

# BHPMF – a hierarchical Bayesian approach to gap-filling and trait prediction for macroecology and functional biogeography

Franziska Schrodt<sup>1,2,3,\*</sup>, Jens Kattge<sup>1,2</sup>, Hanhuai Shan<sup>4,5</sup>, Farideh Fazayeli<sup>4</sup>, Julia Joswig<sup>1</sup>, Arindam Banerjee<sup>4</sup>, Markus Reichstein<sup>1</sup>, Gerhard Bönisch<sup>1</sup>, Sandra Díaz<sup>6</sup>, John Dickie<sup>7</sup>, Andy Gillison<sup>8</sup>, Anuj Karpatne<sup>4</sup>, Sandra Lavorel<sup>9</sup>, Paul Leadley<sup>10</sup>, Christian B. Wirth<sup>2,11</sup>, Ian J. Wright<sup>12</sup>, S. Joseph Wright<sup>13</sup> and Peter B. Reich<sup>3,14</sup>

<sup>1</sup>Max Planck Institute for Biogeochemistry, Hans-Knöll-Strasse 10, 07745 Jena, Germany, <sup>2</sup>German Centre for Integrative Biodiversity Research (iDiv) Halle-Jena-Leipzig, Deutscher Platz 5, 04103 Leipzig, Germany, <sup>3</sup>Department of Forest Resources, University of Minnesota, St Paul, MN 55108, USA, <sup>4</sup>Department of Computer Science and Engineering, University of Minnesota, Twin Cities, USA, 5Microsoft Corporation, One Microsoft Way, Redmond, WA 98052, USA, <sup>6</sup>Instituto Multidisciplinario de Biología Vegetal (IMBIV - CONICET) and Departamento de Diversidad Biológica y Ecología, FCEFyN, Universidad Nacional de Córdoba, CC 495, 5000, Córdoba, Argentina, <sup>7</sup>Royal Botanic Gardens Kew, Wakehurst Place, RH17 6TN, UK, 8Center for Biodiversity Management, Yungaburra 4884, Queensland, Australia, <sup>9</sup>Centre National de la Recherche Scientifique, Grenoble, France, <sup>10</sup>Laboratoire ESE, Université Paris-Sud, UMR 8079 CNRS, UOS, AgroParisTech, 91405 Orsay, France, <sup>11</sup>University of Leipzig, Leipzig, Germany, <sup>12</sup>Department of Biological Sciences, Macquarie University, NSW 2109, Australia, <sup>13</sup>Smithsonian Tropical Research Institute, Apartado 0843-03092, Balboa, Republic of Panama, <sup>14</sup>Hawkesbury Institute for the Environment, University of Western Sydney, Locked Bag 1797, Penrith, NSW 2751 Australia

\*Correspondence: Franziska Schrodt, Max Planck Institute for Biogeochemistry, Hans-Knöll-Strasse 10, 07745 Jena, Germany. E-mail: f.i.schrodt@gmail.com

#### ABSTRACT

**Aim** Functional traits of organisms are key to understanding and predicting biodiversity and ecological change, which motivates continuous collection of traits and their integration into global databases. Such trait matrices are inherently sparse, severely limiting their usefulness for further analyses. On the other hand, traits are characterized by the phylogenetic trait signal, trait–trait correlations and environmental constraints, all of which provide information that could be used to statistically fill gaps. We propose the application of probabilistic models which, for the first time, utilize all three characteristics to fill gaps in trait databases and predict trait values at larger spatial scales.

**Innovation** For this purpose we introduce BHPMF, a hierarchical Bayesian extension of probabilistic matrix factorization (PMF). PMF is a machine learning technique which exploits the correlation structure of sparse matrices to impute missing entries. BHPMF additionally utilizes the taxonomic hierarchy for trait prediction and provides uncertainty estimates for each imputation. In combination with multiple regression against environmental information, BHPMF allows for extrapolation from point measurements to larger spatial scales. We demonstrate the applicability of BHPMF in ecological contexts, using different plant functional trait datasets, also comparing results to taking the species mean and PMF.

**Main conclusions** Sensitivity analyses validate the robustness and accuracy of BHPMF: our method captures the correlation structure of the trait matrix as well as the phylogenetic trait signal – also for extremely sparse trait matrices – and provides a robust measure of confidence in prediction accuracy for each missing entry. The combination of BHPMF with environmental constraints provides a promising concept to extrapolate traits beyond sampled regions, accounting for intraspecific trait variability. We conclude that BHPMF and its derivatives have a high potential to support future trait-based research in macroecology and functional biogeography.

#### Keywords

Bayesian hierarchical model, gap-filling, imputation, machine learning, matrix factorization, PFT, plant functional trait, sparse matrix, spatial extrapolation, TRY.

# INTRODUCTION

Functional trait measurements and analyses have been the focus of numerous studies in recent decades (e.g. Reich *et al.*, 1997; Wright *et al.*, 2004; Chave *et al.*, 2009; Schrodt *et al.*, 2015). However, due to the time and resources required and the sheer number of species on earth, only a small number of species and their traits could be captured to date, especially in tropical and remote ecosystems. In addition, trait data are highly dispersed among numerous datasets and are often not accessible to the wider scientific community. The integration of databases is thus becoming increasingly important for the consolidation of globally dispersed data, as a source of standardized data for further applications, such as model building and validation, and to coordinate future measurement efforts.

Combining trait observations from studies with different research foci produces matrices with substantial gaps. For example, the largest database for plant traits to date, TRY (Kattge *et al.*, 2011), currently contains 215 datasets with 5.6 million trait entries for 1100 traits of 2 million individuals, representing 100,000 plant species. On average only 2 of the 1100 traits represented in TRY are measured for any individual, restricting the usefulness of combined datasets especially for multivariate analyses.

### General characteristics of traits

Some characteristics inherent to functional traits may support statistical gap-filling of sparse trait matrices: a strong phylogenetic trait signal, functional and structural trade-offs between traits and trait–environment relationships.

The phylogenetic trait signal is an effective aid in predicting trait values (e.g. Lovette & Hochachka, 2006; Swenson, 2014): the closer two individuals are related, the more similar their traits will be - with exceptions due to convergent evolution and environmental diversification. This phylogenetic trait signal is reflected in the taxonomic hierarchy: on average individuals within a species are more similar than individuals within genera, families or phylogenetic groups (Kerkhoff et al., 2006; Swenson & Enquist, 2007). In general, most of the trait variance is observed between species (Kattge et al., 2011), differences between species are consistent across spatial scales (Kazakou et al., 2014) and mean trait values at native ranges are appropriate estimates at invaded areas (McMahon, 2002; Ordonez, 2014). Due to these frequently observed patterns, mean trait values of species and even genera are often used to fill gaps in trait matrices (Fried et al., 2012; Cordlandwehr et al., 2013).

Traits possessed by any individual are not independent – functional and structural trade-offs cause correlations between traits (Reich *et al.*, 1997). This has been characterized at the global scale, for example for the leaf and wood economic spectra, respectively (Wright *et al.*, 2004; Chave *et al.*, 2009). Trait variability is also influenced by the environment which, on the one hand, causes large-scale patterns, for example latitudinal gradients of leaf N/P stoichiometry (Reich & Oleksyn, 2004) or changes in the competitive success of *Drosophila* species along a

temperature gradient (Davis *et al.*, 1998), and on the other hand small-scale intraspecific variation, for example in plant traits (Albert *et al.*, 2010; Clark, 2010) and mammalian body size (Diniz-Filho *et al.*, 2007).

### Gap-filling in trait ecology

Many disciplines, for example psychology (e.g. Schafer & Graham, 2002), sociology (e.g. Johnson & Young, 2011) or biogeochemistry (e.g. Moffat et al., 2007; Lasslop et al., 2010) have developed their own set of accepted techniques for predicting missing values tailored to their data structure. In trait ecology, due to the relatively recent advent of large databases, gap-filling methods have been adopted from other disciplines, often without adjusting them to differences in data structure. In general three approaches are used within the community of trait ecologists: (1) deleting rows with missing cases or pair-wise analysis; (2) predicting trait values based on the taxonomic trait signal at species or genus level (mean traits); and (3) predicting values for individual missing entries based on structure, e.g. multiple imputation (MI) (Rubin, 1987; Su et al., 2011) and multivariate imputation using chained equations (MICE) (Rubin, 1987; van Buuren & Groothuis-Oudshoorn, 2011), or/and spatial correlations within the trait matrix (e.g. kriging; Lamsal et al., 2012; Räty & Kangas, 2012). These approaches provide useful imputations in some cases, but whilst deleting missing cases can introduce bias in model parameters (Nakagawa & Freckleton, 2008), taking the 'mean' adds new data points without adding new information, which results in incorrect confidence limits. The techniques mentioned in (3) on the other hand were developed to fill gaps in databases with 10-30% missing entries, but not for a sparsity as high as that observed in combined trait databases such as TRY.

A recent exercise comparing different approaches (amongst them 'mean' and MICE) to filling gaps in a plant trait matrix although at the species rather than the individual level -showed that all methods were indeed only effective up to 30% gaps, that genus mean produced the least reliable results and that including ecological theory, for example by taking into account traittrait correlations, will substantially improve the accuracy of gapfilling approaches (Taugourdeau et al., 2014). Ogle (2013) developed a hierarchical Bayesian model, which could be used for gap-filling of individual trait values based on taxonomic hierarchy and covariates. However, this method only allows for one trait to be predicted at a time, ignoring the correlation structure within a multi-trait matrix. In contrast, some phylogenetic imputation methods, as reviewed recently by Swenson (2014), do allow for multiple trait predictions but only give species or higher-order means and lack any accounting for intraspecific variability. While this may be resolved, for example by adding 'twigs' on the ends of phylogenies with terminal nodes as species (Swenson, pers. comm.) to the authors' knowledge, this has not been implemented yet.

Here we present a new approach – Bayesian hierarchical probabilistic matrix factorization (BHPMF) – which imputes trait values based on the taxonomic hierarchy, structure within

the trait matrix and trait–environment relationships at the same time as providing uncertainty estimates for each single trait prediction. An extension of the method provides a concept for out-of-sample predictions – the extrapolation of point measurements to spatial scales beyond measured areas. We evaluate trait predictions by BHPMF under various aspects and provide perspectives for future developments.

# METHODS

#### Probabilistic matrix factorization

The method we present is a development of probabilistic matrix factorization (PMF) (Salakhutdinov & Mnih, 2008). PMF is a recommendation system developed on the example of predicting users' preferences in movies from other users' movie ratings (Netflix, 2009). Due to its good scalability and predictive accuracy, even for highly sparse datasets, PMF has become a standard technique for imputing missing data (Koren *et al.*, 2009).

PMF models a sparse matrix, such as the TRY database, as the scalar (or inner, dot) product of two latent matrices with the aim of finding a factorization that minimizes the error between predicted and observed data. This technique is closely related to principal components analysis (PCA), which converts a set of observations of correlated variables into a set of values of linearly uncorrelated variables called principal components. Like PCA, PMF is efficient if the original matrix is of low rank, i.e. if the axes of the original matrix provide strong correlations.

In the original case, each column represents a video and each row a user, providing a video ranking (Salakhutdinov & Mnih, 2008). In the case of a trait matrix, videos are replaced by traits and users by entities. In our case, entities are defined as individual plants. They could equally represent the level of an organ or the average over different organisms, for example nitrogen content of a single leaf, motility of a phytoplankton species or the average beak length of several individuals of an avian species. For simplicity, we refer to the individual plant as plant from now on.

In a first step, independent latent vectors are generated over each row (individual, u) and column (trait, v) of the plant × trait matrix  $X \in \mathbb{R}^{N \times M}$  with N rows and M columns (Fig. 1). Any missing entry (n, m) in the original matrix X can be predicted as the inner product of these latent vectors  $x_{nm} = \langle u_n, v_m \rangle$  (Fig. 1).

PMF has been shown to be applicable to biological data, for example in population genetics (Duforet-Frebourg and Blum, 2014). However, our first experiments indicated that for plant traits the prediction accuracy of PMF was insufficient: the accuracy was worse than using species mean trait values to fill the gaps (see Results). We therefore developed an extension of PMF, which accounts for a plants' taxonomic hierarchy to improve prediction accuracy (Bayesian hierarchical PMF; BHPMF). The concept was developed as HPMF in the context of machine learning (Shan *et al.*, 2012). We here introduce the additional application of a Gibbs sampler in order to provide a measure of uncertainty for each imputed trait value (BHPMF) (see 'Gibbs



**Figure 1** Schematic of the probabilistic matrix factorization (PMF) model. *u* denotes the latent vector on the individual plant side, *v* the latent vector on the functional trait side, both of which have a Gaussian normal distribution with a mean of 0 and a variance of  $\sigma^2$ . Each missing entry  $X_{ij}$  can be approximated by the product of the transposed latent vector *U* and the latent vector *V*.

sampler – uncertainty quantified trait prediction'), as well as an extension to facilitate out-of-sample predictions (aHPMF).

# Bayesian hierarchical probabilistic matrix factorization

BHPMF exploits the taxonomic hierarchy of the plant kingdom as a proxy for the phylogenetic trait signal, with the individual plant being nested in species, species in genus, genus in family and family in phylogenetic group (Fig. 2).

BHPMF sequentially performs PMF at the different hierarchical levels, using latent vectors of the neighbouring level ( $\ell$ ) as prior information at the current level. For example, trait data averaged at species level are used to optimize latent vectors at species level, which in turn act as priors for latent vectors at the individual level, which finally are optimized against the observed trait entries in the trait matrix (Fig. 2, equation 1). The sequential approach across the taxonomic hierarchy turned out to be most effective if applied iteratively top down and bottom up.

After transformation of traits to approximate normal distributions and *z*-score transformation, the cost function is developed as the sum of absolute deviations of predictions versus observations for traits (m) of entities (n) (first summand in equation 1) and the sum of absolute deviations of posterior and prior of the latent factors *u* and *v* (second and third summand of equation 1) across all hierarchical levels (L):

$$E = \sum_{\ell=1}^{L} \left\{ \sum_{nm} \delta_{nm}^{(\ell)} \left( \mathbf{x}_{nm}^{(\ell)} - \langle \mathbf{u}_{n}^{(\ell)}, \mathbf{v}_{m}^{(\ell)} \rangle \right)^{2} + \lambda_{u} \sum_{n} \left\| \mathbf{u}_{n}^{(\ell)} - \mathbf{u}_{p(n)}^{(\ell-1)} \right\|_{2}^{2} + \lambda_{v} \sum_{m} \left\| \mathbf{v}_{m}^{(\ell)} - \mathbf{v}_{m}^{(\ell-1)} \right\|_{2}^{2} \right\},$$
(1)

where  $\lambda_u = \sigma^2 / \sigma_u^2$ ,  $\lambda_v = \sigma^2 / \sigma_v^2$  with  $\sigma$  being the standard deviation of the imputations. {·} denotes the set of data at all *L* levels, and  $\delta_{mm}^{(\ell)} = 1$  when the entry (n, m) of  $X^{(\ell)}$  is non-missing and 0 otherwise. Replace  $\ell - 1$  with  $\ell + 1$  and the parent node (p(n)) with the child node (c(n)) for the bottom-up approach. For details see BHPMF in Appendix S1 in Supporting Information.

# Gibbs sampler – uncertainty quantified trait prediction

The parameters of our BHPMF model are optimized against the observations in the matrix using a Gibbs sampler (Fazayeli *et al.*, 2014). The Gibbs sampler is a Markov chain Monte Carlo (MCMC) method, which samples the probability density distributions of model parameters (here the latent vectors) and model predictions (here entries in the plant × trait matrix) (see the grey inset in Fig. 2 and 'Model evaluation', as well as Gibbs sampler results in Appendix S10). The Gibbs sampler-inferred density distributions of trait values are then used to infer the most likely imputation value, as well as the associated uncertainty for each prediction.

# aHPMF – extrapolation from point measurements to regional scales

If BHPMF is stopped at the species level, i.e. without accounting for trait variability specific to individual plants, the residual error represents the intraspecific variability and modelling/ measurement errors. aHPMF focuses on explaining this residual



**Figure 2** Schematic of the Bayesian hierarchical probabilistic matrix factorization (BHPMF) model. *N* denotes the entity (individual plant) side and *U* the corresponding matrix of latent vectors on the row side, *M* the trait side and *V* the corresponding matrix of latent vectors on the column side. *x* denotes an entry in the original plant × trait matrix *S*. The numbers in parentheses show the taxonomic level *L*. For example (4) is the species level whereas (2) is the family level. The grey inset provides a schema for the Gibbs sampler where p(n) is the parent node of *n* in the upper level and c(n) is the set of child nodes *n* in the lower level.

trait variability based on environmental variables, such as soil and climate characteristics of their growth environment in order to enable out-of-sample prediction, i.e. trait predictions for individual plants where the only known factors are species identity and location but no traits have actually been measured on the given individual (Fig. 3).

To capture trait variability that can be attributed to environmental factors, we utilize a hierarchical regression framework, taking into account the taxonomic structure of plants to regularize the regression model. The regression framework takes as independent predictor variables the climatic and soil variables mentioned below at locations with georeferenced trait measurements. The residuals of BHPMF for the 13 plant traits of each observation are considered as the target dependent variables to be predicted. We treat each plant trait independently of every other while regressing them using climate and soil features.

In essence, combining BHPMF with least squares regression over the residuals against environmental factors, we can model the unknown value for species n and trait m in a probabilistic model as

$$k_{nm} = \alpha u_n^{\mathrm{T}} v_m + \beta w^{\mathrm{T}} x + e_{nm} \tag{2}$$

where  $(u_n, v_m)$  are the latent factors, with  $u_n$  having a hierarchical prior from the taxonomy and *x* being the environmental condition with *w* as the regression coefficient.  $e_{nm}$  is the zero mean Gaussian noise. Note that  $\alpha$ ,  $\beta$  are scalar parameters: for BHPMF set ( $\alpha = 1$ ,  $\beta = 0$ ), for aHPMF set ( $\alpha = 1$ ,  $\beta = 1$ ). For details see 'aHPMF' in Appendix S1.

#### Data: traits, climate and soil

We demonstrate the applicability of the methods introduced above on the example of a trait matrix derived from TRY. For details on data standardization see Kattge *et al.* (2011). The spatial distribution of measurement sites and detailed information on the original datasets are shown in Fig. S4.1 (Appendix S4) and Tables S3.1 & S3.2 in Appendix S3.

We extracted a matrix of 13 georeferenced traits consisting of 204,404 trait measurements on 78,300 individuals, spanning 14,320 species, 3793 genera, 358 families and 6 phylogenetic



**Figure 3** Schematic of the advanced hierarchical probabilistic matrix factorization (aHPMF) model. *w* denotes the regression coefficient at different levels of the hierarchy and *Q* the corresponding matrix of latent vectors. The numbers in parentheses shows the taxonomic level *L*.

Global Ecology and Biogeography, © 2015 John Wiley & Sons Ltd

**Table 1** Number of entries, sparsity and root mean square error (RMSE) of species mean (MEAN), probabilistic matrix factorization (PMF), Bayesian hierarchical PMF (BHPMF) and advanced hierarchical PMF (aHPMF) by trait, as well as  $R^2$  values of the regression of imputed versus measured traits. The lowest RMSE and highest  $R^2$  are shown in bold.

Trait	Entries	Sparsity	MEAN		PMF		BHPMF		aHPMF	
			RMSE	$R^2$	RMSE	$R^2$	RMSE	$R^2$	RMSE	$R^2$
SLA	33001	57.9	0.53	0.85	0.88	0.49	0.46	0.88	0.53	0.89
Plant height	16465	79.0	0.47	0.90	0.91	0.40	0.40	0.92	0.44	0.93
Seed mass	7311	90.7	0.37	0.91	0.77	0.40	0.36	0.92	0.36	0.92
LDMC	17331	77.9	0.53	0.83	0.87	0.41	0.43	0.88	0.49	0.89
SSD	9191	88.3	0.51	0.86	1.01	0.19	0.44	0.87	0.51	0.87
Leaf area	39438	49.6	0.50	0.87	0.91	0.41	0.37	0.93	0.41	0.93
Leaf N	26882	65.7	0.67	0.77	0.99	0.28	0.53	0.86	0.59	0.86
Leaf P	11975	84.7	0.72	0.69	0.78	0.62	0.52	0.83	0.62	0.83
Leaf N/area	8180	89.6	0.79	0.65	0.80	0.64	0.51	0.82	0.72	0.82
Leaf fresh mass	11484	85.3	0.47	0.89	0.71	0.71	0.27	0.96	0.39	0.96
Leaf N/P ratio	5999	92.3	0.76	0.69	0.90	0.44	0.49	0.85	0.67	0.84
Leaf C/dry mass	8123	89.6	0.70	0.74	0.884	0.35	0.61	0.61	0.62	0.78
Leaf $\delta^{15}N$	9022	88.5	0.63	0.79	1.02	0.02	0.50	0.87	0.53	0.88
Average	15723	79.9	0.58	0.80	0.88	0.41	0.45	0.88	0.53	0.87

SLA, specific leaf area; LDMC, leaf dry matter content; SSD, stem-specific density; Leaf N and Leaf P, leaf nitrogen and phosphorus concentrations per dry mass, respectively; Leaf N/area, leaf nitrogen concentration per leaf area; Leaf C/dry mass, leaf carbon concentration per dry mass. For definitions of all traits and data sources as well as corresponding references see the Supporting Information (Appendices S2, S3 and S11 respectively).

groups. The sparsity ranged from 49.63% for leaf area to 92.33% for the leaf N to P ratio, with an average sparsity of 79.9% across the trait matrix (Table 1). All traits were log- and z-transformed to improve normality and equalize traits in the cost function during optimization.

For out-of-sample predictions by aHPMF, climate data for mean annual precipitation, mean annual temperature, isothermally and precipitation seasonality were extracted from the WorldClim dataset (Hijmans *et al.*, 2005) and soil texture (sand, silt, clay) and soil organic carbon content in the top soil from the Harmonized World Soil Database v1.2 (FAO *et al.*, 2012).

#### Model evaluation

We ran PMF, BHPMF and aHPMF on the test dataset extracted from TRY. Given the plant × trait matrix, we randomly selected 80% of entries for training (parameter setting), 10% for validation (parameter adjustment by optimizing performance) and 10% for test (independent performance testing after parameter adjustment and learning). This cross-validation improves model fidelity by ensuring that none of the observations are known by the model when performing new predictions. Test entries without training data in the same row would have highly inflated variance. Such cases were prevented by adjusting the splitting accordingly (see 'BHPMF' in Appendix S1).

We evaluated the predicted trait values, using the root mean squared error (RMSE; see equation S13 in Appendix S1) and the correlation coefficient ( $R^2$ ) of *z*-transformed predicted versus observed traits as indicators of overall prediction accuracy. We compared the performance of PMF, BHPMF and aHPMF with a baseline of species mean trait values (MEAN), which uses the overall trait mean of all individual plants within a species for prediction. The effectiveness of capturing the phylogenetic trait

signal was explored by performing BHPMF including increasingly detailed taxonomic information (Fig. 2).

In order to evaluate how well not only predicted versus measured but also trait-trait correlations are preserved in BHPMF, we performed standardized major axis (SMA) regression, the first principal component vector of a correlation matrix fitted through the data centroid (Taskinen & Warton, 2011), on the measured and imputed trait values for some key trait correlations. We also performed a Procrustes analysis with PROTEST (using the R package 'vegan') on a PCA of a subset of the original data versus a PCA based on the estimated values for artificially introduced gaps. Due to its good data cover, we performed this test on the RAINFOR extract from the TRY database (see below). Procrustes is a statistical shape analysis tool (leastsquares orthogonal mapping) which compares two 'superimposed' matrices for overlap, with placement in space and object size being adjustable. We show how uncertainty in trait predictions is accounted for using the Gibbs sampler, comparing prediction confidence (SD) with prediction accuracy (RMSE).

The sensitivity of BHPMF to the fraction of gaps and the effect of using a global database to fill gaps in local or regional datasets were explored using two approaches. First by 'cutting out' a local dataset with high coverage, adding additional gaps (0, 10, 30, 60 and 80%; see Table S8.1 in Appendix S8) and second by using a regional gappy dataset, filling gaps in each of these 'cut-outs' using (1) the global data with information from the local/regional data and (2) just the local/regional data. For our local example, we extracted TRY trait data contributed by the RAINFOR group (Fyllas *et al.*, 2009), which shows a good coverage (sparsity 11%) and covers most of the Amazon (Fig. S8.1 in Appendix S8). For our regional example, we extracted all of the European data (sparsity 72%) (Fig. S8.2 in Appendix S8). For details on methodology please refer to

Appendices S8 & S1. Finally, we provide an example for out-ofsample prediction, extrapolating leaf nitrogen concentration (leaf N) from point measurements to the whole species range of *Acer saccharum* using aHPMF.

Probabilistic matrix factorization and subsequent regression were developed and applied in MATLAB version 2012a (MATLAB, 2012). All other analyses were performed using the statistical platform R version 2.15 (R Core Team, 2014). The maps reported here were produced in ARCMAP 10.1 (ArcGIS Desktop, 2011) and R, using the tree species distribution map of *A. saccharum* from the US Geological Survey (Little, 1971). R scripts to implement BHPMF are available from the authors by request.

### RESULTS

#### Predicted versus observed trait values

To analyse prediction accuracy we compare RMSE and the coefficient of determination  $(R^2)$  for MEAN, PMF, BHPMF and aHPMF averaged across traits and for each trait separately (Table 1; for scatterplots of observed versus predicted for all traits see Fig. S9.1 in Appendix S9). On average, across all traits, BHPMF outperforms PMF, MEAN and aHPMF, with MEAN being significantly more accurate than PMF. This holds after statistical evaluation using a paired *t*-test with *P*-values smaller than 10<sup>-5</sup> at all levels, and is supported by the evaluation of the correlation coefficient  $R^2$  (Table 1). As the RMSE is calculated from z-transformed approximate normal distributions of traits, a RMSE of 0.45 for BHPMF indicates that the average error of predictions is about half a standard deviation, or about 10% of the 95% CI. BHPMF outperforms MEAN and PMF in all traits, while aHPMF shows the same or higher RMSE and higher  $R^2$ than BHPMF for SLA, plant height, leaf dry matter content (LDMC), leaf carbon (C) per dry mass and leaf  $\delta$  <sup>15</sup>N (D15N) (Table 1). The advantage of BHPMF over MEAN is largest for 'physiological traits', such as leaf N and leaf phosphorus concentration (leaf P), and smaller for more 'structural traits' such as seed mass or plant height. The prediction accuracy of BHPMF varies across traits: from RMSE = 0.36 ( $R^2 = 0.92$ ) for seed mass to RMSE = 0.61 ( $R^2 = 0.61$ ) for leaf C content per dry mass. Interestingly, prediction accuracy is not related to the number of entries per trait (Table 1).

#### Accounting for taxonomic hierarchy

The RMSE of MEAN and BHPMF decreases with increasing taxonomic information, indicating that both methods can utilize the hierarchical structure to their advantage (Table S7.1 in Appendix S7). This is also supported by the scatter plot of measured versus predicted specific leaf area (SLA) and leaf N shown in Fig. 4. With increasing taxonomic information, the scatter plot approaches the 1:1 line, i.e. prediction accuracy improves.



**Figure 4** Scatter plots of predicted versus true values for two traits with increasing taxonomic information. Left column, leaf nitrogen concentration per dry weight; right column, specific leaf area. Row 1, no phylogenetic information is used; row 2, only the phylogenetic group is used; row 3, phylogenetic group and family are used; row 4, phylogenetic group, family and genus are used; row 5, phylogenetic group, family, genus and species are used. Predictions are based on Bayesian hierarchical probabilistic matrix factorization. The data are presented in *z*-transformed space. Dotted lines indicate the 1:1 correlation.

#### Trait-trait correlations

Although the presence of strong trait-trait correlations is a prerequisite for the accuracy of BHPMF, such correlations are not provided a priori and are thus not part of the objective function used (equation 1). This turns them into a suitable evaluation measure. An important quality criterion is to what extent the imputed values reflect the observed bivariate correlations, as this is a first indication of the extent to which the overall correlation structure of the n-dimensional trait matrix is maintained by imputation. Our dataset shows on average strong trait-trait correlations, with some exceptions (Fig. S9.5 in Appendix S9). BHPMF and MEAN capture these general trait-trait correlations, but BHPMF reproduces extreme values more accurately than MEAN and is therefore generally better at capturing the shape of the scatter of observed trait data, which is confirmed by more similar SMA R<sup>2</sup> values (Fig. S9.2 in Appendix S9). Looking at the multivariate preservation of trait-trait correlations using Procrustes analysis, our results indicate again that BHPMF does



**Figure 5** Procrustes analysis errors for the first and second principal component axes comparing a princpal components analysis (PCA) performed on the original, gappy RAINFOR data with a PCA performed on the RAINFOR data with artificially introduced gaps being filled using Bayesian hierarchical probabilistic matrix factorization.

not significantly alter the correlation structure of the gap-filled matrix (Fig. 5). The first four principal component axes explain 83.4% and 83.4% of the variability in the dataset for the original and gap-filled data, respectively. None of the principal component axes are significantly different between the gappy and gap-filled data for any of the traits. The traits stem specific density and leaf carbon differ – but not significantly – along the third and fourth axes (see Fig. S9.3 in Appendix S9).

#### **Uncertainty quantified predictions**

The Gibbs sampler provides a probability distribution for every single prediction, as shown in the example of Gibbs samplergenerated density plots of BHPMF-estimated LDMC, leaf N and SLA for *A. saccharum* and *Pinus sylvestris* trees (Fig. S10.1 in Appendix S10). This distribution can be exploited to calculate indices for the best estimate (e.g. mean) and variability (e.g. SD).

This provides an additional means to evaluate our imputation model by comparing prediction confidence (SD) with prediction accuracy (RMSE): when we are confident about our predictions (small SD), these predictions should also be accurate (small RMSE) and vice versa. Figure S10.2 in Appendix S10 shows that this is indeed the case for the whole 13-trait dataset, implying that our model is appropriate. This remains true when we evaluate the Gibbs sampler on each trait separately (Fig. S10.3 in Appendix S10).

#### Gap-filling of regional/local data using BHPMF

As expected, increasing the number of gaps in the RAINFOR dataset generally resulted in a decrease of prediction accuracy (Fig. 6), although less so for structural traits, such as stemspecific density (SSD) and plant height. Reproducibility was high in all cases (Fig. 6). Prediction accuracy of BHPMF was generally approximately equal, no matter whether the regional



Figure 6 Root mean square error (RMSE) of performing Bayesian hierarchical probabilistic matrix factorization (BHPMF) on the RAINFOR cutout (red points in Fig. S8.1 in Appendix S8) for the whole dataset (Total), specific leaf area (SLA), plant height (PlantHt), stem-specific density (SSD), leaf nitrogen (LeafN), leaf phosphorus (LeafP), leaf nitrogen per area (LeafNArea), leaf carbon (LeafC) with increasing number of gaps added to the original RAINFOR data (inherent gappiness of 11%). For the total number of gaps for each trait and added gaps per dataset, see Table S8.1 in Appendix S8. Left- and right-hand sections for each trait (separated by a dotted line) show results when using only the RAINFOR data (RforR) or using all available data (WforR), respectively, to fill the gaps.

(RforR) or global (WforR) datasets were used to fill the gaps (Figs 6 & S8.3 in Appendix S8). This was particularly the case if gap sizes were large (above 10%), whilst RforR outperformed WforR for the imputations of plant height, leaf N, SLA and leaf carbon only where additional gap sizes were small (0 and 10%).

#### **Out-of-sample prediction (aHPMF)**

We illustrate the extension of BHPMF towards out-of-sample prediction with the example of leaf N across the species range of *A. saccharum* (Fig. 7).

F. Schrodt et al.



**Figure 7** Advanced hierarchical probabilistic matrix factorization (aHPMF)-predicted leaf nitrogen concentration (mg  $g^{-1}$ ) of *Acer* saccharum (a), measured values for leaf nitrogen (mg  $g^{-1}$ ) (b), MAT (mean annual temperature) (c), and MAP (mean annual precipitation) (d) across the species range of *A. saccharum*. For a map of the geographic location see Figure S5.1 in Appendix S5.

BHPMF was stopped at species level, followed by a multivariate regression of residuals of predicted versus observed traits at the level of the individual plants against environmental conditions. Trait predictions follow the trends in both measured leaf N and environmental conditions. However, the variability within aHPMF-predicted leaf N is low (18.69–18.80 mg g<sup>-1</sup>) compared with the range of actually measured values (8.65– 28.23 mg g<sup>-1</sup>). While the amplitude of predicted traits for the grid elements is much smaller than the observed ranges, it is notable that *A. saccharum* occurs over a wide range of environmental conditions and grid averages may not reflect variation at local scales. The flat response surface Reich & Oleksyn (2004) found for the correlation between leaf N in the genus *Acer* and mean annual temperature may further support the validity of our results.

#### DISCUSSION

We demonstrate that BHPMF provides accurate and robust uncertainty-quantified trait predictions, even for sparse matrices with up to 80% missing entries. BHPMF outperforms the species MEAN baseline in all aspects: RMSE and  $R^2$  of predicted versus observed entries are smaller and larger, respectively, for each individual trait (Table 1) and trait–trait correlations are better retrieved (Fig. S9.2 in Appendix S9). Prediction accuracy is high (small RMSE) when prediction uncertainty is small (small SD; Fig. S10.2 in Appendix S10), providing a measure of confidence for prediction accuracy. In addition, aHPMF provides a concept to extrapolate from point measurements to species ranges accounting for intraspecific variability (Fig. 7). These results give rise to three major questions: (1) Why does BHPMF provide accurate and robust trait predictions even for sparse trait matrices? (2) Is the prediction accuracy of BHPMF sufficient for applications in ecological contexts? (3) What are the prospects for BHPMF?

#### Why are BHPMF predictions robust and accurate?

BHPMF is a Bayesian hierarchical approach that simultaneously takes the taxonomic trait signal, the correlation structure within the trait matrix and environmental constraints into account to fill gaps in trait matrices. A comparison of BHPMF-predicted trait values with results based on PMF, MEAN and aHPMF indicates the relevance of all three aspects.

PMF has been shown to be accurate even for the imputation of sparse datasets (Koren *et al.*, 2009). However, in the case of our test dataset, PMF performs worse than the baseline MEAN approach. This indicates that our test dataset is not of sufficiently low rank to allow for robust trait predictions based on the correlation structure in the matrix alone.

BHPMF converts PMF into a hierarchical Bayesian model. This has two effects: (1) the higher levels of taxonomy (e.g. phylogenetic group, family) provide almost complete information (very low sparsity), which enables efficient PMF; (2) approximations of the matrices at higher taxonomic level provide excellent prior information for the approximations at lower levels (e.g. genus, species), thus constraining imputations due to the taxonomic signal in trait variation at all levels.

This is achieved without a priori assuming a phylogenetic signal in the trait variability, but rather by opening the door for our model to extract a signal, if it should be there. Thus, in some cases, BHPMF might not put any constraint on the imputed trait values from the taxonomy side, whereas in others, this signal might be stronger, hence the constraint from the taxonomy. The hierarchical taxonomic structure in combination with a consistent phylogenetic trait signal is therefore the key to facilitate *robust* gap-filling by BHPMF – despite high sparsity – at the level of the individual observations. On the other hand BHPMF outperforms the MEAN approach in all aspects. This indicates that the capture by PMF of the correlation structure of the matrix on top of the phylogenetic trait signal is the key to providing *accurate* predictions.

A comparison of BHPMF with aHPMF indicates that explicitly taking environmental constraints into account surprisingly adds little or no improvement to BHPMF: the average  $R^2$  was only higher in 5 out of 13 traits with the RMSE being consistently smaller for BHPMF compared with aHPMF (Table 1). At the individual level, PMF was replaced by multiple regressions against environmental constraints, based on the assumption that the phylogenetic trait signal is mainly observed at the species level whilst environmental constraints add to the trait variability at the individual level.

The fact that BHPMF largely outperforms aHPMF is an indication that, by taking into account trait–trait correlations, PMF seems to be more appropriate than the multiple regression at the individual level. Also, BHPMF implicitly takes environmental information into account via the correlation of measured to predicted traits. This environmental constraint is related to the immediate environment experienced by the observed plant and propagated via trait–trait correlations to the predicted traits, while environmental information incorporated in the multiple regression model by necessity represents only the average over larger scales. Taking environmental information explicitly into account may therefore not substantially improve BHPMF-based gap-filling. Rather, it may be a tool to extrapolate traits from point measurements to the regional scale, i.e. perform out-ofsample predictions.

# Is the prediction accuracy sufficient for applications in ecological contexts?

Our results indicate that BHPMF outperforms the species MEAN baseline in all aspects, but is the prediction accuracy indeed appropriate for use in ecological contexts, and to what extent are the results a special case of our test dataset?

The test dataset is an extract of 78,300 individual observations and the 13 best covered traits from the TRY database, still with 80% of trait entries missing. Our dataset is not typical for common datasets of traits, which generally originate from specific measurement campaigns with only 5–30% missing entries. In these cases, gap-filling may not be such a challenge and common approaches may be sufficient. However, given the experience that BHPMF substantially outperforms both PMF and MEAN, we also expected improved trait predictions in such cases. Indeed, our sensitivity test using 'cut outs' from Europe and data across the Amazon showed that BHPMF was able to impute gaps even in smaller and extremely sparse databases.

When taking advantage of using the global database to fill gaps in a smaller 'cutout', possible confounding factors introduced by global trait variability influencing the sometimes more constrained trait space of a local dataset should be considered. For example in the case of the RAINFOR cut-out, plant height was better predicted using just the local dataset. One likely explanation is that a large amount of new information was included from the global database, amounting to more than 60% more data with a high standard deviation within species. For SSD, on the other hand, only 30% more data were contributed by the global dataset, which were also less plastic within species compared with plant height. Thus, a global dataset may easily introduce 'false' information in the case of plastic traits that are highly influenced by local environmental conditions but provide a lot of valuable additional information in the case of traits that are mainly determined by phylogeny. We recommend, depending on the number and sparsity of the data, using both approaches wherever possible - local and global gap-filling and select the best-fitting approach depending on the target trait and aim.

BHPMF trait prediction uncertainties are well correlated with their error, turning uncertainty into a good surrogate for prediction error (Fig. S10.2 in Appendix S10). This offers the opportunity to select trait predictions with high probability of low errors or weight results in further analyses according to their uncertainty.

#### Perspectives

Statistical approaches to gap-filling and improvement of ecological understanding of species occurrence and dynamics have been criticized for being too complex and including parameters and assumptions without appropriate prior validation (Lavine, 2010). In contrast, BHPMF is a simple and generic model, which has the advantage of incorporating factors known to influence trait expression, such as phylogenetic trait signals, trait-trait correlations and trade-offs without implicitly requiring prior assumptions. This simplicity opens the opportunity to add complexity to improve trait predictions, for example by replacing the taxonomic hierarchy by a hierarchy of phylogenetic distances. BHPMF has been explicitly developed for gap-filling (matrix completion), not for the prediction of trait values in individuals where no trait has ever been measured before, i.e. out-of-sample predictions. Advancing BHPMF to account for phylogenetic distances instead of taxonomy will improve opportunities for efficient out-of-sample predictions, at least in well-resolved clades.

Uncertainty quantified predictions support the validity of the BHPMF approach, where predictions with low uncertainty (small SD) also show high accuracy (small RMSE), and vice versa. Using the Gibbs sampler to get an indication as to where uncertainties are largest merits special attention. Data coverage for traits such as SLA, which are relatively fast and easy to collect, is much better than for other traits. In addition, some traits are highly variable across ecosystems and plant populations. Using the Gibbs sampler, one can statistically define where more sampling effort is probably going to significantly improve our understanding of variation in any specific trait and where the currently available data are sufficient.

The combination of BHPMF with environmental covariates may not necessarily improve gap-filling of trait matrices, but seems to be a promising concept for the extrapolation of traits from point measurements to regional scales. The concept of out-of-sample prediction presented here illustrates a fundamental problem in trait ecology, separating the phylogenetic signal and environmental impact on intraspecific trait variation. The regression of trait values against the environment after gapfilling by BHPMF provides an advantage over using non-gapfilled data of an improved number of data points, which might be essential as trait predictions are often limited by data availability (Verheijen *et al.*, 2013). BHPMF-filled trait matrices could thus become an invaluable tool for the parameterization and validation of global vegetation models.

The TRY database is a typical example of a combination of datasets which have been collected for different purposes. Other disciplines besides plant ecology which have also started to combine their (trait) data are faced with the same problem of sparsity and restrictions in the context of multivariate analyses. BHPMF provides an opportunity to extract the entire information content from such databases combining data from various aspects measured at locations all over the world.

#### Conclusion

BHPMF is a hierarchical Bayesian implementation of probabilistic matrix factorization, which for the first time simultaneously utilizes the taxonomic trait signal, the correlation structure within trait matrices and - implicitly through traitenvironment relationships - environmental constraints for gapfilling of trait matrices. We demonstrate using the example of different plant trait datasets that BHPMF provides robust and accurate predictions even for sparse matrices. In addition, Gibbs sampler-calculated uncertainties indicate how accurate each imputed trait value is, and thus how to treat it in further analyses. The combination of BHPMF with environmental information provides the opportunity to extrapolate from point measurements to continuous trait surfaces across large spatial scales whilst accounting for intraspecific trait variability. We therefore conclude that BHPMF-based gap-filling and trait prediction has a high potential to support future trait-based research in macroecology and functional biogeography.

#### ACKNOWLEDGEMENTS

F.S. was supported by the University of Minnesota, Institute on the Environment via the grant to P.R.: 'Transformational steps in synthesis science', the Max Planck Institute for Biogeochemistry and iDiv, the German Centre for Integrative Biodiversity Research. H.S. was supported by NSF grants IIS-0812183, IIS-0916750, IIS-1029711, IIS-1017647, and NSF CAREER award IIS-0953274. This study has been performed with support by the TRY initiative on plant traits (https://www.try-db.org). TRY is hosted and developed at the Max Planck Institute for Biogeochemistry, with support from DIVERSITAS and iDiv. We thank Ulrich Weber for help with the preparation of soil data and Maarten Braakhekke, Nate Swenson and two anonymous referees for helpful comments and suggestions.

#### REFERENCES

- Albert, C.H., Thuiller, W., Yoccoz, N.G., Soudant, A., Boucher, F., Saccone, P. & Lavorel, S. (2010) Intraspecific functional variability: extent, structure and sources of variation. *Journal* of Ecology, **98**, 604–613.
- ArcGIS Desktop, E. (2011) *Release 10.1*. Environmental Systems Research Institute, Redlands, CA.
- van Buuren, S. & Groothuis-Oudshoorn, K. (2011) mice: multiple imputation by chained equations in r. *Journal of Statistical Software*, **45**, 1–67.
- Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009) Towards a worldwide wood economics spectrum. *Ecology Letters*, **12**, 351–366.
- Clark, J.S. (2010) Individuals and the variation needed for high species diversity in forest trees. *Science*, **327**, 1129– 1132.
- Cordlandwehr, V., Meredith, R., Ozinga, W., Bekker, R., van Groenendael, J. & Bakker, J. (2013) Do plant traits retrieved from a database accurately predict on-site measurements? *Journal of Ecology*, **101**, 662–670.
- Davis, A., Lawton, J., Shorrocks, B. & Jenkinson, L. (1998) Individualistic species responses invalidate simple physiological models of community dynamics under global environmental change. *Journal of Animal Ecology*, **67**, 600–612.
- Diniz-Filho, J.A.F., Bini, L.M., Rodríguez, M.A., Rangel, T.F.L.V.B. & Hawkins, B.A. (2007) Seeing the forest for the trees: partitioning ecological and phylogenetic components of Bergmann's rule in European Carnivora. *Ecography*, **30**, 598– 608.
- Duforet-Frebourg, N. & Blum, M.G.B. (2014) Bayesian matrix factorization for outlier detection: an application in population genetics. The Contribution of Young Researchers to Bayesian Statistics. *Springer Proceedings in Mathematics & Statistics*, **63**, 143–147.
- FAO, IIASA, ISRIC, ISSCAS, JRC (2012) Harmonized World Soil Database (version 1.2). FAO, Rome, Italy and IIASA, Laxenburg, Austria.
- Fazayeli, F., Banerjee, A., Kattge, J., Schrodt, F. & Reich, P. (2014) Uncertainty quantified matrix completion using Bayesian hierarchical matrix factorization. Proceedings of the 13th International Conference on Machine Learning and Applications.
- Fried, G., Kazakou, E. & Gaba, S. (2012) Trajectories of weed communities explained by traits associated with species response to management practices. *Agriculture, Ecosystems and Environment*, **158**, 147–155.
- Fyllas, N.M., Patiño, S., Baker, T.R. *et al.* (2009) Basin-wide variations in foliar properties of Amazonian forest: phylogeny, soils and climate. *Biogeosciences*, **6**, 2677–2708.

- Hijmans, R.J., Cameron, S.E., Parra, J.L., Jones, P.G. & Jarvis, A. (2005) Very high resolution interpolated climate surfaces for global land areas. *International Journal of Climatology*, 25, 1965–1978.
- Johnson, D.R. & Young, R. (2011) Toward best practices in analyzing datasets with missing data: comparisons and recommendations. *Journal of Marriage and Family*, **73**, 926– 945.
- Kattge, J., Díaz, S., Lavorel, S., *et al.* (2011) TRY a global database of plant traits. *Global Change Biology*, **17**, 2905–2935.
- Kazakou, E., Violle, C., Roumet, C., Navas, M.L., Vile, D., Kattge, J. & Garnier, E. (2014) Are trait-based species rankings consistent across datasets and spatial scales? *Journal of Vegetation Science*, 25, 235–237.
- Kerkhoff, A., Fagan, W., Elser, J. & Enquist, B. (2006) Phylogenetic and growth form variation in the scaling of nitrogen and phosphorus in the seed plants. *The American Naturalist*, **168**, E103–E122.
- Koren, Y., Bell, R. & Volinsky, C. (2009) Matrix factorization techniques for recommender systems. IEEE Computer, 42(8), 30–37.
- Lamsal, S., Rizzo, D.M. & Meentemeyer, R.K. (2012) Spatial variation and prediction of forest biomass in a heterogeneous landscape. *Journal of Forestry Research*, **23**, 13–22.
- Lasslop, G., Reichstein, M., Papale, D., Richardson, A., Arneth, A., Barr, A., Stoy, P. & Wohlfahrt, G. (2010) Separation of net ecosystem exchange into assimilation and respiration using a light response curve approach: critical issues and global evaluation. *Global Change Biology*, **16**, 187–208.
- Lavine, M. (2010) Living dangerously with big fancy models. *Ecology*, **91**, 3487.
- Little, E.L.J. (1971) Atlas of United States trees, volume 1, conifers and important hardwoods. Miscellaneous Publication 1146. US Department of Agriculture, Forest Service, Washington, DC.
- Lovette, I.J. & Hochachka, W.M. (2006) Simultaneous effects of phylogenetic niche conservatism and competition on avian community structure. *Ecology*, **87**, 14–28.
- McMahon, R.F. (2002) Evolutionary and physiological adaptations of aquatic invasive animals: *r* selection versus resistance. *Canadian Journal of Fisheries and Aquatic Sciences*, **59**, 1235– 1244.
- MATLAB (2012) Computer software. The MathWorks Inc., Natick, MA.
- Moffat, A., Papale, D., Reichstein, M., Hollinger, D., Richardson,
  A., Barr, A., Beckstein, C., Braswell, B., Churkina, G., Desai, A.,
  Falge, E., Gove, J., Heimann, M., Hui, D., Jarvis, A., Kattge, J.,
  Noormets, A. & Stauch, V. (2007) Comprehensive comparison
  of gap-filling techniques for eddy covariance net carbon
  fluxes. Agricultural and Forest Meteorology, 147, 209–232.
- Nakagawa, S. & Freckleton, R.P. (2008) Missing inaction: the dangers of ignoring missing data. *Trends in Ecology and Evolution*, **23**, 592–596.
- Netflix (2009) Netflix prize. Available at: http://www .netflixprize.com.

- Ogle, K. (2013) Feedback and modularization in a Bayesian metaanalysis of tree traits affecting forest dynamics. *Bayesian Analysis*, **8**, 133–168.
- Ordonez, A. (2014) Functional and phylogenetic similarity of alien plants to co-occurring natives. *Ecology*, **95**, 1191–1202.
- R Core Team (2014) R: a language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria. Available at: http://www.R-project.org/.
- Räty, M. & Kangas, A. (2012) Comparison of k-msn and kriging in local prediction. *Forest Ecology and Management*, 263, 47–56.
- Reich, P.B. & Oleksyn, J. (2004) Global patterns of plant leaf N and P in relation to temperature and latitude. *Proceedings of the National Academy of Sciences USA*, **101**, 11001–11006.
- Reich, P.B., Walters, M.B. & Ellsworth, D.S. (1997) From tropics to tundra: global convergence in plant functioning. *Proceedings of the National Academy of Sciences USA*, **94**, 13730– 13734.
- Rubin, D.B. (1987) *Multiple imputation for nonresponse in surveys*. Wiley and Sons, New York.
- Salakhutdinov, S. & Mnih, A. (2008) Probabilistic matrix factorization. Advances in Neural Information Processing Systems 20 (NIPS 07). Available at: http://www.cs.toronto.edu/ ~rsalakhu/papers/nips07\_pmf.pdf
- Schafer, J.L. & Graham, J.W. (2002) Missing data: our view of state of the art. *Psychological Methods*, **7**, 147–177.
- Schrodt, F., Domingues, T.F., Feldpausch, T.R. *et al.* (2015) Foliar trait contrasts between African forest and savanna trees: genetic versus environmental effects. *Functional Plant Biology*, 42, 63–83.
- Shan, H., Kattge, J., Reich, P.B., Banerjee, A., Schrodt, F. & Reichstein, M. (2012) Gap filling in the plant kingdom – trait prediction using hierarchical probabilistic matrix factorization. Proceedings of the 29th International Conference on Machine Learning (ed. by J. Langford and J. Pineau), pp. 1303–1310. Omnipress, Madison, WI.
- Su, Y.S., Gelman, A., Hill, J. & Yajima, M. (2011) Multiple imputation with diagnostics (mi) in r: opening windows into the black box. *Journal of Statistical Software*, **45**, 55–64.
- Swenson, N. & Enquist, B. (2007) Ecological and evolutionary determinants of a key plant functional trait: wood density and its community-wide variation across latitude and elevation. *American Journal of Botany*, **94**, 451–459.
- Swenson, N.G. (2014) Phylogenetic imputation of plant functional trait databases. *Ecography*, **37**, 105–110.
- Taskinen, S. & Warton, D.I. (2011) Robust estimation and inference for bivariate line-fitting in allometry. *Biometrical Journal*, 53, 652–672.
- Taugourdeau, S., Villerd, J., Plantureux, S., Huguenin-Elie, O. & Amiaud, B. (2014) Filling the gap in functional trait databases: use of ecological hypotheses to replace missing data. *Ecology and Evolution*, **4**, 944–958.
- Verheijen, L.M., Brovkin, V., Aerts, R., Bönisch, G., Cornelissen, J.H.C., Kattge, J., Reich, P.B., Wright, I.J. & van Bodegom, P.M. (2013) Impacts of trait variation through observed

#### F. Schrodt et al.

trait–climate relationships on performance of an earth system model: a conceptual analysis. *Biogeosciences*, **10**, 5497–5515.

Wright, I.J., Reich, P.B., Westoby, M. *et al.* (2004) The worldwide leaf economics spectrum. *Nature*, **428**, 821–827.

Additional references are in the supplementary file at: (weblink)

# SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article at the publisher's web-site.

Appendix S1 Supplementary methods.

Appendix S2 Definition of traits used in this study.

**Appendix S3** References for contributing databases and number of traits contributed.

Appendix S4 Map of TRY measurement sites.

**Appendix S5** Location of *Acer saccharum* range map and soil and climate across the range of *Acer saccharum* 

**Appendix S6** Correlation between traits and environmental variables used in aHPMF.

**Appendix S7** Root mean squared error comparison between MEAN, BHPMF and aHPMF across the taxonomic hierarchy. **Appendix S8** Sensitivity analysis.

**Appendix S9** Bi- and multivariate relationships between traits, measured and imputed trait values.

Appendix S10 Gibbs sampler results.

**Appendix S11** Additional references of data contributors. **Appendix S12** Author contributions.

# BIOSKETCH

**Franziska Schrodt** is a post-doctoral researcher at the Max Planck Institute for Biogeochemistry in Jena and the German Centre for Integrative Biodiversity Research iDIV Leipzig, Jena, Halle. Her work focuses on the application of machine learning and nonlinear statistical tools to the study of biogeochemical patterns. She is especially interested in plant functional trait/biodiversity–environment correlations and the associated implications for ecosystem structure and functioning.

Editor: José Alexandre Diniz-Filho