Interpreting Stellar Spectra with Unsupervised Domain Adaptation

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Abstract
We discuss how to achieve mapping from large sets of imperfect simulations and observational data with unsupervised domain adaptation. Under the hypothesis that simulated and observed data distributions share a common underlying representation, we show how it is possible to transfer between simulated and observed domains. Driven by an application to interpret stellar spectroscopic sky surveys, we construct the domain transfer pipeline from two adversarial autoencoders on each domains with a disentangling latent space, and a cycle-consistency constraint. We then construct a differentiable pipeline from physical stellar parameters to realistic observed spectra, aided by a supplementary generative surrogate physics emulator network. We further exemplify the potential of the method on the reconstructed spectra quality and to discover new spectral features associated to elemental abundances.

1. Introduction
In an effort to understand the nearby galactic dynamics, and how stellar populations form and evolve, large dedicated sky surveys (Buder et al., 2018; Holtzman et al., 2018) are currently accumulating millions of stellar spectra. Large statistical samples of survey data enable galactic archaeologists to construct tools to study the history of galaxy formation or stellar chemical evolution of the Milky Way and the nearby Universe. From the large collections of spectra, estimates of stellar properties such as temperature, surface gravity, metallicities or elemental abundances are routinely produced from data reduction pipelines. The production of these stellar parameter databases requires computationally demanding simulations (e.g., Kurucz, 1970; Mészáros et al., 2012), and implementing boutique data analysis pipelines (e.g., Pérez et al., 2016; Ting et al., 2019). However simulations of synthetic spectra are naturally limited to the assumptions behind the stellar physics that we can currently model (e.g., Bialek et al., 2019; Shetrone et al., 2015). Imperfect corrections of the instrumental and earth-atmosphere signatures also are often left-over systematics from the data-reduction pipelines. It is thus difficult to have a clear understanding of the systematic uncertainties involved with either the data reduction, the synthetic spectral modelling, or from the algorithms limitations.
Thus, in this paper, we propose an unsupervised domain adaptation method, that learns these corrections without human intervention.

2. Methods
Our work is largely based on the UNIT framework (Liu et al., 2017), briefly summarized here. The key UNIT idea is to share a latent representation between two unpaired domains of interest. In each domain, a Variational AutoEncoder (VAE) is trained to generate fake samples, sharing the common latent space. The generated samples are thereafter challenged with adversarial classifier networks, trained to discriminate between pairs of reconstructed samples originating from either domain.
We set the domains of interest to be a large sample \(X_{\text{synth}}\) simulation data and a large sample \(X_{\text{obs}}\) of observed data. For our stellar physics purpose, domain data are synthetic and observed spectra. Applying UNIT requires technical modifications to adapt to noisy 1D data, but we also made important architectural choices. We summarize the architecture in the diagram in Fig. 1. The main high level differences between the UNIT and our current method are: 1) a disentangled latent representation, and 2) stitching a supplementary generative surrogate physics emulator network to the main UNIT framework. Another minor difference is that we have our autoencoders to be deterministic.

Disentangled Latent Representation Even good physics simulators are capable of only partly representing the complexity of all the observations. Thus, it will always suffer from the gap between theory and practice. We model this gap by splitting the latent space in two: one shared latent space between simulations and observations, and one
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The architecture of the unsupervised domain transfer between simulations and observations. Synthetic spectra are generated on the fly with a physics simulator with input parameters \( \theta \). Both synthetic and observed spectra are encoded into a common latent space \( Z_{\text{shared}} \). The observed spectra are also encoded simultaneously to a split latent space \( Z_{\text{split}} \), coding other observation-specific factors. The rest of the architecture is similar as the UNIT (Liu et al., 2017) framework.

Surrogate Physics Simulator The samples in the synthetic domain are generated by a physical simulator. In our application, the physical simulator is computationally expensive, thus producing a stochastic simulator would be very costly. We replace it with a fast neural-network trained to emulate the physics. We carefully selected physical ranges and sampling strategies of the physical parameter space, within the same “expected” range of parameters covered by the observed samples. We then ran the costly simulations, and trained a network capable of emulating the generated samples from the physical parameters. In our particular case, a simple MLP was enough to ensure 0.1% accuracy at emulating the physical simulator over the full spectral range. Once the differentiable emulator network is trained, the main network can be trained from generating samples on-the-fly.

Besides on-the-fly generation of synthetic spectra, the introduction of the surrogate simulator network allows us to constitute an end-to-end system from interpretable physical parameters to realistic spectra which implicitly learns the non-modelled physics. Being end-to-end, the system is also fully differentiable, and allows one to differentiate the the domain-adapted synthetic spectra with respect to physical parameters. We, in fact, utilize this property to infer physics from spectra.

3. Experiments

We present two case studies to demonstrate how we generate systematic-corrected synthetic models from unlabelled observed spectra through domain adaptation.

3.1. Data

For these two experiments, we use the publicly available APOGEE (Holtzman et al., 2018) survey data, which com-
prises of 250,000 high-resolution \( R = \lambda / \Delta \lambda \sim 22,000 \) infrared spectra, where \( \lambda \) designates wavelength.

- For the synthetic domain, we use the Kurucz ATLAS12/SYNTHE models (see Ting et al., 2019, for details).
- As for the observed domain, we adopt the APOGEE Data Release 14 spectra. The APOGEE spectra are wavelength calibrated to vacuum to be consistent with the Kurucz models. Multiple visits to the same object are pre-processed and co-added. We corrected the spectra for spectral redshift and self-consistently continuum normalized both the Kurucz models and APOGEE spectra using the same routine as laid out in Ting et al. (2019).

The data set provides estimated stellar parameters for both dwarf and giant stars. We further use the samples in the dataset in a random order to make sure spectra from the two domains are not paired.

APOGEE spectra are adopted from half of the objects as the observed domain, and for the other half, we generate synthetic Kurucz spectra via the surrogate emulator assuming only their stellar parameters. We inherit the 25 stellar parameters derived in Ting et al. (2019): \( T_{\text{eff}}, \log g, \) microturbulence \( v_{\text{turb}}, \) additional broadening \( v_{\text{broad}}, \) and 20 elemental abundances, namely, C, N, O, Na, Mg, Al, Si, P, S, K, Ca, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Ge, and the isotopic ratio, C12/C13. This procedure yields a training set of about 100,000 spectra in each domain. We adopt 80,000 spectra as the training set, 10,000 spectra as the validation set. The rest (7,000 spectra) are held out as the test set. All results below are based on the test set.

### 3.2. Stellar Spectra Translation

As a natural application, we investigate the quality of the translated spectra in the observed domain to that originated from the synthetic domain. Once the framework is trained, the generation is fast and allows us to perform a maximum-likelihood fit between the translated and the observed spectra to get the best-fit stellar parameters.

In Fig 3, we compare the average residuals between our transferred best-fit model (bottom) to a best-fit emulated (non-translated) physics model (top). Fitting spectra is a typical process for stellar parameter analysis. The residuals are normalised with the spectra uncertainties provided by the APOGEE survey data release. Well calibrated uncertainties with unbiased best-fit models would show a distribution of \( z \)-score residuals following \( \mathcal{N}(0, 1) \). We show the sample mean \( \bar{m} \) and standard deviation \( s \) in each case. The systematics-learned model clearly outperforms the stellar physics-only model.

![Figure 3. Residuals between physical (Kurucz) best-fit modelled spectra and the APOGEE observed spectra. The top panel shows the difference between the 10,000 APOGEE test spectra and their corresponding best-fit Kurucz models. The bottom panel is a similar comparison but with the transferred spectra by our network, showing smaller mean bias \( \bar{T} \) and standard deviation \( s \) of the residuals.](image)

### 3.3. Spectral Lines

Here, we study the derivatives of the full network, in particular the derivatives with respect to elemental abundances. The derivatives (i.e., flux “response”) of elemental abundances examine which spectral features are associated with a particular element. Unfortunately, for the APOGEE observed spectra, it is impossible to know the ground truth — we simply do not know what might be missing in the Kurucz models. Therefore, as a proof of concept, a mock “observed” data set is drawn from the Kurucz models instead, of which we know the ground truth derivatives. More advanced applications are deferred to future studies.

For the observed domain, instead of adopting the APOGEE spectra, an “observed” data set is synthesized with our physics simulator. We further add noise to these observed mock spectra to mimic the same noise distribution of the APOGEE spectra. As for the synthetic domain, we mask approximately 30% of the spectral features randomly by setting them to the continuum level. In short, in this controlled experiment, we assume two sets of unpaired Kurucz models.

To demonstrate that we can learn actual physics, the goal here is to correctly identify the physical source (i.e., which element) of these masked spectral features, which act as missing information in the synthetic models that we want to learn and recover from the data.

We calculate the difference between the flux derivatives of the synthetic emulator, \( \partial G(Y) / \partial Y \) and the derivatives of the domain transferred spectra, \( \partial T_{\text{synth}\rightarrow\text{obs}}(G(Y)) / \partial Y \), where \( G \) is the surrogate emulator, and \( T_{\text{synth}\rightarrow\text{obs}} \) the domain adaptation network and \( Y \) the physical stellar param-
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Figure 4. Generated Kurucz spectral models but 30% masked of the spectral features in the synthetic domain, and original Kurucz models (with noise) as the observed domain. The top panel shows the Kurucz spectrum with missing spectral features, whereas the second panel shows the systematic-corrected synthetic spectra. The third panel shows the differences between the synthetic and observed spectrum, demonstrating the missing features; the missing features of Mg, Si, Fe, and C are annotated in blue, orange, green, and red respectively. The final panel shows the differences in the derivatives between the synthetic domain and the transferred models.

4. Conclusion

In this study, we illustrated that domain-adaptation framework can have broad implications for simulation-based physics. As a proof of concept, we focused on stellar spectral modeling. Synthetic spectral models are auto-calibrated through a domain adaptation network, thereby reducing the gap between theory and observations. The same network can also predict the elemental sources of unknown spectral features in the synthetic models.

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