectrograph/GIRAFFE (FLAMES/GIRAFFE) (Pasquini et al. 2002). The atmospheric stellar parameters of effective temperature ($T_{\text{eff}}$), surface gravity (log $g$), global metallicity ($[\text{M/H}]$), $\alpha$ element to iron ratio ([α/Fe]) and radial velocity ($V_{\text{rad}}$) will be derived for each of the archived stellar spectra. These will be delivered to ESO for inclusion in the ESO database and then made available to the astronomical community via the ESO archive. It is intended that the availability of stellar parameters for each spectra will encourage further use of the archived spectra. Previously unconsidered samples can be found through searches on the parameters, for example extracting all the spectra in a particular metallicity range, or with very similar temperatures and gravities.

The analysis of the archived spectra of these four spectrographs presents a unique opportunity to test the performance of MATISSE on large datasets of real spectra. In particular key instrument configurations of FLAMES/GIRAFFE cover the Gaia RVS wavelength domain and resolutions. Rigorous testing of MATISSE is necessary in order to optimise its performance in the Gaia-RVS analysis pipeline that is being compiled at the Centre National d’Etudes Spatiales (CNES). As such the AMBRE Project has been formally designated as a sub-work package under GSP-spec. The stars analysed by AMBRE will also be available for use as standard or calibration stars for the Gaia-ESO survey, and as secondary standards for the Gaia Mission.

Figure 1 shows a histogram of the number of FEROS observations per year that were received by OCA, the number for 2005 representing only 3 months of observations. The number counts for the number of distinct objects that have been observed

### Table 1. Details of the four ESO spectrographs and their publicly available archived spectra sample that are part of the AMBRE Project.

| Spectrograph | Resolution | $\lambda$ (nm) | No. spectra |
|--------------|------------|----------------|-------------|
| FEROS        | 48 000     | 350–920        | 21 551      |
| HARPS        | 115 000    | 378–690        | 126 694     |
| UES          | 20 000–110 000 | 300–1100      | 78 593      |
| FLAMES/GIRAFFE | 5 600–46 000 | 370–900        | > 100 000   |

1 http://www.lsw.uni-heidelberg.de/projects/instrumentation/Feros/
2 http://www.eso.org/sci/facilities/lasilla/instruments/feros/index.html
by FEROS in this timeframe and the number with repeated observations are as indicated in the key.

The analysis procedure and results for these FEROS spectra are presented in this paper. This analysis has been the testbed for producing many of the tools that will also be used in the analysis of the UVES, HARPS and FLAMES/GIRAFFE archived spectra. These tools have been integrated into an analysis pipeline that feeds the processed spectra into MATISSE for derivation of the spectral parameters.

3. MATISSE and the synthetic spectra grid

MATISSE (MATrix Inversion for Spectral SynthEsis) is an automated stellar parameterisation algorithm based on a local multilinear regression method. It derives stellar parameters ($\theta = T_{\text{eff}}, \log g, [\text{M/H}], \text{individual chemical abundances}$) by the projection of an input observed spectrum on a vector function $B_\theta(\lambda)$,

$$\hat{\theta} = \sum_{i} B_\theta(\lambda) \cdot O_i(\lambda)$$

with $\hat{\theta}$ being the derived value. The $B_\theta(\lambda)$ vector function is an optimal linear combination of theoretical spectra, $S_j(\lambda)$, calculated from a synthetic spectra grid.

$$B_\theta(\lambda) = \sum_{j} \alpha_{ij} S_j(\lambda)$$

Key features in the observed spectrum due to a particular $\theta$ are reflected in the corresponding $B_\theta(\lambda)$ vector indicating the particular regions which are sensitive to $\theta$ (Recio-Blanco et al. 2006; Bijaoui et al. 2008).

MATISSE also generates synthetic spectra interpolated to each set of output stellar parameters. For each set of parameters, the synthetic spectrum on the grid, $S_\theta(\lambda)$, with the closest stellar parameters is identified along with the associated $B_\theta(\lambda)$ function. The $B_\theta(\lambda)$ functions located locally about this point are used to estimate the variations in the flux between the synthetic spectra at the corresponding grid points and the final interpolated synthetic spectrum corresponding to the required stellar parameters ($\hat{\theta}_k$) (Kordopatis et al. 2011).

This interpolated spectrum is then used to calculate a $\chi^2$ between the interpolated and input normalised spectrum. This provides a measure of the goodness of the fit of the derived stellar parameters to the observed spectrum. Stellar parameters, with $\log \chi^2$, and corresponding interpolated synthetic spectra are generated for every spectrum that is analysed in MATISSE.

3.1. Grid of synthetic spectra for the classification algorithm

Within the AMBRE Project, the adopted procedure for the automatic classification of stellar spectra relies on a library of reference spectra in order to derive their atmospheric parameters and their chemical abundances. Due to the lack of a library of observed spectra that covers a large range of atmospheric parameters and chemical abundances over a very large spectral domain and resolution, the only solution was to compute large grids of synthetic spectra.

For the AMBRE application, the computed grid of theoretical stellar spectra has to cover the whole optical spectral range at very high resolution in order to be used for the analysis of the majority of the spectroscopic data. For that reason, we have computed a synthetic spectra grid covering the wavelength range between 300 and 1200 nm with a wavelength step of 0.001 nm (900 000 pixels in total) (de Laverny et al. 2012). Since this project is mostly devoted to the analysis of FGKM stars, this grid is based on the latest generation of MARCS model atmospheres presented in Gustafsson et al. (2008). An extension to hotter effective temperatures with Kurucz stellar atmosphere models (Kurucz 1979) is planned for the near future. The considered parameters of the spectral grid are the effective temperature ($T_{\text{eff}}$ in K), the stellar surface gravity ($\log g$ in dex), the mean metallicity ([M/H] in dex) and the enrichment in $\alpha$-elements with respect to iron ([Fe/H] in dex).

The metallicity definition here is different to the classical use of [Fe/H] to denote the metallicity of a star. For [Fe/H] the metallicity is defined using the derived abundances from Fe lines only. MATISSE provides the opportunity to use all the available metal lines (those atoms heavier than He) to define a metallicity providing a global metallicity designated as [M/H]. The opportunity also exists with MATISSE to derive a global $\alpha$ element abundance using as many $\alpha$ element spectral lines as possible, where we assumed that in the generation of the synthetic spectra the abundances of the different $\alpha$ elements vary in lockstep. Throughout the AMBRE Project, the following chemical species are assumed to be $\alpha$-elements: O, Ne, Mg, Si, S, Ar, Ca and Ti, although for any selected wavelength region spectral features for all of these elements may not necessarily be present.

From the selected MARCS model atmospheres, the synthetic spectra were computed with the turbospectrum code (Alvarez & Plez 1998, and further improvements by Plez) in plane-parallel and spherical geometry assuming hydrostatic and local thermodynamic equilibrium. Atomic lines have been recovered from the Vienna Atomic Line Database (in August 2009; Kupka et al. 1999). The molecular line list has been provided by B. Plez. It includes transitions from ZrO, TiO, VO, OH, CN, C2, CH, SiH, CaH, MgH and FeH with their corresponding isotopic variations (see Gustafsson et al. 2008, for a list of references).

This grid covers the following ranges of atmospheric parameters: $T_{\text{eff}}$ between 2500 K and 8000 K, $\log g$ from −0.5 to +5.5 dex, and [Fe/H] from −5.0 to +1.0 dex, although not all combinations of the parameters are available within the grid. The selected MARCS models have $[\alpha/\text{Fe}] = 0.0$ for $[\text{M/H}] \geq 0.0$, $[\alpha/\text{Fe}] = +0.4$ for $[\text{M/H}] \leq −1.0$ and, in between, $[\alpha/\text{Fe}] = −0.25 \times [\text{M/H}]$. For the spectra computation from each
of these MARCS models, we considered an [$\alpha$/Fe] enrichment from $-0.4$ to $+0.4$ dex with respect to the canonical values that correspond to the original abundances of the MARCS models. The final AMBRE synthetic spectra grid consists of 16 783 flux normalized spectra (see de Laverny et al. 2012, for a complete description of this grid).

The microturbulence ($\xi$) is not a free parameter in this synthetic spectra grid. For the atmospheric models with high log $g$ ($+3.5 \leq \log g \leq +5.5$) $\xi$ is set at 1.0 km s$^{-1}$. For low log $g$ ($\log g < +3.0$) $\xi$ is set at 2.0 km s$^{-1}$, being typical values for dwarfs and giants respectively. These values reflect the MARCS model atmospheres configurations, and further details on the model selection is given in de Laverny et al. (2012). Due to the microturbulence being hardwired into the synthetic grid it was not possible to carry out tests on the effects of variations in $\xi$ on the stellar parameter determination across the whole synthetic spectra grid. However the effects of changes in $\xi$ are most prominent for strong lines. By using the global metallicity [M/H] rather than [Fe/H] we expect the contribution from strong lines on the [M/H] to be negligible due to the significantly larger quantity of non-$\xi$ sensitive small metallic lines.

Section 6 describes the extensive testing of the pipeline that was carried out using observed spectra from which an external error on the resulting parameters is derived by comparison to literature values. To gauge the effect of microturbulence on the derived stellar parameters a key sample of 66 Main Sequence (MS) stars (Bensby et al. 2003), and two key samples of 16 Red Giant Branch (RGB) stars (McWilliam 1990; Hekker & Meléndez 2007) were considered from within the test sample.

The most noticeable effect within the MS sample was an underestimation of the $T_{\text{eff}}$ for the hotter MS stars, for which the constant $\xi$ of 1.0 km s$^{-1}$ underestimates the accepted value. Hence at $T_{\text{eff}} \sim 6500$ K, where $\Delta T_{\text{eff}} \sim +0.8$ km s$^{-1}$ the effect was $\Delta T_{\text{eff}} \sim -100$ K. This is well within the derived $T_{\text{eff}}$ external error of 120 K (see Sect. 6).

The most noticeably effect in the RGB sample was on the derivation of log $g$. At the base of the RGB, where the constant $\xi$ of 2.0 km s$^{-1}$ is an overestimation, the derived log $g$ was observed to be overestimated by $-0.3$ dex for $\Delta g \sim +0.7$ km s$^{-1}$. At the RGB tip, where $\xi \sim 2.0$ km s$^{-1}$ is an underestimated, log $g$ was observed to be underestimated by $-0.3$ dex for $\Delta g \sim -0.7$ km s$^{-1}$. These variations are within the derived log $g$ external error of 0.37 dex. Hence based on these samples, variations of $\xi$ from the assumed constant values do have an effect on the derived parameters that varies in magnitude with stellar evolutionary stage. However the external error derived from the entire reference sample takes account of them in a global sense (see Sect. 6).

3.2. AMBRE:FEROS subgrid

The FEROS spectra cover most of the optical wavelengths but only a subset of these wavelengths was required for the MATISSE analysis. In order to carry out the MATISSE training phase, where the $B_0(\lambda)$ vectors are generated, for the FEROS analysis the optimum resolution, wavelength regions, and sampling of the FEROS spectra were determined. The full AMBRE synthetic spectra grid was then adapted to the same specifications creating the AMBRE:FEROS $B_0(\lambda)$ vectors.

In analysis, only $B_0(\lambda)$ functions computed from the direct numerical inversion of the correlation matrix (see Recio-Blanco et al. 2006) were considered. These functions are thus not optimized for very low SNR spectra. However, due to the large number of spectral features available in the selected spectral domains, it was not necessary to use the approximated $B_0(\lambda)$ functions that are computed with the Landweber algorithm as in Kordopatis et al. (2011). As will be shown in Sect. 5 the estimated internal errors based on the AMBRE:FEROS grid are indeed already very small (see Fig. 10).

3.2.1. Selection of wavelength regions

A detailed analysis of the full wavelength range of the FEROS spectra was undertaken in order to select wavelength regions that would provide the greatest amount of information for each of the derived parameters while also minimising the number of pixels in order to reduce computing time.

FEROS disperses the spectra into 39 orders of varying wavelength interval from $\sim3500$ Å to $\sim9200$ Å at high resolution ($R \sim 48000$). The optimum spectral regions for the determination of the stellar parameters were selected taking into account the need to avoid excessive computing time and to avoid low spectral information region. To this end, the signal-to-noise (SNR) per pixel ($0.03$ or $0.06$ Å per pixel for FEROS) as a function of the wavelength was derived for each archived spectra by determining the SNR profile of each spectral order and thus providing a SNR profile over the entire spectral domain. Wavelengths regions with low SNR were rejected, typically the start and end of each order. The wavelengths regions affected by sky absorption and telluric features were also rejected, and also regions where continuum placement proved too difficult owing to wide spectral features in the region. Some key spectral features, such as H$_\alpha$ and the Ca II H & K lines, were also discarded as these features were found to be poorly synthesised based on our current understanding of stellar atmospheres as well as being difficult to normalise automatically.

The remaining spectral regions were then investigated for their intrinsic sensitivity to three of the four stellar parameters to be determined by MATISSE: $T_{\text{eff}}$, log $g$, and [M/H]. This was carried out using preliminary $B_0(\lambda)$ vectors that covered the full optical domain at very high resolution generated using a reduced synthetic grid. Due to the underlying equation for calculating the $B_0(\lambda)$ vectors, the high resolution $B_0(\lambda)$ vectors could not be simply degraded to lower resolutions to investigate the sensitivity, as this would not accurately reflect the information at that resolution. However the high resolution $B_0(\lambda)$ vectors are useful to give a general sense of the sensitivity.

For stellar parameters at:

1. $T_{\text{eff}} = 6000$ K, log $g = 4.0$ dex, [M/H] = 0.0 dex (Sun);
2. $T_{\text{eff}} = 6000$ K, log $g = 2.0$ dex, [M/H] = 0.0 dex;
3. $T_{\text{eff}} = 4500$ K, log $g = 2.0$ dex, [M/H] = 0.5 dex (Arcturus);
4. $T_{\text{eff}} = 4500$ K, log $g = 4.0$ dex, [M/H] = 0.5 dex,
the $B_0(\lambda)$ vectors at the same $T_{\text{eff}}$, log $g$, and [M/H] have been analysed for their extrema distribution. This selection explores the sensitivity of both dwarfs and giants that are similar to the two standard stars, the Sun and Arcturus. The panels in Fig. 2 show how the spectral domain changes in sensitivity with the different stellar parameters. The standard deviation ($\sigma$) for each $B_0(\lambda)$ vector was calculated as the spread of the pixel values in the $B_0(\lambda)$ vector. Then for each $B_0(\lambda)$ vector the pixels outside 2$d\sigma$ were binned in $\sim200$ Å bins over the wavelength range from $\sim4000$ Å to $\sim9500$ Å. Hence the bars in Fig. 2 show the number of pixels per bin which have a high sensitivity to the respective $\theta$. Clearly the bluer wavelengths show the greatest
sensitivity to all three parameters, reflecting the greater quantity of spectral features in the blue.

This sensitivity to \( \theta \) for different ranges in \( \theta \) was used to select the wavelengths regions in the FEROS spectra to be used in the AMBRE analysis. As the greatest sensitivity was located towards the blue, regions were selected, where possible, to capture this sensitivity. The grey regions in Fig. 2 represent the final wavelength regions selected for the AMBRE:FEROS analysis (see Table 2).

Fig. 3. As for Fig. 2 but for the corresponding \( B_\theta(\lambda) \) vectors of the AMBRE:FEROS synthetic spectra grid. The y-axis is scaled down compared to Fig. 2.

Table 2. FEROS échelle order, starting wavelength and finishing wavelength for each region used in the analysis of the FEROS spectra.

| Region | \( \lambda \) min | \( \lambda \) max | Region | \( \lambda \) min | \( \lambda \) max |
|--------|-----------------|-----------------|--------|-----------------|-----------------|
| 1      | 6660 Å          | 6730 Å         | 10     | 4585 Å          | 4670 Å         |
| 2      | 5655 Å          | 5725 Å         | 11     | 4505 Å          | 4580 Å         |
| 3      | 5390 Å          | 5430 Å         | 12     | 4405 Å          | 4492 Å         |
| 4      | 5260 Å          | 5320 Å         | 13     | 4300 Å          | 4405 Å         |
| 5      | 5120 Å          | 5190 Å         | 14     | 4245 Å          | 4300 Å         |
| 6      | 4990 Å          | 5055 Å         | 15     | 4140 Å          | 4220 Å         |
| 7      | 4910 Å          | 4960 Å         | 16     | 4075 Å          | 4140 Å         |
| 8      | 4800 Å          | 4890 Å         | 17     | 4019 Å          | 4075 Å         |
| 9      | 4674 Å          | 4740 Å         |        |                  |                 |

Table 3 reflects the sensitivity of the \( B_\theta(\lambda) \) vectors at the actual resolution of the AMBRE:FEROS analysis. Due to the reduced number of points in sampling and resolution the 2\( \sigma \) limit is significantly reduced. While some regions show there are subtle differences in the sensitivity of some regions, overall the distribution of greater sensitivity in the blue is in good agreement with the sensitivity at high resolution.

3.2.2. Convolution and sampling of synthetic spectra grid

At the high resolution of \( R \sim 48000 \), the FEROS spectra have typical wavelength sampling (\( \Delta \lambda \)) of 0.03 Å (1 \times 1 binning) or 0.06 Å (1 \times 2 binning). The final selected wavelengths corresponded to a total of \( \sim 1500 \) Å and at this sampling this translated to a (maximum) total of \( \sim 50000 \) pixels. Creating the FEROS synthetic spectra subgrid of \( \sim 16000 \) spectra at this resolution and sampling would result in excessively large memory and computing requirements when creating the \( B_\theta(\lambda) \) vectors in the training phase, and also when loading these vectors during the MATISSE analysis. It was necessary to reduce the resolution and sampling to a more reasonable pixel total, but without sacrificing the key spectral information. The most reasonable number of pixels per spectra was determined to be \( \sim 15000 \) in order to ease the use of the available computing power.

By optimising the memory and spectral information requirements, the final specifications for the FEROS synthetic spectra subgrid was determined to be 11 890 pixels with \( R \sim 15000 \) at \( \lambda \sim 4500 \) Å with a sampling of 0.1 Å over \( \sim 15000 \) Å. Hence the full AMBRE synthetic spectra grid (de Laverny et al. 2012) was then sliced to the wavelengths specified in Table 2, convolved by a Gaussian kernel with constant \( \sigma \) and resampled to the optimised FEROS pixels. The training phase was then carried out whereby the \( B_\theta(\lambda) \) vectors were generated for use in the MATISSE analysis of the FEROS archived spectra.

4. FEROS analysis pipeline

A complex analysis pipeline (written in shell, Java and IDL) has been built to wavelength slice, radial velocity correct, normalise and convolve the FEROS spectra and then feed them into MATISSE for the determination of their stellar parameters. Figure 4 shows a flowchart of the key stages in the analysis pipeline. The pipeline has been developed so that it can be easily adapted to the remaining three instrument datasets of ESO archived spectra. The following sections outline the procedures carried out for each of the key stages in the pipeline.
4.  Spectral Processing A: spectra processing for radial velocity determination

Spectral Processing A (SPA) is the initial stage that prepares the FEROS spectra in order to determine the radial velocities. The procedures used here are also used in the later stages of the pipeline. Key flags were defined within the pipeline to be attached to any spectra that satisfied the conditions of the flag. The flags were for “Faulty Spectra” (e.g. missing spectral orders), “Extreme Emission Features” (spectral emission features in the blue with width greater than 50 Å), “Excessive Noise” (extremely noisy spectra with the number of negative flux pixels >25%) and “Poor Normalisation” (spectra with a total flux less than 1.2 Å (40 pixels) are cleaned. This limit of 1.2 Å (40 pixels) in width are flagged as large emission features and are not cleaned. This limit of 1.2 Å (40 pixels) was determined by an examination of many examples of cosmic rays in the FEROS spectra to get a sense of the number of pixels over which they could be dispersed.

Each wavelength region is treated individually for each spectrum, and the regions are then combined into one vector. This merged spectrum is tested for the goodness of the normalisation and flagged if the normalisation is poor. The resulting spectrum is used in the determination of the radial velocity. At this stage the original resolution is retained. None of the FEROS spectra were rejected at this stage in case later testing and processing recovered a previously rejected spectra (i.e. mis-calculated radial velocity is corrected).

4.2. Radial velocity determination

A completely automatic radial velocity ($V_{\text{rad}}$) programme has been established that can analyse spectra across a very wide range of stellar parameters and is based on a cross-correlation algorithm that compares the normalised observed spectrum to binary masks (C. Melo, priv. comm.; Melo et al. 2001). To complement this programme a procedure was developed at OCA that computes a set of binary masks to match the observations in terms of wavelength range and resolution, and are used as input to the radial velocity programme. Masks covering the parameter space of the AMBRE:FEROS analysis were computed using synthetic spectra taken from the high resolution, full optical domain, AMBRE synthetic spectra grid (see Sect. 3.1). Six masks for stars hotter that 8000 K were also computed using synthetic spectra from the POLLUX database (Palacios et al. 2010). For the AMBRE:FEROS analysis 51 masks were computed at the stellar parameters given in Table 3. Also listed are 5 standard masks for dwarf stars that were supplied with the radial velocity programme (C. Melo, priv. comm.) and used in the AMBRE:FEROS analysis.

For each spectrum a cross-correlation function (CCF) ranging from $-500$ to $+500$ km s$^{-1}$ was calculated for each of the 56 masks. The CCF was calculated in radial velocity steps ($\Delta V_{\text{rad}}$) based on the specified wavelength step ($\Delta \lambda$) of the spectrum. For spectra with $\Delta \lambda = 0.03$ Å the CCF step was calculated to be $\Delta V_{\text{rad}} \approx 1.8$ km s$^{-1}$, and for $\Delta \lambda = 0.06$ Å the CCF step was $\Delta V_{\text{rad}} \approx 3.6$ km s$^{-1}$. For each CCF two separate gaussian fits were made in order to determine the minimum of the profile, and hence the radial velocity. The first fit used the majority of the profile, while the second fit used a section of the profile centred near the minimum with a width equal to the full-width-at-half-maximum (FWHM) of the profile.

Figure 5a shows an example of a noisy CCF although the key profile is distinct. However, as seen in Fig. 5b, the fit to the full profile (black) provides a minimum skewed away from the actual minimum due to asymmetries of the full profile. The use of the central core of the profile provides a more accurate fit (red). Figure 5c shows an example of a CCF for which two profiles...
Table 3. List of the stellar parameters for the binary masks used in the AMBRE:FEROS radial velocity determination that were built from the full AMBRE synthetic grid (45 masks) and the POLLUX database (6 masks).

| Teff | log g | [M/H] | [α/Fe] | Teff | log g | [M/H] | [α/Fe] | Teff | log g | [M/H] | [α/Fe] |
|------|-------|-------|--------|------|-------|-------|--------|------|-------|-------|--------|
| 5000 | 0.5   | −2.00 | 0.40   | 6500 | 1.0   | 0.00  | 0.00   | 9000 | 4.5   | 0.00  | 0.00   |
| 2000 | 1.5   | −0.25 | 0.10   | 2000 | 2.5   | −2.00 | 0.40   | 10000| 4.5   | 0.00  | 0.00   |
| 5000 | 0.5   | 0.00  | 0.00   | 5750 | 2.5   | −2.00 | 0.40   | 12000| 4.5   | 0.00  | 0.00   |
| 4000 | 0.5   | 0.00  | 0.00   | 6500 | 1.0   | 0.00  | 0.00   | 15000| 4.5   | 0.00  | 0.00   |
| 5000 | 0.5   | 0.00  | 0.00   | 5750 | 2.5   | −2.00 | 0.40   | 30000| 3.2   | 0.00  | 0.00   |
| 4000 | 2.5   | 0.00  | 0.00   | 2500 | 2.5   | 0.00  | 0.00   | 37500| 3.5   | 0.00  | 0.00   |

Notes. Five standard masks for dwarf stars provided by C. Melo are also listed.

Fig. 5. CCFs produced for calculating radial velocities. a) Noisy CCF with an asymmetric profile. b) Core of profile showing skewed full profile gaussian fit (black). The secondary gaussian fit (red) provides a better estimate. c) CCF of the spectroscopic binary, HD 135728, showing two profiles.

Fig. 6. Examples of cross-correlation functions of the best fit masks for three of the FEROS spectra. The first row shows the full CCFs while the second row shows the profiles from which the radial velocities were determined. The final AMBRE:FEROS stellar parameters (Teff, log g, [M/H], [α/Fe]) and radial velocities are stated for each spectrum. The red lines are used to calculate the contrast and FWHM for each profile.

are observed. This is the spectroscopic binary HD 135728. The pipeline selects the most prominent of the two profiles in the determination of the radial velocity. The CCF seems the most likely tool with which to be able to identify the spectra of spectroscopic binaries. However at this stage we have not developed such a routine and so binaries are not specifically detected in the pipeline. The stellar parameters for such spectra are likely to be poorly determined and so will be at least identified as having large associated errors from the radial velocity and stellar parameter analysis. Quality control flags, such as the \( \chi^2 \), may also be used to help identify spectroscopic binaries.

Figure 6 shows the CCF corresponding to the best fit mask for three FEROS spectra at different stellar parameters. The full CCF for each spectrum is shown in the first row, while the second row shows more clearly the profile used to calculate the radial velocity. The full CCF can show varying degrees of noise and gradient depending on the quality of the spectra as is shown in each of these three examples. The respective profiles used to determine the radial velocity reflect the characteristics of the different spectra. The two cooler stars have very prominent inverted peaks, reflecting the numerous spectral lines available with which to make the cross-correlation. The hotter star, with fewer spectral lines, has a much less pronounced profile. These differences can be quantified by the measurement of the contrast of the profile.

The contrast was defined here as the amplitude of the profile multiplied by 100 (hence a percentage) where the amplitude of the profile is the length of the vertical red line as shown in each example profile in Fig. 6. The amplitude of the CCF, the continuum placement of the CCF and their associated errors are calculated automatically when using the GAUSSFIT routine in IDL. The sign of the amplitude does change depending on whether the CCF has an absorption or an emission profile. The expected profile in this analysis is an absorption profile.

The two conditions that were required for a profile to have been well fitted by a gaussian were first: the contrast must be less than zero ensuring an absorption profile; and second, the error on the amplitude must be less than 20% of the amplitude ensuring that the CCF profile dominates above the noise. CCFs that satisfied these conditions were considered to be well-defined. Finally the error on the radial velocity \( \sigma_{\text{Vrad}} \) was calculated using the prescription outlined in Tonry & Davis (1979).

This prescription makes use of the relative heights of the primary and secondary peaks in the CCF profile hence \( \sigma_{\text{Vrad}} \) reflects the goodness of the definition of the primary peak.
The radial velocity programme carried out the calculations for each of the 56 masks, resulting in 56 determinations of the radial velocity for each spectrum. The errors derived from the quality of the gaussian fit and $\sigma_{V_{\text{rad}}}$ were used to select the best fit radial velocity. Typically the majority of these determinations were in good agreement at the correct radial velocity. However for masks with parameters far from the true parameters of the star, a radial velocity would be determined that was incorrect but sometimes had sufficiently small errors such that it would be incorrectly selected as the best fit. If the smallest error was the only selection criteria. To avoid this a binning procedure was implemented as follows that discarded such outliers.

For a single spectrum, all of the $V_{\text{rad}}$ determinations with well-defined CCFs were binned by $V_{\text{rad}}$ into a maximum of 3 bins in correlation with the overall spread of the $V_{\text{rad}}$ values. The bin size was set in each instance by the difference in the maximum and minimum $V_{\text{rad}}$ values divided by the number of bins. The bin with the largest number of members was retained and the outlying bins were discarded. The $V_{\text{rad}}$ within accepted bin with the smallest $\sigma_{V_{\text{rad}}}$ was selected as the final $V_{\text{rad}}$. In the case of an equal number of members within 2 or more bins, the bin with the smallest spread in $V_{\text{rad}}$ values (so best agreement) was selected. Typically, most of the radial velocities for a spectrum were in good agreement, so the maximum bin was easily identified. Otherwise the radial velocities were highly dispersed, and this would be reflected in the $\sigma_{V_{\text{rad}}}$ of the final selected value.

The errors on the gaussian fit of the CCF and the $\sigma_{V_{\text{rad}}}$ values were used later in the analysis pipeline to identify problematic spectra and select alternate radial velocities where necessary, and also as quality criteria for the final set of results to be delivered to ESO.

The S4N (Allende Prieto et al. 2004) and Crifo et al. (2010) libraries provided good samples of stars with which to validate the radial velocities determined in the AMBRE:FEROS pipeline. The S4N library comprises of 118 F and G dwarf stars for which a detailed analysis was carried out using high resolution high SNR spectra. All four parameters ($T_{\text{eff}}$, log $g$, [Fe/H] and [$\alpha$/Fe]) as well as radial velocities are available for each of the S4N stars. Crifo et al. (2010) is the preliminary list of 1420 candidate standard stars to be used to calibrate the radial velocities obtained with Gaia RVS. Section 6 gives further discussion on the use of these libraries to validate the AMBRE:FEROS results.

Within the FEROS archived dataset 30 stars (338 spectra) were found that are also within the S4N library, and 183 stars (411 spectra) were found that are also within Crifo et al. (2010). Figures 7a and b compare the S4N and Crifo et al. (2010) radial velocities values with those determined for AMBRE:FEROS. The mean and standard deviation of the differences between the two sets for each sample are also shown. There is a small offset in both sets with the S4N having a greater offset of 1.8 ± 1.4 km s$^{-1}$ compared to Crifo et al. (2010) of 0.52 ± 0.35 km s$^{-1}$. The FEROS spectra were sampled at either 0.03 Å or 0.06 Å from the ESO:FEROS reduction pipeline corresponding to expected $V_{\text{rad}}$ accuracies of 1.6 km s$^{-1}$ and 3.3 km s$^{-1}$ respectively. Hence these offsets indicate very good agreement for both samples, particularly for the Crifo et al. (2010) values.

All of the FEROS spectra were analysed for their radial velocity. However for 4217 spectra (19.6%) poorly defined or non-standard CCFs lead to unreliable estimates of the radial velocity. This is most likely due to some peculiar or non-stellar properties of the spectra (i.e. nova). Of these 4217 spectra, 549 also failed the conditions of the rejection flags as defined in Sect. 4.1.

![Fig. 7. Comparison of AMBRE:FEROS radial velocities: a) S4N library for 29 stars (338 spectra); and b) Crifo et al. (2010) for 158 stars (318 spectra).](image1)

![Fig. 8. Histogram of the radial velocity values calculated for all of the FEROS archived spectra defined as good spectra with well-defined CCFs: a) $V_{\text{rad}}$; and b) FWHM of the $V_{\text{rad}}$ CCF.](image2)

Figure 8a is a histogram of the radial velocities calculated for the FEROS spectra defined as good quality with well-defined CCF (15,513 spectra). The majority of the spectra have radial velocities between −100 and +50 km s$^{-1}$. Figure 8b is a histogram of the FWHM of the CCF, calculated from the full CCF profile, for the same sample of spectra. It shows that the majority of spectra returned a FWHM of less than 50 km s$^{-1}$. The effects of the CCF on the selection of the final dataset will be discussed in Sect. 8.

4.3. Spectral Processing B: spectra reduction, convolution and first parameter estimation

Spectral Processing B (SPB) proceeds under two iterations through the normalisation procedure and the MATISSE analysis. These are necessary as key tests on the normalisation and radial velocity correction are carried out between the two iterations allowing alterations to be made that provide better first
estimates of the stellar parameters for the remaining pipeline procedures. For both iterations the original archived spectra are re-analysed using the same procedures as outlined in SPA in terms of wavelength selection, cosmic ray cleaning and normalisation. However in both iterations of SPB the measured \( V_{\text{rad}} \) correction is automatically applied to each spectrum prior to normalisation. This allows each spectrum to be shifted to the laboratory rest frame at which the synthetic grid, and so the \( B_{\lambda}(\lambda) \) vector functions, have been calculated.

The AMBRE:FEROS synthetic grid was set to a resolution lower than the resolution of the observed FEROS spectra. In order to convolve the observed spectra to the appropriate resolution three factors needed to be taken into account. First, the observed spectra are observed with constant resolving power, which corresponds to increasing FWHM of the spectral features with wavelength. Second, the synthetic spectra grid was convolved using a gaussian profile with constant FWHM hence the synthetic spectra have constant spectral FWHM with wavelength, not a constant resolution. Third, the construction of the full AMBRE synthetic spectra grid did not include astrophysical broadening such as those due to \( V \sin i \) and \( \xi \) (de Laverny et al. 2012). Hence a straight-forward convolution using the nominal instrument FWHM and the FWHM applied to obtain the AMBRE:FEROS synthetic spectra grid would result in convolved observed spectra that had increasing FWHM with wavelength in disagreement with the synthetic spectra. Also for stars with astrophysical broadening greater than the broadening due to the resolving power of the instrument the spectra would be too broad for the appropriate parameter range of synthetic spectra.

Therefore a procedure was implemented that measured the spectral FWHM for each of the 17 selected wavelength regions (see Table 2) for each spectrum and then applied an appropriate gaussian profile to smooth each wavelength region in order that the convolution resulted in observed spectra that had spectral FWHM as close as possible to the constant FWHM of the synthetic spectra grid.

The FWHM values were measured as part of SPB and this was found to significantly increase the processing time of this stage of the pipeline. Hence, due to it being primarily a testing phase, for the first iteration of SPB the observed spectra are convolved using predetermined “mean” spectral FWHM values that were determined for each of the 17 wavelength regions. These default spectral FWHM values were determined by a statistical analysis of a subsample of 384 of the FEROS archived spectra, whereby the FWHM for as many weak and medium strength spectral lines as possible were measured in each of the sample spectra.

The FWHM was measured by fitting a gaussian function to each spectral feature and calculating the width of the gaussian profile at the midpoint of the depth of the profile. For each spectrum the FWHM values increased with wavelength hence assuming one FWHM for the entire spectrum was not reasonable across the wavelength domain of the AMBRE:FEROS analysis. Hence a mean FWHM value for each of the 17 wavelength sections was determined and these were used in the convolution of the corresponding wavelength section. The convolution was carried out using a transformation gaussian profile for which the standard deviation (\( \sigma_X \)) was defined as:

\[
\sigma_X = \sqrt{\frac{\text{FWHM}_{\text{grid}}^2}{\sigma_{\text{FWHM}}} - \frac{\text{FWHM}_{\text{obs}}^2}{\sigma_{\text{FWHM}}} + 2.0 \ln(2.0)}
\]

\[
\sigma_{\text{FWHM}} = \sqrt{\frac{2.0 \ln(2.0)}{2}}
\]

where FWHM\(_{\text{obs}}\) is the default FWHM for the region, FWHM\(_{\text{grid}}\) = 0.33 mA and \( \sigma_{\text{FWHM}} \) transforms the FWHMs to \( \sigma_X \). Therefore each region of the observed spectrum was convolved such that the FWHM of the convolved spectrum matched the FWHM of the synthetic spectrum.

Part of the convolution process is to resample the observed spectra to the same wavelength bins as the FEROS synthetic spectra grid. Also for SPB, the same quality tests outlined in SPA are used but again no spectra were rejected based on these tests in this first iteration.

The convolved FEROS spectra were then analysed using MATISSE to obtain the first estimate of the stellar atmospheric parameters (\( T_{\text{eff}}, \log g, [\text{M/H}] \) and \([\alpha/\text{Fe}] \)). MATISSE also outputs synthetic spectra interpolated to the derived stellar parameters. Within the MATISSE algorithm the determination of the stellar parameters undergoes 10 iterations during which the algorithm converges on the final parameters for each spectrum. This number is set empirically and the majority of spectra converge to their parameters in much less than 10 iterations. After the MATISSE analysis is complete each spectrum is tested for whether convergence on the final \( B_{\lambda}(\lambda) \) function occurred within the 10 iterations and also the spectrum is tested as to the goodness of the fit between the normalised and the interpolated synthetic spectra in a \( \log(\chi^2) \) calculation. In the case of non-convergence within MATISSE, the parameters determined at the ninth and tenth iterations are compared and the solution with the lowest \( \log(\chi^2) \) is selected as the final set of parameters. If there is no convergence and/or a high \( \log(\chi^2) \), the spectra are flagged for further investigation. The majority of these instances were attributable to incorrect selection of the radial velocity. A separate routine was developed that carries out an automated visual inspection of these spectra in order to select, where possible, an improved radial velocity correction from the full list of radial velocity masks. In some instances the normalisation of the spectra was poor due to noisy spectra, strange spectral features or an invalid radial velocity correction. In the latter case the adjustment to the radial velocity would correct the poor normalisation. In the former cases these spectra were captured by quality/rejection flags in the normalisation process or in the construction of the final ESO dataset.

After these adjustments were made the spectra were reprocessed in the second iteration of SPB. This process is exactly the same as the first iteration except the mean spectral FWHM of the absorption lines for each wavelength section is now measured directly for each individual spectrum. This second iteration of SPB provides the first scientific estimate of the parameters, and hence convolving the spectrum with spectral FWHM values measured specifically for each spectrum provides a more robust analysis. The measured spectral FWHM values were used to convolve each region of each spectrum to match the FWHM of the AMBRE:FEROS synthetic spectra grid. Therefore in Eq. (3) the FWHM\(_{\text{obs}}\) is the measured spectral FWHM for each wavelength section for each spectrum.

Extensive testing was carried out which compared spectral lines that were in common between the observed and synthetic spectra both before and after convolution in order to ensure that the transformation calculation resulted in appropriately convolved observed spectra. This included testing the procedure on the original high resolution synthetic spectra to ensure the AMBRE:FEROS resolution was obtained after convolution by this procedure. The FWHM\(_{\text{grid}}\) (0.33 mA) was calculated based on a sample of the AMBRE:FEROS synthetic spectra for which the FWHM values were determined by the same method as for
the observed spectrum and were found to be constant with wavelength as expected due to how the synthetic spectra were generated. This ensured consistency in the calculation and application of the FWHM in the convolution process.

Part of the calculation of the spectral FWHM was to classify each spectrum using the number of spectral lines and the widths that were measured. Three categories were used: “Weak” for less than 25 lines measured which were typically found to have \( \text{FWHM} < 0.11 \) (FWHM\text{weak}) and were generally noisy spectra; “Medium” for more than 25 lines measured, where the line depths of the measured lines were between 0.5 and 0.95 in normalised intensity, and no more than 1 strong line was identified (FWHM\text{medium}); and “Strong” for more than 1 strong line identified where the line depth of the “Strong” line was between 0.35 and 0.5 of the normalised intensity (FWHM\text{strong}). The spectral lines were identified by locating the minima and maxima in the spectra, fitting a gaussian at each minima and then discarding those lines which failed the quality tests of the gaussian fit. For a spectrum typically classified as FWHM\text{medium} 1000 to 2000 lines were identified which were reduced to ~400 lines with good gaussian fits.

The “Strong” classification was an attempt to identify the spectra with large, broad spectral features as this provided another estimate of the spectral classification of the star. This was accomplished by applying a high FWHM smoothing function to each spectrum and then testing for which spectral features still remained. A spectrum without “Strong” lines would be reduced to a line near the continuum, while “Strong” features would otherwise still be prominent. This method had to be tailored to the FEROS wavelength region in order to place relevant limits on the possible number of “Strong” lines and their location within the wavelength regions. If, by this method, more than 1 “Strong” line were found then the spectrum was classified as FWHM\text{strong} and the mean spectral FWHM of the “Strong” lines was recorded.

In the cases where the spectrum was designated as FWHM\text{weak} or FWHM\text{strong}, then default spectral FWHM values were used instead. The spectral FWHM values for each spectrum that were used in the convolution, measured or mean values, were saved to an external file to be used in the next stage of the pipeline. Extensive testing was carried out using corresponding spectral lines in the original, convolved and synthetic spectra to confirm that the spectral FWHM procedure correctly convolved the archived spectra to the resolution of the synthetic grid across the wavelength range.

At this stage the spectra which failed the tests identifying unanalysable spectra (extreme emission features, extreme noise-dominated spectra etc.) were rejected from the analysis process. This reduced list was analysed once more in MATISSE in order to determine the first estimate of the stellar parameters to be used in the next stage of the analysis pipeline. The convergence and \( \log \chi^2 \) were tested again to catch any further mis-identified radial velocity corrections. At the end of the second iteration of SPB 2370 FEROS spectra were rejected, 11% of the total number of FEROS archived spectra.

4.4. Spectral Processing C: iterative spectra normalisation and parameterisation

The final number of FEROS spectra that were passed to the final stage of the analysis pipeline, Spectral Processing C (SPC), was 19 181. SPC again uses the same procedures as SPA and SPB with two differences. The first difference is that the spectra are convolved using the spectral FWHM values determined in the second iteration of SPB rather than re-measuring the FWHM. The second difference is the use of a more robust method for the normalisation of the spectra.

Previously, in SPA and SPB, a “rough” normalisation process was carried out that used pre-selected continuum regions to normalise the spectra to unity. But this is an invalid assumption for many stars, in particular for cool stars which have depressed continuum regions due to molecular bandheads that should not be normalised to unity. However at the beginning of SPC we have an estimate of the stellar parameters from the second iteration of SPB that gives us some valid information about each spectrum that we did not previously possess.

We take advantage of this information by using the interpolated synthetic spectra generated for each set of stellar parameters as a better estimate of the continuum placement for the normalisation of the observed spectra. In an iterative process that discards absorption and emission features by polynomial fitting and sigma clipping to leave only “continuum” regions, the synthetic spectrum is divided out of the corresponding observed spectrum leaving a residual of continuum lines. A continuum profile is fitted to this residual which is then divided out of the observed spectrum, normalising it to a pseudo-continuum that better represents the parameters of the star. This process overrides any residual curvature that remains after, or was introduced by, the normalisation to the hot star continuum.

These pseudo-continuum normalised spectra are reanalysed in MATISSE to obtain a second estimate of the stellar parameters and another set of synthetic spectra. The “normalisation treatment & MATISSE analysis” cycle is repeated 9 times, which had been determined to be a sufficient number of iterations within which the normalised spectra and stellar parameters could converge to their optimum state. This typically occurred within five iterations. This approach provides a robust incremental adjustment of the parameters and normalisation process that hones in on a realistic estimate of the stellar parameters of each spectra, a process which is not possible using the normalisation method of SPA and SPB.

Figure 9 shows the normalisation procedure for two FEROS spectra in the wavelength region about two of the magnesium triplet at 5160 Å as an example of depressed continuum regions. The two stars correspond to temperatures of 4600 K (\( \text{SNR} = 115 \)) and 5300 K (\( \text{SNR} = 204 \)) respectively. For each spectrum the normalisation by SPB, the first iteration of SPC and the final solution that was found in SPC are shown. The stellar parameters and \( \log \chi^2 \) (which is calculated over all the AMBRE:FEROS wavelengths) determined at each stage are shown as well as the synthetic spectrum interpolated to the stellar parameters. The difference between the observed and synthetic spectra is also included to show how it converges.

For the cooler star this rough normalisation of SPB is not a good fit as the observed spectrum is set too high compared to the synthetic spectrum generated at the corresponding stellar parameters. This shows how inadequate normalisation to unity is for cool star spectra. However the rough normalisation does provide a reasonable fit to the synthetic spectrum for the hot star although the placement of the continuum is still not ideal.

At the initial iteration of SPC the spectrum undergoes its first normalisation to the previous solution synthetic spectrum. For the cooler star the observed spectrum now sits below unity in better agreement with the new synthetic spectrum, but there are still mismatches in the relative placement of the two spectra and the line depths. For the hotter star there is very good agreement
between the observed and synthetic spectra. In both cases the improvement in the match are shown by the lower log $\chi^2$ values.

The final solution for the cooler star ($i = 6$) shows excellent agreement between the observed and the synthetic spectra, which is reflected in the much lower log $\chi^2$ value. Indeed the visual inspection of the spectra show that they are in good agreement in this section of wavelength. For the hotter star the solution ($i = 7$) is very close to the solution determined in the first iteration of SPC, indeed the log $\chi^2$ does not change. Visually this good fit is obvious between the observed and synthetic spectra. The comparison of these two stars illustrates the greater difficulty there is in the normalisation and parameterisation of cool stars. However the procedure developed for the AMBRE pipeline can successfully manage both cases and the normalisation procedure in SPC is particularly effective in the case of cool stars where the continuum regions are more heavily obscured by spectral features.

Typically convergence occurred by the fourth or fifth iteration of the normalisation and stellar parameter determination cycle. When convergence has occurred the final set of parameters are selected as those from the converged solutions with the lowest log $\chi^2$. In the case of non-convergence within the 9 iterations of SPC, it is assumed that for the final six iterations any bias introduced by the initial rough normalisation has been erased. Of these final six iterations the solution with the lowest log $\chi^2$ is selected as the final stellar parameters for the spectra along with the corresponding normalised and synthetic spectra.

Hence the radial velocity, stellar parameters ($T_{\text{eff}}$, $\log g$, [M/H] and [$\alpha$/Fe]), normalised spectra and corresponding interpolated synthetic spectra were determined in SPC for 19 181 of the 21 551 FEROS spectra.

Extensive testing of the FEROS analysis pipeline was carried out in order to optimise the procedures and to validate the MATISSE results by making comparison to literature values. The following sections describe the different tests and validations that were carried out.

5. AMBRE:FEROS internal error analysis

The internal errors can be used to determine how well MATISSE will derive the stellar parameters of similar stellar types which have different levels of noise and different uncertainties in radial velocity. In order to test for noise and radial velocity effects a sample of 500 synthetic spectra were generated at random stellar parameters covering the entire stellar parameter range, spectral resolution and wavelength domains of the AMBRE:FEROS synthetic spectra grid.

5.1. Internal errors: SNR

To test the effects of noise the sample of synthetic spectra was reproduced five times by adding differing levels of noise per pixel (0.1 Å per pixel for AMBRE:FEROS synthetic grid)
at SNR = 10, 20, 50, 100. Each sample was then re-analysed in MATISSE and the derived stellar parameters were compared to the original values.

Figures 10a to d show how the difference in stellar parameters changes with increasing SNR for $T_{\text{eff}}$, log $g$, [M/H] and [$\alpha$/Fe] respectively. For each noise-added sample the difference between the original and derived parameters was calculated and then the difference value at the 70th percentile (~1σ) was determined. For example, in Fig. 10a at a SNR of 10 (the noisiest), 70% of the synthetic spectra returned $T_{\text{eff}}$ values within 14 K of the original $T_{\text{eff}}$ value. At a SNR of 100, 70% of the sample returned values within 2 K of the original $T_{\text{eff}}$ value. Similarly small errors were also found for log $g$, [M/H] and [$\alpha$/Fe] that diminished to negligible levels at SNR of 100 in all cases. This analysis shows that for two synthetic spectra with very similar minished to negligible levels at SNR of 100 in all cases. This analysis shows that the internal error due to uncertainties in radial velocity on the determination becomes significant (>1%) for each of the stellar parameters. However if the uncertainty in the radial velocity is less than 5 km s$^{-1}$ the MATISSE parameters compare well to the true values.

The internal uncertainty in each stellar parameter due to radial velocity was calculated using the equations that connect the points in Fig. 11. The error on the radial velocity, $\sigma_{\text{V}}(\text{rad})$, determined in the radial velocity programme was used as the input to determine the internal error due to the $V_{\text{rad}}$ uncertainty ($\sigma_{\text{int, V}}$) for each parameter for each spectrum according to the following conditions:

$$\sigma(T_{\text{eff}})_{\text{int, V}} = \begin{cases} 
70.5 & \text{if } \sigma(V_{\text{rad}}) < 0.5, \\
9.60\sigma(V_{\text{rad}}) + 0.40 & \text{if } 0.5 \leq \sigma(V_{\text{rad}}) < 1.0, \\
10.05\sigma(V_{\text{rad}}) - 0.05 & \text{if } 1.0 \leq \sigma(V_{\text{rad}}) < 5.0, \\
13.24\sigma(V_{\text{rad}}) - 16.0 & \text{if } 5.0 \leq \sigma(V_{\text{rad}}) < 10.0.
\end{cases}$$

$$\sigma(\log g)_{\text{int, V}} = \begin{cases} 
0.015 & \text{if } \sigma(V_{\text{rad}}) < 0.5, \\
0.028\sigma(V_{\text{rad}}) + 0.001 & \text{if } 0.5 \leq \sigma(V_{\text{rad}}) < 1.0, \\
0.0287\sigma(V_{\text{rad}}) + 0.00025 & \text{if } 1.0 \leq \sigma(V_{\text{rad}}) < 5.0, \\
0.0558\sigma(V_{\text{rad}}) + 0.135 & \text{if } 5.0 \leq \sigma(V_{\text{rad}}) < 10.0.
\end{cases}$$

$$\sigma([\text{M/H}])_{\text{int, V}} = \begin{cases} 
0.008 & \text{if } \sigma(V_{\text{rad}}) < 0.5, \\
0.012\sigma(V_{\text{rad}}) + 0.002 & \text{if } 0.5 \leq \sigma(V_{\text{rad}}) < 1.0, \\
0.0213\sigma(V_{\text{rad}}) - 0.0073 & \text{if } 1.0 \leq \sigma(V_{\text{rad}}) < 5.0, \\
0.0510\sigma(V_{\text{rad}}) - 0.156 & \text{if } 5.0 \leq \sigma(V_{\text{rad}}) < 10.0.
\end{cases}$$

$$\sigma([\alpha/\text{Fe}])_{\text{int, V}} = \begin{cases} 
0.006 & \text{if } \sigma(V_{\text{rad}}) < 0.5, \\
0.010\sigma(V_{\text{rad}}) + 0.001 & \text{if } 0.5 \leq \sigma(V_{\text{rad}}) < 1.0, \\
0.0128\sigma(V_{\text{rad}}) - 0.0018 & \text{if } 1.0 \leq \sigma(V_{\text{rad}}) < 5.0, \\
0.0204\sigma(V_{\text{rad}}) - 0.0400 & \text{if } 5.0 \leq \sigma(V_{\text{rad}}) < 10.0. 
\end{cases}$$
The rotational velocity ($V\sin i$) of a star can have an impact on the broadening of the observed spectral features depending on the magnitude of the rotational velocity and the resolution of the instrument. The AMBRE synthetic spectra grid was generated with no variations in $V\sin i$, assuming all stars to be slow rotators. However, it is important to consider whether variations in $V\sin i$ will have an effect on the determined stellar parameters. Gazzano et al. (2010) carried out a robust investigation into the $V\sin i$ limits of reliable stellar parameter determination using MATISSE and found that spectra with $V\sin i < 11$ km s$^{-1}$ produced good results in the MATISSE analysis of a sample of FGK dwarfs. Based on this, Gazzano et al. (2010) accepted all CCF FWHM $\leq 20$ km s$^{-1}$ for spectra at a resolution of $R \sim 26000$. The FEROS sample and the AMBRE synthetic spectra grid are of much broader range in stellar parameters and wavelength, and of lower resolution ($R \sim 15000$) than the Gazzano et al. (2010) study. Hence the effects of $V\sin i$ on the range of stars for which the synthetic grid has been optimised, would be well-masked for the AMBRE:FEROS configuration. In Sect. 7.2.3 we examine the FEROS sample in the context of the CCF FWHM to identify spectra which are not well-represented by the AMBRE synthetic spectra grid.

5.3. Internal errors: normalisation

The effects of normalisation on the MATISSE determination were measured using the iterative normalisation process (SPC) of the FEROS analysis pipeline. Using a test sample of 384 FEROS spectra ($\langle$SNR$\rangle = 100 \pm 45$) the changes in normalisation between iterations of SPC was explored.

Figure 12 shows the progression of the goodness of fit between the reconstructed spectra and the normalised spectra at each iteration of SPC as a Box & Whisker graph. Assuming that by the 9th iteration convergence has occurred, the absolute difference in the log $\chi^2$ between the 9th and i-th iteration was calculated for each of the 384 test spectra. The spread in values is largest at $i = 0$ and there is a large decrease in the spread for $i = 1$. The following iterations show a more sedate decrease in spread and appear to have converged from $i = 6$.

To measure the internal error associated with the normalisation process we investigated the difference in stellar parameters between the 9th and 7th iteration for the test sample. Hence we assumed that the solutions have converged from at least the 7th iteration and the subsequent variations in the stellar parameters are due to minor adjustment of the normalisation of the observed spectra. The variation from convergence ($i = 7$) to the final iteration ($i = 9$) was used to give the internal error due to the normalisation process.

Figure 13 shows the difference in each stellar parameter between the 9th and 7th iterations against SNR for the 384 test spectra as black points. These values were binned into two SNR bins ($\Delta$SNR = 30) centred at SNR = 65 and 100, and a single bin for all points with SNR $> 125$. The 2σ uncertainty for each bin is shown as red squares at the bin centre. Red lines connect the bin uncertainty values. For $T_{\text{eff}}$ the spread is constant for each bin and so a constant value for the internal uncertainty due to normalisation was adopted. For log $g$, [M/H] and [$\alpha$/Fe] the spread diminishes with increased SNR until SNR = 125 after which a constant spread is found. The adopted internal errors for these three parameters were therefore a piecewise function of SNR. The following equations define the internal errors due to normalisation for each parameter ($\sigma_{\text{int,norm}}$):

$$
\sigma(T_{\text{eff}})_{\text{int,norm}} = 11.0 \text{ K}
$$

$$
\sigma(\log g)_{\text{int,norm}} = \begin{cases} 
0.0017 \text{ SNR} + 0.2414 \text{ dex} & \text{if SNR} < 100, \\
0.0018 \text{ SNR} + 0.2500 \text{ dex} & \text{if 100} \leq \text{SNR} < 125, \\
0.025 & \text{if SNR} \geq 125, 
\end{cases}
$$

$$
\sigma([\text{M/H}])_{\text{int,norm}} = \begin{cases} 
0.0007 \text{ SNR} + 0.0964 \text{ dex} & \text{if SNR} < 100, \\
0.0004 \text{ SNR} + 0.0690 \text{ dex} & \text{if 100} \leq \text{SNR} < 125, \\
0.014 & \text{if SNR} \geq 125, 
\end{cases}
$$

$$
\sigma([\text{Fe}])_{\text{int,norm}} = \begin{cases} 
0.0006 \text{ SNR} + 0.0671 \text{ dex} & \text{if SNR} < 100, \\
0.0002 \text{ SNR} + 0.0260 \text{ dex} & \text{if 100} \leq \text{SNR} < 125, \\
0.006 & \text{if SNR} \geq 125, 
\end{cases}
$$

5.4. Total internal error

The internal errors due to the SNR, $V_{\text{rad}}$ and Normalisation calculated above were combined in quadrature to give the total internal error for each parameter ($\sigma(\theta)_{\text{int}}$) for each spectrum. This was calculated as follows:

$$
\sigma(\theta)_{\text{int}} = \sqrt{\sigma^2(\theta)_{\text{int,norm}} + \sigma^2(\theta)_{\text{int,vrad}} + \sigma^2(\theta)_{\text{int,SNR}}} 
$$

These values were reported for each spectrum in the ESO dataset as described in Sect. 8.

6. AMBRE:FEROS external error analysis

The external error was quantified by comparing the AMBRE:FEROS stellar parameter values to literature values for key reference stars that exist within the AMBRE:FEROS dataset. A list of stars that had been observed with FEROS but were also part of other high quality spectroscopic studies were identified and this reference sample was used to provide quality criteria for each derived stellar parameter. Unfortunately, reference stars covering the whole range of possible parameters could not be found in the literature. In several cases, estimates of the mean stellar metallicity were lacking and the situation was even worse for the [$\alpha$/Fe] chemical contents since too few reference stars with known [$\alpha$/Fe] content could be found in the literature.
6.1.1. Stellar atlases: Sun, Arcturus, Procyon

Table 4 lists the standard stars and the corresponding spectral atlases used to test the FEROS analysis pipeline.

| Standard star/ATLAS | \(T_{\text{eff}}\) (K) | \(\log g\) (dex) | \([\text{Fe/H}]\) (dex) | \(\Delta\sigma_{\text{Fe}}\) (dex) |
|---------------------|----------------------|-----------------|---------------------|---------------------|
| SUN                 | 5770                 | 4.44            | 0.00                | 0.00                |
| Wallace et al. (1998) | 5732                 | 4.42            | −0.03               | 0.05                |
| Braudt & Neckel (1999) | 5674                 | 4.32            | 0.00                | 0.08                |
| BASS2000            | 5841                 | 4.49            | −0.01               | 0.06                |
| SN2000             | 5735                 | 4.43            | −0.03               | 0.05                |
| ARCTURUS            | 4300                 | 1.70            | −0.60               | 0.20                |
| Hinkle et al. (2000) | 4537                 | 2.12            | −0.49               | 0.28                |
| UVES-POP            | 4468                 | 2.03            | −0.47               | 0.29                |
| PROCYON             | 6600                 | 4.00            | 0.00                | 0.00                |
| UVES-POP            | 6538                 | 4.00            | −0.25               | 0.07                |

Notes. The accepted parameters for the standard stars are listed (bold) as well as the final stellar parameters with bias corrections derived for each atlas in the AMBRE:FEROS analysis.

(a) ftp://ftp.noao.edu/catalogs/arcturusatlas/visual/
(b) ftp://ftp.hs.uni-hamburg.de/pub/outgoing/FTS-Atlas/
(c) Delbouille et al. (1973); http://bass2000.obspm.fr/
(d) Allende Prieto et al. (2004); http://hebe.as.utexas.edu/s4n/
(e) Bagnulo et al. (2003); http://www.sc.eso.org/santiago/uvespop/
(f) Mean of parameters listed on SIMBAD.

6.1.2. Spectral libraries reference sample

Preliminary investigations showed that merely making comparison between the MATISSE results and all of the stellar parameter values that were available in SIMBAD for the FEROS stars resulted in a great deal of disparity. There was no quality determination on the values with which we made the comparison. Homogeneously analysed large sample datasets were necessary in order to ensure that a valid comparison was being made. To this end key spectral libraries were investigated in order to locate within the FEROS archive dataset stars that had been analysed in high quality studies. This list of spectral libraries is given in Table 5.
Table 5. List of spectral libraries used to select comparison stars from the FEROS archived spectra.

| Source                  | Papers                                                                 | Abbrev. | No. stars | No. spectra | Methods applied                                              |
|-------------------------|------------------------------------------------------------------------|---------|-----------|-------------|--------------------------------------------------------------|
| S4N                     | Allende Prieto et al. (2004) \(T_{\text{eff}}, \log g, [\text{Fe/H}], [\alpha/\text{Fe}], V_{\text{rad}}\) | AP04    | 30        | 338         | Alonso et al. (1996, 1999) calibration, isochrones, Allende Prieto (2003) GA, spectroscopic inversion |
|                         |                                                                        |         |           |             |                                                              |
| Gaia RVS Standards      | Crifo et al. (2010) \(V_{\text{rad}}\)                                | C10     | 158       | 318         | Convolution, CORAVEL cross-correlation                       |
|                         | Bensby et al. (2003) \(T_{\text{eff}}, \log g, [\text{Fe/H}], [\alpha/\text{Fe}]\) | B03     | 68        | 68          | Fe I EP balance, parallax, equivalent widths                |
|                         |                                                                        |         |           |             |                                                              |
| PASTEL                  | \(T_{\text{eff}}, \log g, [\text{Fe/H}]\)                           | F97-08  | 49        | 206         | Balmer line wings, Fe IP balance, Fe I/Fe II line profiles |
|                         | Fuhrmann et al. (1997, 1998); Fuhrmann (1998a,b, 2004, 2008)          | G82-03  | 23        | 118         | Fe I EP balance, Fe I/Fe II IP balance, equivalent widths   |
|                         | Gratton et al. (1982); Gratton (1989); Gratton et al. (1996, 2003)    | H&M07   | 6         | 27          | Fe I EP balance, Fe I/Fe II IP balance, equivalent widths   |
|                         | Hekker & Meléndez (2007)                                              | L&H83-06| 41        | 224         | Fe I EP balance, Fe I/Fe II IP balance, equivalent widths   |
|                         | Luck & Bond (1983, 1991); Luck & Heiter (2005, 2006)                  | McWilliam| 10       | 41          | IRFM+colour calibration, parallax, equivalent widths        |
|                         | McWilliam (1990)                                                      | McW90   | 10        | 41          | IRFM+colour calibration, parallax, equivalent widths        |
|                         | Mishenina & Kovtyukh (2001)                                           | M&K01;M04-08 | 42   | 227         | Wings of H\(\alpha\), Fe I EP balance, Fe I/Fe II IP balance, equivalent widths |
|                         | Mishenina et al. (2003, 2004, 2006)                                   | R07     | 26        | 151         | Colour calibration, parallax, equivalent widths             |
|                         | Ramírez et al. (2007)                                                 | V&F05   | 81        | 310         | “Spectroscopy Made Easy”                                    |
|                         | Valenti & Fischer (2005)                                              |         |           |             |                                                              |
|                         | \(T_{\text{eff}}\) only                                               | A96,99  | 22        | 147         | Infra-Red Flux Method (IRFM)                                |
|                         | Alonso et al. (1996, 1999)                                            | B&LG98  | 20        | 152         | IRFM                                                        |
|                         | Blackwell & Lynam-Gray (1998)                                         | dB98    | 18        | 163         | Surface brightness and \(V - K\) calibration               |
|                         | di Benedetto (1998)                                                   | GH&B09  | 26        | 162         | Line depth ratios                                           |
|                         | González Hernández & Bonifacio (2009)                                 | K03-06  | 48        | 354         | SED fit + \(V - K\) calibration                            |
|                         | Kovtyukh et al. (2003, 2004, 2006)                                    | M06     | 57        | 306         | IRFM                                                        |
|                         | Masana et al. (2006)                                                  | R&M05   | 37        | 197         | IRFM + colour calibration                                  |
|                         | Ramírez & Meléndez (2005)                                             |         |           |             |                                                              |
|                         | PASTEL Total                                                          |         | 148       | 618         |                                                              |

Notes. The number of stars from each library and the corresponding number of FEROS spectra are also listed. \(^6^{6}\) FEROS Observations 2000–2001. GA = Genetic Algorithm, EP = Excitation Potential, IP = Ionisation Potential.
The primary spectral library that was used was the PASTEL database (Soubiran et al. 2010), which provides a selection of key papers that carried out detailed spectroscopic studies at high resolution and high SNR to derive high quality stellar parameters. From these papers we were able to identify a total of 148 stars corresponding to 618 FEROS archived spectra (Table 5). There was significant crossover of stars between the studies which provided another level of comparison for this work. From the PASTEL sample ~120 stars (~300 spectra) had values for three stellar parameters, $T_{\text{eff}}$, log $g$ and [Fe/H], while the remaining stars had $T_{\text{eff}}$ values only. In the following comparison we assume the literature values to be correct. However we note that the literature values, while they are high quality studies, were determined using a range of techniques (Soubiran et al. 2010), in particular, several studies determined photometric $T_{\text{eff}}$ values using the Infra-Red Flux Method (Alonso et al. 1996, 1999; Ramírez & Meléndez 2005; González Hernández et al. 2010), in particular, several studies determined photometric $T_{\text{eff}}$ values for three stellar parameters, $T_{\text{eff}}$, log $g$ and [Fe/H], while the remaining stars had $T_{\text{eff}}$ values only. In the following comparison we assume the literature values to be correct. However we note that the literature values, while they are high quality studies, were determined using a range of techniques (Soubiran et al. 2010), in particular, several studies determined photometric $T_{\text{eff}}$ values using the Infra-Red Flux Method (Alonso et al. 1996, 1999; Ramírez & Meléndez 2005; González Hernández et al. 2005; González Hernández & Bonifacio 2009).

Figures 14a to b compare the $T_{\text{eff}}$, log $g$, [M/H] and [$\alpha$/Fe] values determined by AMBRE:FEROS with the literature values of the reference sample and stellar atlases. The legend in each provides the bias and $\sigma$ for each subsample. The axes represent the limits of the final accepted parameters and clearly the reference sample does not cover the entire grid as would be ideal. Despite this limitation the reference sample was sufficient to define bias corrections which were necessary to apply to the AMBRE:FEROS stellar parameters.

6.1.3. Reference sample: $T_{\text{eff}}$

Figure 14a compares the $T_{\text{eff}}$ for each star. It is important to note that many of the literature $T_{\text{eff}}$ values were not determined spectroscopically, as stated above. Despite the differences in measurement technique, overall there is excellent agreement between the AMBRE:FEROS results and the PASTEL values, and also good agreement for the stellar atlases. There is a slight turning over of the distribution from $T_{\text{eff}} \gtrsim 5750$ K for which a small bias correction was applied that will be quantified at the end of this section.

6.1.4. Reference sample: log $g$

Figure 14b shows the comparison of the log $g$ values for which there is reasonably good agreement. However the majority of the sample are dwarfs (high log $g$), and the giants (low log $g$) are under-represented. There is a slight gradient within the sample which resulted in a small bias correction being applied at high log $g$ for the final results (see end of the section). The lack of a significant sample of giants made it difficult to accurately define a bias at low log $g$, hence the correction was only

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Footnote:

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applied at high log $g$. However the uncertainty on the giants is definitely higher.

6.1.5. Reference sample: [M/H]

Figure 14c shows the comparison of the AMBRE:FEROS [M/H] values with the literature [Fe/H] values. The difference in definition between [M/H] and [Fe/H], as discussed previously, means that this is not an accurate comparison, as so many more elements (albeit of lesser contribution) than just Fe are included in the MATISSE metallicity. The comparison shows an overall systematic offset between the AMBRE:FEROS and literature values reflected in the subsample biases. The sources of this bias are difficult to quantify so we assumed a direct comparison between [Fe/H] and [M/H] then made a systematic bias correction to the final [M/H] results as outlined at the end of this section.

6.1.6. Reference sample: [$\alpha$/Fe]

Unfortunately very few comparison stars with published values of their [$\alpha$/Fe] exist in the literature. The stellar parameters in the S4N library (Allende Prieto et al. 2004) also include $\alpha$ element abundances where possible, which provided some comparisons with the AMBRE:FEROS [$\alpha$/Fe] results. The S4N $T_{\text{eff}}$ values were determined using photometric calibrations while the log $g$ were determined from Hipparcos parallaxes. The chemical abundances were determined using $\chi^2$ minimisation of the spectral line profile between the observed spectrum and a grid of synthetic spectra (Allende Prieto et al. 2004).

A key study that was investigated specifically in order to validate the AMBRE:FEROS [$\alpha$/Fe] values was Bensby et al. (2003). This study was a detailed analysis of 66 F and G dwarf stars in the galactic disc, and accurate abundances were determined for the $\alpha$ elements, Mg, Si, Ca, Ti. The mean of these abundances was taken as the value for the global [$\alpha$/Fe] with which to compare the AMBRE:FEROS [$\alpha$/Fe] values. This study was particularly useful because the spectra analysed in Bensby et al. (2003) were observed with FEROS. However the observations took place in 2000 and 2001 and so the spectra were not part of the archived sample delivered to AMBRE. The original spectra were obtained (Bensby, priv. comm.) and analysed in the AMBRE:FEROS pipeline. Hence a direct comparison between AMBRE:FEROS and Bensby et al. (2003) could be made for all four parameters (see Figs. 14a to c).

Figure 14d shows the comparison of the AMBRE:FEROS [$\alpha$/Fe] results with the values from the S4N library, Bensby et al. (2003) and the stellar atlases. The biases are also listed for each sample. There is reasonably good agreement with these samples. In particular there is excellent agreement in the results for the Bensby et al. (2003) stars, which as stated above, is a study where the $\alpha$ element abundances were carefully determined, thereby providing an excellent validation of the AMBRE:FEROS [$\alpha$/Fe] results.

6.2. Bias corrections

The sample of reference stars and the spectral atlases provided a crucial comparison at all stages of the development of the AMBRE:FEROS analysis pipeline. Although it was not possible to cover the entire parameter space, and the reference sample was ultimately biased towards metal-rich dwarfs over a small temperature range, the results provided sufficient information with which to identify biases within the analysis, assuming the literature values to be correct. The following corrections were made in order to remove these biases:

$$T_{\text{eff}}(\text{cor}) = \begin{cases} T_{\text{eff}} - 35 & \text{if } T_{\text{eff}} < 5300 \text{ K}, \\ T_{\text{eff}} + 0.21 \times T_{\text{eff}} - 1141.4 & \text{if } 5300 \leq T_{\text{eff}} \leq 6000 \text{ K}, \\ T_{\text{eff}} + 110 & \text{if } T_{\text{eff}} > 6000 \text{ K}. \end{cases}$$

$$\log g(\text{cor}) = \begin{cases} \log g - 0.296 \times \log g + 1.388 & \text{if } \log g \geq 4.0, \\ \log g + 0.204 & \text{if } 3.75 \leq \log g < 4.0, \\ \log g + 0.817 \times \log g - 2.860 & \text{if } 3.5 \leq \log g < 3.75, \\ \log g & \text{if } \log g < 3.5. \end{cases}$$

$$[\text{M/H}](\text{cor}) = [\text{M/H}] + 0.15$$
$$[\alpha/\text{Fe}](\text{cor}) = [\alpha/\text{Fe}]$$

There are several potential sources of the bias corrections for the AMBRE:FEROS analysis. First, the differing techniques, as well as differing spectral domains and resolutions, used in the determination of the stellar parameters by the reference studies compared with AMBRE:FEROS analysis may make a significant contribution to the difference in derived parameters. Second, while the normalisation process within the AMBRE analysis pipeline was designed to be as robust as possible, the normalisation of stellar spectra is an inherently complex problem, particularly over large wavelength ranges with many spectral features. For spectra of particular spectral types (e.g. cool, low gravity, metal-rich) the normalisation procedure may not be as robust as for less detailed spectra. Third, due to the length of the atomic and molecular linelist that was required to synthesise the AMBRE:FEROS wavelength regions, it was inefficient to carry out a calibration of the linelist to standard stars (i.e. the Sun). It was assumed that the number of lines was statistically sufficient to dampen the noise from ill-fitted spectral features but non-calibration of the line list may play some role in the degree of the bias corrections. Individually, and in combination, these are the most likely sources of the bias corrections for the AMBRE:FEROS stellar parameters.

6.3. External error

The external error for each stellar parameter was determined from the above reference sample analysis. For each individual study the mean difference and spread in differences ($\sigma$) between the literature and AMBRE:FEROS values was calculated. The mean of these $\sigma$ values for each stellar parameter was taken as the global external error ($\sigma(\theta)_{\text{ext}}$) for all of the spectra as follows:

$$\sigma(T_{\text{eff}})_{\text{ext}} = 120 \text{ K}$$
$$\sigma(\log g)_{\text{ext}} = \begin{cases} 0.20 \text{ dex} & \text{if } \log g \geq 3.2, \\ 0.37 \text{ dex} & \text{if } \log g < 3.2, \end{cases}$$
$$\sigma([\text{M/H}]_{\text{ext}} = 0.10 \text{ dex}$$
$$\sigma([\alpha/\text{Fe}]_{\text{ext}} = 0.10 \text{ dex}$$

7. ESO table: rejection criteria

The construction of the ESO Table of stellar parameters that would be delivered to the ESO Archive was the final stage of the pipeline. At this stage the quality flags and tests included throughout the pipeline were drawn together to provide the final set of stellar parameters. The columns headings, their definitions, range of values, null values and rejection conditions used to construct the ESO Table are listed in Table A1.
There were two phases of rejecting spectra in the AMBRE:FEROS analysis. First, prior to analysis in SPC spectra were rejected due to spectral quality issues. This will be described in the Sect. 7.1. Second, the remaining spectra were all analysed in SPC for their stellar parameters and then another set of rejection criteria were applied to construct the final table. These criteria are described and discussed in Sects. 7.2 to 7.4.

7.1. Pre-SPC: non-standard spectra

Prior to SPC, 11% of the FEROS archived spectra were rejected as being non-analyisable primarily due to being non-standard spectra. The rejection flags, as defined in Sect. 4.1, were for “Faulty Spectra”, “Extreme Emission Features”, “Poor Normalisation” and “Excessive Noise”. Often the nature of a spectrum rejected at this stage meant the conditions for two or more of these flags were met. For instance, an extremely noisy spectrum was also consequently poorly normalised. Quality flags were also attached for “Large Emission Features” and “Instrumental Relics” but these spectra were not rejected based solely on these flags. The number of spectra identified for each rejection flag and for each quality flag are listed in Table 6.

7.2. Rejection criteria post-SPC

The remaining 89% of the FEROS spectra were analysed in SPC and so stellar parameters were obtained for each of these spectra. Figure 15a shows the HR diagram for all the stellar parameters obtained in SPC. The giant branch and main sequence can be observed but there is a great deal of mis-classification at hot temperatures for log \( g \) ≥ 2 dex, and an overdensity at \( T_{\text{eff}} \approx 3000 \) K and log \( g \) ≥ 2 dex. Most noticeable is that there are many spectra (8947) for which the derived parameters lie outside the \( T_{\text{eff}} \) and log \( g \) boundaries of the synthetic grid which are indicated in red.

![Fig. 15. HR diagrams of the FEROS stellar parameters as each rejection criterion is applied as follows: a) all spectra analysed in SPC; b) spectra with \( V_{\text{rad}} \text{CCF} \) with positive contrast, \( \delta \alpha_{\text{rad}} \leq 0.2 \) and \( \delta \alpha_{\text{rad}} \leq 0.1 \); c) \( \sigma_{\text{rad}} \leq 10 \text{ km s}^{-1} \); d) CCF \( \text{FWHM} \leq 40 \text{ km s}^{-1} \); e) \( \text{FWHM} \) of strong spectral lines \( \leq 0.8 \) mÅ; f) \( \text{FWHM} \) of strong spectral lines \( \leq 8 \) mÅ. The \( T_{\text{eff}} \) and log \( g \) boundaries of the synthetic grid are indicated in red.

![Table 6. Number of spectra rejected (Rej) prior to SPC based on the defined rejection flags and the number of spectra accepted (Acc) for SPC but identified by the quality flags.]

| Spectra | No. Spectra | % of FEROS | % of Rej/Acc sample |
|---------|-------------|------------|---------------------|
| FEROS   | 21,551      |            |                     |
| Rejected before SPC | 2370 | 11 | 3.0 |
| Faulty files | 71 | 0.33 | 3.0 |
| Extreme emission features | 1455 | 6.75 | 61.4 |
| Excessive noise | 109 | 0.51 | 4.6 |
| Poor normalisation | 735 | 3.41 | 31.0 |
| Accepted for SPC | 19,181 | 89 | |
| Large emission features | 3177 | 14.7 | 16.6 |
| Instrumental relics | 58 | 0.27 | 0.30 |
Table 7. Summary of the number of spectra flagged as satisfying the conditions of the post-SPC rejection criteria.

| Spectra rejected | pre-SPC | post-SPC |
|------------------|---------|----------|
| Total            | 2370    | 12673    |
| PD CCF\(^a\)     | 549     | 3668     |
| \(\sigma_{\text{V}_{\text{rad}}}\)\(^b\) | 309     | 6348     |
| CCF\(^b\)        | 469     | 10062    |
| FWHM\(^a\)       | 7       | 1199     |
| Grid\(^a\)       | –       | 8494     |

Notes. The spectra that were rejected prior to the SPC analysis were rejected due to other quality issues (see Sect. 7.1). The spectra analysed in SPC were accepted or rejected based on the listed criteria.

(a) Poorly-Defined CCF (Sect. 7.2.1); (b) \(\sigma_{\text{V}_{\text{rad}}} > 10 \text{ km s}^{-1}\) (Sect. 7.2.2); (c) FWHM CCF > 40 km s\(^{-1}\) (Sect. 7.2.3); (d) FWHM\(_{\text{medium}}\) > 0.8 mÅ or FWHM\(_{\text{long}}\) > 8 mÅ (Sect. 7.2.4); (e) parameters lie outside grid parameter space (Sect. 7.4).

Indicators relating to the measurement of the spectral FWHM and the radial velocity proved to be the least affected by spectral type and better identified the mis-classifications. The key indicators that were explored were: the \(V_{\text{rad}}\) CCF contrast, the error on the amplitude and the error on continuum of the \(V_{\text{rad}}\) CCF; the \(V_{\text{rad}}\) error; the FWHM of the \(V_{\text{rad}}\) CCF; the spectral FWHM; and the AMBRE:FEROS synthetic grid boundaries in \(T_{\text{eff}}, \log g, [\text{M/Fe}]\), \([\text{O/Fe}]\).

Table 7 summarises the number of spectra that satisfied the conditions of each rejection criteria for the spectra rejected before and after SPC. The post-SPC rejection criteria were applied to the spectra rejected before SPC to further characterise that sample. To illustrate the effects of the rejection criteria on the sample of spectra analysed in SPC, Fig. 15 shows progressively the HR diagram of the AMBRE:FEROS stellar parameters as each criteria is applied. The limits of the synthetic grid in \(T_{\text{eff}}\) and \(\log g\) are also shown. This process is discussed in the following sections.

7.2.1. CCF contrast, amplitude and continuum

As described in Sect. 4.2, the radial velocity was determined using binary masks to calculate a CCF. The contrast, the depth (amplitude) of the CCF, the continuum placement of the CCF and their associated errors (as calculated in the IDL:GAUSSFIT routine) are measures of the quality of the CCF. The rejection criteria were derived from these quantities as follows:

1. A negative contrast means that the CCF profile is inverted, which is contrary to the expected result, hence all spectra with a negative contrast were rejected.
2. The relative error on the amplitude of the CCF (\(\frac{\sigma_{\text{ampl}}}{\text{ampl}}\)) is mainly a measure of how much noise there is in the CCF. If the \(\frac{\sigma_{\text{ampl}}}{\text{ampl}}\) ≤ 0.2 then the noise makes up less than 20% of the profile depth. We rejected spectra with \(\frac{\sigma_{\text{ampl}}}{\text{ampl}} > 0.2\).
3. The placement of the continuum is also an indication of how well the CCF was defined. The relative error on the continuum (\(\frac{\sigma_{\text{cont}}}{\text{cont}}\)) gives a measure of the noise in the continuum placement. We rejected all spectra with \(\frac{\sigma_{\text{cont}}}{\text{cont}} > 0.1\).

These three criteria identify spectra with poorly-defined CCFs. Combined they are the first criteria to be applied to the SPC dataset resulting in the rejection of a further 3668 spectra from the analysis. Figure 15b shows the resulting HR diagram after these spectra were removed. The hot star mis-classifications and overdensity at \(T_{\text{eff}} \approx 3000 \text{ K}\) are diminished.

Figures 16a and b show histograms of the SNR per pixel (0.03 or 0.06 Å per pixel) for the full sample of FEROS archived spectra. The SNR has been calculated within the analysis pipeline using sigma clipping on either the predefined continuum regions in SPB or the pseudo-continuum residual in SPC. The highest SNR values do not necessarily mean optimum spectra and are identified by the quality flags within the pipeline as required. Figure 16a is the histogram of the spectra that were accepted as being of good quality and with a well-defined CCF, whereas Fig. 16b is the histogram of the spectra rejected as being of poor quality or with poorly-defined CCF. Note that the y-axis scale is different between the figures. The histogram in Fig. 16a peaks between 50 and 100 SNR. There are significantly fewer spectra with SNR < 50 in Fig. 16a compared with Fig. 16b, indicating the cleaning thus far has indeed identified low SNR spectra. However, as Figs. 10a to d shows, even at SNR of 50 the internal error is negligible.

7.2.2. \(V_{\text{rad}}\) error

The next criterion to be applied was the \(V_{\text{rad}}\) error (\(\sigma_{\text{V}_{\text{rad}}}\)) which had been calculated using the prescription in Tonry & Davis (1979). Figure 17 shows histograms of the \(\sigma_{\text{V}_{\text{rad}}}\) determined for each of the FEROS archived spectra of good quality with a well-defined CCF. Figure 17a is a histogram of \(\sigma_{\text{V}_{\text{rad}}} < 10 \text{ km s}^{-1}\).
in bins of 1 km s\(^{-1}\), and Fig. 17b is a histogram of \(\sigma_{V_{\text{rad}}} > 10\) km s\(^{-1}\) in bins of 10 km s\(^{-1}\). The majority of the spectra have a low \(\sigma_{V_{\text{rad}}} (<0.5\) km s\(^{-1}\)), and based on the synthetic spectra analysis in Sect. 4.2, Fig. 11 shows that \(\Delta V_{\text{rad}} \leq 5\) km s\(^{-1}\) correspond to reasonable variations in the stellar parameters (\(\theta\)). However there are a significant number of spectra with \(\sigma_{V_{\text{rad}}} > 10\) km s\(^{-1}\) (see Fig. 17b) which correspond to much larger uncertainties in the \(\theta\) determination.

Hence we decided to reject all spectra with \(\sigma_{V_{\text{rad}}} > 10\) km s\(^{-1}\). This resulted in a further 3550 spectra being rejected from the SPC dataset. The resulting HR diagram is shown in Fig. 15c. Again more of the mis-classifications at hot \(T_{\text{eff}}\), particular for high \(\log g\), have been removed and the overdensity at \(T_{\text{eff}} \approx 3000\) K is considerably diminished.

7.2.3. FWHM of \(V_{\text{rad}}\) CCF

Astrophysically, the FWHM of the \(V_{\text{rad}}\) CCF can be used to calculate the rotational velocity (\(V\sin i\)) of a star. That particular calculation was not carried out here, for the purposes of expediency, but the FWHM of the CCF was used to identify spectra with broadened spectral features that would not be well represented by the synthetic grid. As discussed in Sect. 5.2, the synthetic spectra grid was generated with no variations in \(V\sin i\), assuming all stars to be slow rotators. The analysis in Gazzano et al. (2010) found that for a sample of FGK dwarfs \(V\sin i < 11\) km s\(^{-1}\) produced good results in the stellar parameter determination by MATISSE. At a resolution of \(R \sim 26000\) an upper limit of CCF FWHM = 20 km s\(^{-1}\) was set. Due to the lower resolution (\(R \sim 15000\)), and hence greater masking of the effects of \(V\sin i\), of the AMBRE:FEROS analysis we decided to relax that criterion.

The comparison of spectral type with \(V\sin i\) is used to show the increase in the number of fast rotators with hotter spectral type for both dwarfs and giants (see Chap. 18, Gray 2005). For dwarfs, fast rotators begin to appear at approximately \(F2\) in spec-

FWHM limit of CCF

Fig. 15c. Again more of the mis-classifications at hot \(T_{\text{eff}}\), particular for high \(\log g\), have been removed and the overdensity at \(T_{\text{eff}} \approx 3000\) K is considerably diminished.

7.2.4. FWHM of the stellar spectrum

The calculation of the spectral FWHM was necessary in order to convolve the observed spectra to the resolution of the synthetic grid. However the FWHM values also provide an extra degree of information with which to understand the spectral dataset. The FWHM measurements were separated into FWHM\(\text{weak}\), FWHM\(\text{strong}\) and FWHM\(\text{medium}\) classifications (Sect. 4.3). In particular the FWHM\(\text{strong}\) classification allowed us to identify spectra with broad features, independently of the FWHM of the \(V_{\text{rad}}\) CCF. For each FWHM\(\text{strong}\) spectra, the FWHM of the medium strength lines was measured (where possible) as well as the FWHM of the identified strong lines. Figures 19a–c show the relationships between FWHM\(\text{medium}\) and CCF FWHM, FWHM\(\text{medium}\) and \(T_{\text{eff}}\), and FWHM\(\text{strong}\) and \(T_{\text{eff}}\) respectively for the dataset as of the application of the \(\sigma_{V_{\text{rad}}}\) criterion.

In Fig. 19a there is a clear trend of increasing CCF FWHM with FWHM\(\text{medium}\). The line at FWHM\(\text{medium} = 0.8\) mÅ intersects the trend at the threshold for CCF FWHM = 40 km s\(^{-1}\), where the trend becomes less dense. This provided the first criterion that we applied such that any spectra with a measured FWHM\(\text{medium}\) greater than 0.8 mÅ were rejected.

Figure 15e shows the HR diagram after the removal of spectra based on this criterion. There is some improvement such that the density of the hot mis-classifications is decreased. However there are still a significant number of mis-classifications at temperatures greater than 7000 K. Further investigation showed that these remaining mis-classifications were connected to the “strong” FWHM classification (FWHM\(\text{strong}\)).

There was no clear trend between the CCF FWHM and the FWHM\(\text{strong}\) with which to derive a threshold of rejection. Instead a comparison was made to \(T_{\text{eff}}\). Figure 19b compares the \(T_{\text{eff}}\) with the corresponding FWHM\(\text{medium}\). The lower boundary of possible values of \(T_{\text{eff}}\) does increase with FWHM\(\text{medium}\) and at the derived threshold of FWHM\(\text{medium} = 0.8\) mÅ this trend intersects with a \(T_{\text{eff}} = 7000\) K. For greater FWHM\(\text{medium}\), values the trend of the lowest \(T_{\text{eff}}\) disappears into greater scatter, essentially attributing a temperature value to the FWHM\(\text{medium}\) threshold. This temperature value was used to infer a FWHM\(\text{medium}\) threshold for the strong spectral lines.

Figure 19c shows the measured FWHM\(\text{strong}\) (where possible) with the \(T_{\text{eff}}\). There is distinct trend of increasing temperature with increasing FWHM\(\text{strong}\). The nominal temperature threshold at 7000 K intersects with this trend at a FWHM\(\text{strong} \approx 8\) mÅ. At greater FWHM\(\text{strong}\), there is an increase in scatter and the trend becomes more dispersed. The greater spread in values for spectra with FWHM\(\text{strong} > 8\) mÅ, and for FWHM\(\text{medium} > 0.8\) mÅ, may indicate a greater uncertainty in the parameterisation as the boundary of the grid is approached.

Based on this we can assume that spectra with a FWHM\(\text{strong}\) not more than 8 mÅ have stellar parameters that fall within the synthetic grid and are well-defined. Hence this provided the second criterion to apply such that spectra with a FWHM\(\text{strong} > 8\) mÅ were rejected. Figure 15f shows that indeed the mis-classifications between 7000 K and 8000 K are significantly reduced after the application of this criterion.
7.3. Understanding the rejected spectra

Figure 20 explores the types of stars to which the 11 971 rejected spectra correspond. Figures 20a compares $T_{\text{eff}}$ with $\sigma_{\text{Vrad}}$ for the rejected sample. It is clearly bimodal with the primary peak at 8190 ± 924 K (10 068 spectra) and the secondary peak at 3351 ± 679 K (1903 spectra). The datasets corresponding to the key rejection criteria are also indicated: FWHM$_{\text{medium}}$ rejections (pink), FWHM$_{\text{strong}}$ rejections (yellow) and CCF FWHM rejections (cyan). The spectra rejected on the basis of FWHM$_{\text{medium}}$ and FWHM$_{\text{strong}}$ are concentrated in the two peaks. This argues that at least 84% of these spectra with broad features have returned parameters greater than the 8000 K temperature limit of the synthetic spectra grid.

Figure 20b is the HR diagram of the rejected spectra. The majority have $T_{\text{eff}} > 6000$ K, but also with $\log g > 2$ dex implying these are sub-giants and dwarf stars. It is possible that these spectra have parameters that lie outside the available synthetic grid parameter space and MATISSE has compensated by varying one or all of the stellar parameters away from the true values in order to converge on a solution. Combined with the high number of these spectra that have been measured to have broadened spectral features, it is possible that these are in fact hot and/or fast rotating stars for which the synthetic grid has not been designed.

In Fig. 20a 16% of the rejected sample are located near the cool temperature limit of the synthetic grid (3351 ± 679 K).
Are these truly such cool stars? Figure 20c compares the \( T_{\text{eff}} \) with \([\text{M/H}]\). The cool temperature limit spectra show an interesting trend of decreasing \([\text{M/H}]\) with decreasing temperature. For spectra with \( T_{\text{eff}} < 3000 \) K the \([\text{M/H}]\) extends significantly outside the grid range. A possible answer is that these are hot or fast rotating stars (based on the FWHM measurements) for which MATISSE has converged to a significantly cooler temperature with significantly decreased metallicity. A similar but less dramatic argument can be made for the spectra rejected at the hot temperature limit. These lie within the \([\text{M/H}]\) range for the synthetic grid although many are metal-poor to \(-3.5\) dex. Combined with the high gravity values of these spectra the low metallicity emphasises the possibility that these stars have been mis-classified due to their parameters lying outside the grid parameter space.

The comparison of \([\alpha/\text{Fe}]\) to \( T_{\text{eff}} \) in Fig. 20d adds weight to this, as the cool temperature limit group have very high \([\alpha/\text{Fe}]\) values. For the hot temperature limit group the \([\alpha/\text{Fe}]\) are typically within the range of the synthetic grid.

From this examination of the key variables, the main argument that can be drawn is to why \(-66.1\%\) of the spectra analysed in SPC have been rejected by these very strict criteria is that they are most likely hot and/or fast rotating stars that are not well-represented by the current synthetic grid. Hence MATISSE has converged on solutions which mis-classify the spectra in the absence of a grid covering the required parameter range. This is not an unreasonable conclusion given that FEROS was designed and built by researchers within the hot star community (Kaufer et al. 1999). A visual inspection was carried out on large random samples of the spectra that were rejected based on each of the rejection criteria. This inspection supported the appropriate application of the criteria as the rejected spectra corresponded to noisy, peculiar or strong-lined spectra, and there was an over-representation of spectra comprising solely of strong balmer lines. However to confirm this conclusion these spectra can be re-analysed at a later date when a synthetic grid with \( T_{\text{eff}} > 8000 \) K and a synthetic grid with a range in \( V \sin i \) values will be available.

### 7.4. Synthetic grid boundaries

The above rejection criteria succeeded in cleaning the stellar parameter sample without losing the well-defined branches of stellar evolution that are observed in the HR diagram. Finally, the stellar parameters determined near the boundaries of the synthetic grid are inherently less reliable due to there being less synthetic spectra present to aid in the definition of the parameters. Hence the final rejection criteria that were applied are based on the stellar parameters themselves. Taking into account boundary effects in the learning phase of MATISSE, the limits on the accepted parameters are defined as follows:

\[
3000 \leq T_{\text{eff}} \leq 7625
\]

\[
1 \leq \log g \leq 5
\]

\[
-3.5 \leq [\text{M/H}] \leq 1
\]

\[
-0.4 \leq [\alpha/\text{Fe}] \leq 0.4 \quad \text{if } [\text{M/H}] \geq 0.0
\]

\[
-0.4 \leq [\alpha/\text{Fe}] \leq 0.8 \quad \text{if } -1.0 < [\text{M/H}] < 0
\]

\[
0.0 \leq [\alpha/\text{Fe}] \leq 0.8 \quad \text{if } [\text{M/H}] \leq -1.
\]

Three categories were established using these definitions:

1. **TON**: \( T_{\text{eff}} \) values only within the accepted limits;
2. **TGM**: \( T_{\text{eff}} \), \( \log g \) and \([\text{M/H}]\) values within accepted limits;
3. **TGMA**: \( T_{\text{eff}} \), \( \log g \), \([\text{M/H}]\) and \([\alpha/\text{Fe}]\) within accepted limits.

The final count of spectra for these three categories were: TON = 394 spectra (~292 stars), TGM = 97 spectra (~87 stars), TGMA = 6017 spectra (~2780 stars).

In total 12 673 spectra were rejected from the SPC analysis. The final set of the stellar parameters accepted for delivery to ESO was comprised of 6508 spectra (~3087 stars). For the radial velocity analysis 11 963 spectra (~4505 stars) were determined to have radial velocities with \( \sigma_{V\sin i} \leq 10 \) km s\(^{-1}\) and these values were also accepted for delivery to ESO. The full breakdown of the spectral number count is given in Table 8.

### 7.5. Repeated observations

The FEROS dataset contains many stars with repeated observations. Figure 21 shows histograms of the number of stars per number of observations for the TGMA sample. Figure 21a shows the histogram for the number of stars with observations less than or equal to 10, while Fig. 21b shows the number of stars with observations greater than 10. There is a significant number of stars with 2 or more observations, and some stars have been observed a great many times. The repeated observations were used to assess whether the AMBRE analysis derived consistent stellar parameters for the same star from different spectra.

All objects with 2 or more observations were extracted from the TGMA sample in order to explore the agreement in parameters for a single object over a number of observations of similar SNR. We also extracted from SIMBAD a list of objects within the FEROS dataset (based on a 10\(^{+}\) coordinate search) that were designated as a binary and/or variable star. We found 17 binary stars and 111 variable stars in the final sample which...
we eliminated from the sample of repeat observations. Hence we removed those stars with possible astrophysical variations in the parameters to be left with variations that are solely due to the analysis process. (It must be noted that, as the list in SIMBAD is not exhaustive, it is likely that there are still stars for which the parameters vary astrophysically within the repeated observations sample.) The repeated observations sample was further constrained such that each set of repeats had a mean SNR less than 200, as the observations above SNR = 200 were sparcely sampled. Due to the prior application of the rejection criteria there were no spectra present with SNR < 40. Finally, the sample of repeats comprised of 584 stars equating to 3438 spectra. The maximum number of repeats was 73 for a single star in the repeats sample.

This sample represents the internal error of the AMBRE analysis hence we compared it to the internal error ($\sigma_{\text{int}}$) that we calculated for each spectra based on the SNR, $V_{\text{rad}}$ and Normalisation (Sect. 5).

Figure 22 replicates the relationships in Fig. 10 but for the repeats sample. For each set of repeats the mean and standard deviation were calculated for the SNR and stellar parameters. The standard deviation of each of the parameters was used to represent the change in the parameters ($\Delta \theta$) for each set. The mean SNR values were binned (binsize = 25) and for each bin the 70th percentile of the $\Delta \theta$ was calculated. These values are shown in black. Also included in grey for each parameter are the internal errors calculated for the TGM spectra ($\sigma_{\text{int}}$, see Sect. 5).

In Fig. 22a the variations in $T_{\text{eff}}$ with SNR for the repeats sample is greater than the calculated internal error distribution and almost traces out the upper limit on the internal error. However, in general these two independent measurements of the internal error are on the same order of magnitude confirming the internal consistency within the AMBRE:FEROS analysis.

As defined in Sect. 5 the internal errors for log $g$, [M/H] and [$\alpha$/Fe] are calculated as a piecewise function of SNR and this is reflected in the distribution of the internal errors for TGM as shown in Figs. 22b to d. The lower limit of the internal errors for each of these three parameters decreases with increasing SNR to the defined constant value at SNR = 125. This distribution is replicated in the variation of each parameter with SNR for the repeats sample. At low SNR there is an offset in log $g$ and [M/H] between the two sets of values where the internal errors are of greater value. This implies that the internal errors may be over-estimated at low SNR in light of the repeats sample analysis. For [$\alpha$/Fe] the variations due to the repeats sample trace the internal errors very well across the range of SNR values. But overall, for $T_{\text{eff}}$, these two independent measurements of the internal error for each parameter are in very good agreement.

The distribution of the variation of the parameters with SNR for the repeats sample in Fig. 22 also replicates the analysis of the internal error due to SNR that was carried out using the synthetic spectra sample as shown in Fig. 10. For the lowest SNR bin for each parameter the $\Delta \theta$ are only 2 or 3 times the equivalent value in the synthetic case which is reasonable, and as the SNR increases the $\Delta \theta$ decreases significantly also replicating the synthetic case.

The comparison between the two measures of the internal error from the observed spectra and with the synthetic spectra analysis shows that the analysis carried out within the pipeline is consistent such that closely comparable parameters are found for different spectra of the same star, and so for different stars of similar stellar parameters, at a particular SNR value.

8. The AMBRE:FEROS stellar parameters

The final FEROS stellar parameters are shown in Fig. 23 in six combinations with which to examine the distribution of the parameters.

Figure 23a shows the Hertzsprung-Russell (HR) diagram of the TGM (red) and TGMA (black) samples. The branches of stellar evolution are distinct although broad in width with some small scatter evident throughout the figure. Figure 23b compares [M/H] with $T_{\text{eff}}$ and the majority of the spectra show near solar metallicities ($-1.0 < [\text{M/H}] < 0.5$ dex) consistently across the temperatures.

Figure 23c compares [$\alpha$/Fe] with $T_{\text{eff}}$ and shows an interesting distribution of low (<0.0 dex) and high (>0.2 dex) [$\alpha$/Fe] at low temperatures but an even distribution of solar [$\alpha$/Fe] across all temperatures.

Figure 23d compares [M/H] with log $g$ and the separation between the giants and dwarfs is evident. There is larger scatter in [M/H] at low log $g$ (giants).

Figure 23e compares [$\alpha$/Fe] with log $g$ and again the scatter is larger at low log $g$ (giants).

Figure 23f compares [$\alpha$/Fe] with [M/H] for which there is a clear trend of enhanced [$\alpha$/Fe] (~0.35 dex) at low [M/H] (<-1.0 dex). For [M/H] of solar and above the [$\alpha$/Fe] decreases to solar and depleted values.

Combining three parameters in one graph provides another perspective with which to further explore this dataset. Figure 24 shows the HR diagram of the TGM and TGMA samples binned by colour in [M/H]. The near solar [M/H] bins contain the majority of the sample as expected from Fig. 23 across the Main Sequence and RGB. The samples within the more metal-poor bins lie to the left of the stellar evolution branches which agrees with stellar evolutionary tracks. Interestingly the majority of the Horizontal Branch (HB) are metal-poor ([M/H] = -1.62 dex).
The stellar parameterisation of the FEROS archive completes the first phase of the AMBRE Project. Of the 21 551 FEROS object spectra that were delivered to OCA, 19 181 could be analysed for their stellar parameters. The quality flags and error analysis resulted in a final total of 6508 spectra (~3087 stars) with stellar parameters to be delivered to ESO. Also delivered to ESO are the radial velocity values for 11 963 spectra (~4505 stars).

Approximately 70% of the spectra were rejected from the stellar parameterisation analysis. While 28% were rejected due to the quality of the spectra being insufficient for analysis, 42% were rejected due to astrophysical reasons. From this analysis it seems very likely that hot and/or fast rotating stars are the favoured observational object for FEROS. As the current synthetic grid is defined for stars cooler than 8000 K only then the high number of rejections was inevitable. In general the AMBRE synthetic spectra grid is currently configured for slow-rotating FGK stars. Non-standard stars, such as binaries and chemically peculiar stars, within the FEROS sample may still pollute the final accepted dataset as we were unable to identify such stars at this stage.

However, stellar parameter extensions to the synthetic spectra grid are under development and identification tools are also being developed such that a larger range of spectra may be reliably analysed at a later date.

In summary the work carried out in the analysis of the FEROS archived spectra has resulted in:

1. The development of a complex and robust analysis pipeline for the determination of stellar parameters of large spectral datasets through automated iterative spectral reduction and MATISSE analysis.
2. The establishment of tools with which to exploit spectral atlases and libraries for the comprehensive testing and validation of the results from the AMBRE analysis pipeline.
3. The determination of stellar parameters ($T_{\text{eff}}$, log $g$, [M/H], [$\alpha$/Fe]) for 6508 of 21 551 FEROS archived spectra to be made available in the ESO Archive, including quality flags.
4. The determination of radial velocities ($V_{\text{rad}}$) for 11 963 of 21 551 FEROS archived spectra to be made available in the ESO Archive.

The AMBRE pipeline is currently being used to determine stellar parameters for the UVES and HARPS archived spectra, the next two phases of analysis in the AMBRE Project. The analysis of the FLAMES/GIRAFFE archived spectra is expected to commence in early 2012.

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Table A.1. Description of columns in the table of FEROS stellar parameters delivered to ESO.

| Keyword | Definition | Value range | Null value | Determination |
|---------|------------|-------------|------------|---------------|
| DP_ID  | ESO data set identifier |            |            |               |
| OBJECT | Object designation as read in ORIGFILE |            |            |               |
| TARG_NAME | Target designation as read in ORIGFILE |            |            |               |
| RAJ2000 | Telescope pointing (right ascension, J2000) | deg |            |               |
| DE2000 | Telescope pointing (declination, J2000) | deg |            |               |
| MJD_OBS | Start of observation date | Julian Day |            |               |
| EXPTIME | Total integration time | sec | 0–∞ | NaN |               |
| SNR_FLAG | Signal-to-noise ratio as estimated by the pipeline | C, R | True: detection therefore no analysis carried out, False: no detection therefore analysis carried out |               |
| EXTREME_EMISSION_LINE_FLAG | Detection of extreme emission lines. | T, F | True: detection therefore no analysis carried out, False: no detection therefore analysis carried out |               |
| EMISSION_LINE_FLAG | Detection of some emission lines | T, F | True: some emission lines detected but analysis carried out, False: no detection therefore analysis carried out |               |
| MEANFWHM_LINES_FLAG | Mean FWHM of absorption lines around 4500 Å | 0–0.33 | NaN | FWHM measured from spectral features (mÅ) |               |
| VRAD | Stellar radial velocity | −500 to +500 | NaN | Units = km s⁻¹ |               |
| ERR_VRAD | Error on the radial velocity | 0–∞ | NaN | If σ₁₀ > 10, null value used for all stellar parameters. Units = km s⁻¹ |               |
| VRAD_CCCF_FWHM | FWHM of the CCF between the spectrum and the binary mask | 0–∞ | NaN | Units = km s⁻¹ |               |
| VRAD_FLAG | Quality flag on the radial velocity analysis | 0, 1, 2, 3, 4, 5 | −99 | 0 = Excellent determination ... 5 = Poor determination |               |
| TEFF | Stellar effective temperature (Tₑₑ) | 3000–7625 | NaN | Units = K. Null value used if Tₑₑ is outside accepted parameter limits or if the spectrum is rejected due to quality flags. |               |
| ERR_INT_TEFF | Effective temperature internal error | 0–∞ | NaN | Units = K. Square root of quadrature sum of internal errors |               |
| ERR_EXT_TEFF | Effective temperature external error | 0–∞ | NaN | Units = K. Maximum expected error due to external sources |               |
| LOG_G | Stellar surface gravity (log g) as estimated by the pipeline | 1–4.9 | NaN | Units = dex. Null value used if log g is outside accepted parameter limits or if the spectrum is rejected due to quality flags. |               |
| ERR_INT_LOG_G | Surface gravity internal error | 0–∞ | NaN | Units = dex. Maximum expected error due to external sources |               |
| ERR_EXT_LOG_G | Surface gravity external error | 0–∞ | NaN | Units = dex. Maximum expected error due to external sources |               |
| M_H | Mean metallicity [M/H] as estimated by the pipeline | 0–∞ | NaN | Units = dex. Null value used if [M/H] is outside accepted parameter limits or if the spectrum is rejected due to quality flags. |               |
| ERR_INT_M_H | Mean metallicity internal error | 0–∞ | NaN | Units = dex. Square root of quadrature sum of internal errors |               |
| ERR_EXT_M_H | Mean metallicity external error | 0–∞ | NaN | Units = dex. Maximum expected error due to external sources |               |
| ALPHA | α-elements over iron enrichment ([α/Fe]) as estimated by the pipeline | −0.4–0.4 | NaN | Units = dex. Null value used if [α/Fe] is outside accepted parameter limits or if the spectrum is rejected due to quality flags. |               |
| ERR_INT_ALPHA | α-elements over iron enrichment internal error | 0–∞ | NaN | Units = dex. Square root of quadrature sum of internal errors |               |
| ERR_EXT_ALPHA | α-elements over iron enrichment external error | 0–∞ | NaN | Units = dex. Maximum expected error due to external sources |               |
| CHI2 | log(χ²) of the fit between the observed and the reconstructed synthetic spectrum at the MATISSE parameters | 0–∞ | NaN | Units = dex. Goodness of fit between final normalised and final reconstructed spectra |               |
| CHI2_FLAG | Quality flag on the fit between the observed and the reconstructed synthetic spectrum at the MATISSE parameters | 0, 1, 2 | −99 | 0 = Good fit ... 2 = Poor fit |               |

Notes. ( vòng 8) = Spectral Processing B; ( vòng 9) = Spectral Processing C.
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