Scaling methods in ecological modelling

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Abstract
1. Modelling is often confronted with scaling problems, because modelling—directly or indirectly—always implies scaling. This is because models simplify. Simplification usually means aggregation, and aggregation is a scaling process. As scaling cannot be avoided in modelling, it should carefully be addressed and resolved, at least to the degree possible.

2. In this paper, we give an overview of scaling approaches in ecological modelling. We propose to classify scaling approaches into pre-model scaling, in-model scaling and post-model scaling depending on the timing of the scaling relative to the main modelling process.

3. We show general approaches, examples and potential application problems for each category. We suggest that scaling problems might be more widespread than previously thought. These scaling problems are matched with a range of solutions, but often these solutions will have to be adapted and tailored to the specific scaling case.

4. Thus, we recommend that ecologists be aware of scaling challenges especially where models do not explicitly aim at scaling. Developing general test systems for scaling methods may help to broaden and enhance the application of scaling methods in ecology.

KEYWORDS
aggregation, ecological model, levels of scale, meta-model, pattern-process relationship, scale transition theory, scaling-down, scaling-up

1 INTRODUCTION

Scaling is ubiquitous and persistent in ecology. Following the acclaimed concept of pattern and scale (Levin, 1992), patterns and processes at a certain spatial or temporal scale or organizational level emerge from patterns and processes at finer scales or levels and these, in turn, are influenced by the large-scale patterns (Figure 1, Lischke, Löffler, Thornton, & Zimmermann, 2007). Due to this, scaling, that is, changing from one scale to another, is not always straightforward, and sometimes can cause problems due to scale breaks, nonlinearities, feedbacks and heterogeneity in such pattern-process relationships (Snell et al., 2014). Additionally, scaling is sometimes not explicit, and confusion in terminology adds to scaling-related problems. Here, we (1) address scaling terminology; (2) define three categories of scaling approaches, (a) pre-model scaling, (b) in-model scaling and (c) post-model scaling; and (3) explore examples, problems, and, where available, potential solutions in each category. We also elaborate our main claim that modelling is often confronted with scaling challenges, because modelling—directly or indirectly—always implies scaling.

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The term 'scale' has several meanings in different scientific fields. Moreover, scale is not a property of the observed system, but of the observer, so that the observed systems may appear different at different scales, although the system does not change. In the ecological context, scale usually refers to the combination of extent and grain. Extent is the overall area of space or overall duration in time of the study, divided into sub-units with dimensions defined by the grain. Sub-units with small dimensions have a fine grain (also 'fine scale'), and sub-units with large dimensions have a coarse grain (also 'coarse scale'). Ecological scales are usually explicit (spatially and or temporally) and thus can be quantified with SI units [e.g. time in (s) or area in (m²)]. On the contrary, hierarchical levels or entities such as individual, population and community or local, regional and landscape usually cannot be converted, combined or divided in the same way. Unfortunately, hierarchical or organizational levels and entities are also referred to as 'scales'. This is a common source of terminological confusion yielding imprecise descriptions such as 'at the landscape scale' instead of an exact scale quantification in terms of extent and grain or simply 'at the landscape level'. Furthermore, different scientific fields accept different terminologies, which may prevent agreement on a standard definition (Schneider, 2009) and hinder interdisciplinary collaborations. For example, cartographers use 'large scale' to describe maps with small extent (e.g. with a side length of 1 km), whereas ecologists often use 'large scale' to refer to large extents of areas (e.g. with a side length of 1,000 km). Here, we follow the ecologists' approach and use large scale and small scale to refer to large and small extents if not indicated otherwise.

Scaling or process scaling is defined as translating information from one scale to the other. These scales are called source and target scale and differ in grain, extent or both. The direction of the information transfer defines whether it is scaling-up or scaling-down. In ecology, scaling is inevitable because ecological studies typically involve a range of scales. Ecosystem patterns and processes cover a wide range of space and time and often have multiple drivers that act at different scales.

Scaling problems arise basically from the loss of fine-scale information involved in scaling-up the grain or scaling-down the extent and the lack of information involved in scaling-up the extent or scaling-down the grain (Figure 2). They are reinforced by the complexity of the involved systems or models. In scaling-up, problems can be caused by fine-scale heterogeneity, unknown factors, temporally changing relationships, interactions, nonlinearity and feedbacks, the latter three leading to scale dependence (partly reviewed by Lischke et al., 2007; Snell et al., 2014). These problems can originate at the fine scale, at the coarse scale, or in between. Fine-scale problems are due to heterogeneity or interactions. These scaling problems occur if the quantity to scale is not proportional to the ratio of area in spatial scaling or the time step ratio in temporal scaling. This means that proportionality to these scale ratios is only given if the process forming the quantity is homogeneous over the scale range and no interactions distort the proportionality. There are a few cases where this proportionality to scale ratios holds true, for example, for allometric relationships in tropical forests (Chave et al., 2005). Allometric relationships include nonlinear relationships that are still monotonic so that scaling problems may be overcome through the derivation of power equations, with further examples ranging from body-size allometry to species-area curves (Miller, Turner, Smithwick, Dent, & Stanley, 2004). However, most real situations are more complex than this (e.g. Muller-Landau et al., 2006) and span several domains of scale, which are characterized by abrupt changes or critical thresholds in the target process. Scaling via power laws (and in fact via most available techniques) is ill-advised across domains of scale. Here, inappropriate scaling equations would result in an aggregation error. At the coarse scale, unknown factors may play a role that were not considered important enough to be included at the fine scale. Scaling-up without considering these factors may thus lead to erroneous coarse-scale results. The same holds for scaling-down, where, for example, microclimate can have a huge effect not visible at the coarse scale. Between the scales, scaling problems arise from feedbacks between processes or patterns at different levels and from scale-dependent processes. Scale dependence occurs where a process or pattern is significantly different at different scales due to local interactions and feedbacks. Such processes at broader scales emerge from the finer-scale processes. For example, following the gap-dynamics theory, the succession in a natural forest stand emerges from the stochastic dynamics of tree groups on many small patches, which show very different trajectories (Botkin, Janak, & Wallis, 1972).

Modelling is often confronted with scaling challenges, because scaling is an integral part of all modelling; otherwise, every process would have to be described on the spatial, temporal and organizational scale of sub-atomic particles. Here, we broadly define models as purposeful simplifications, including conceptual, statistical,
equation-based, process-based, rule-based, simulation-based or agent-based models. Essentially, models have inputs (data, parameters), do something (calculation, simulation) and then produce an output (values, statistics, patterns). Even in the simple case of a statistical model that is calculating the mean value of a sample, the scale or level of the output differs from the scale or level of the input because the model provides aggregated information about the sample. We call this scaling in the broad sense, whereas scaling in the strict sense is translating a result obtained at one scale to another scale (e.g. Barraquand & Murrell, 2013; Denny & Benedetti-Cecchi, 2012; Englund & Leonardsson, 2008; Johst, Lima, & Berryman, 2013). In summary, models, by definition, simplify and simplification is a scaling process.

Scaling can, thus, not be avoided in modelling and should carefully be addressed, at least to the degree possible. Scaling may occur at different stages during the modelling process (Figure 1). We have pointed out that models in themselves are scaling procedures. Such 'in-model scaling' can also be done on purpose, for example, for a controlled simplification (Lischke, Löfler, & Fischlin, 1998), or for including finer-scale processes, for example, hourly transpiration in a vegetation model acting on a yearly time step (Speich, Zappa, & Lischke, 2018; Speich, Zappa, Scherstjanoi, & Lischke, 2019). However, scaling can also happen before and after the actual modelling process. For example, when models are parametrized with empirical data, the parameter values usually are the result of statistical models of these data such as slopes and intercepts of a linear regression model. Since statistical analysis can be an aggregation and thus a scaling process, we suggest to call this 'pre-model scaling'. Pre-model scaling includes any scaling that is done prior to running a model, including the data collection process. It can involve the use of other models, such as statistical models in the example, or climatic relationships in statistical climate downscaling (Karger et al., 2017). 'Post-model scaling' then captures scaling based on model outputs, that is, scaling in the strict sense. Neither pre-model nor post-model scaling are essential, because a model can do without empirical input data if it addresses theoretical questions or without outputs scaled up in space or time if outputs at the spatial and temporal scales of the model are sufficient to answer the model question. We think that the classification into pre-model, in-model and post-model scaling methods facilitates revealing hidden scaling procedures and disentangling some of the confusion and ambiguity in current treatments of scaling methods. Where the application of this classification reveals hidden scaling methods, it becomes apparent that scaling methods and thus scaling problems might be more widespread than previously thought. We assigned some commonly used scaling techniques to those new categories, describing the purpose of the scaling methods and whether they accept heterogeneous input, visualizing their main principle and giving at least one example application from the literature for each of them (Table 1).

In the following, we will specify scaling problems and examples for approaches from pre-model scaling via in-model scaling to post-model scaling. Finally, we will make some practical
**TABLE 1** Overview of scaling methods in the categories pre-model scaling, in-model scaling and post-model scaling (see Figure 1), together with their main purpose and their type of input and including a visualization of the principle of the method and one or more example studies addressing the method (see Supporting Information for enlarged versions of the figures)

| Scaling method                  | Purpose                                                                 | Input | Visualization | Examples                                                                                       |
|---------------------------------|--------------------------------------------------------------------------|-------|---------------|------------------------------------------------------------------------------------------------|
| No scaling                      | To create a model with input and output at the same scale                | Het   |               | Only as ‘null’ reference  |
| Pre-model scaling               |                                                                          |       |               |                                                                                               |
| Statistical models              | To aggregate values to make them usable as model inputs                  | n.a.  |               | Simple averaging, linear and nonlinear regression modelling, Bayesian analysis, generalized linear mixed effects models |
| Coarsening gridded maps         | To aggregate spatial information                                         | Het   |               | Majority rule, simple averaging, area-weighted mean, kriging                                 |
| Refining gridded maps           | To derive finer-scale spatial patterns                                   | Het   |               | Interpolation, statistical scaling-down, with(out) additional information (Karger et al., 2017) |
| In-model scaling                |                                                                          |       |               |                                                                                               |
| Process refinement              | To simulate certain processes on a finer resolution                      |       |               | Smith, Prentice, and Sykes (2008)                                                            |
| State variable refinement       | To apply aggregated model to finer-scale state variables                 | Het   |               | Hickler et al. (2012)                                                                         |
| Effective parameters            | To derive large-scale parameters directly from small-scale parameters to use as input to small-scale model | Non-het |              | Wu, Jones, Li, and Loucks (2006)                                                             |
| Brute force—classic             | To run copies of the small-scale model in each cell of a large-cell grid | Het   |               | Peters et al. (2004)                                                                          |
| Brute force—with interactions   | To add a model of interactions between cells                              | Het   |               | Cipriotti, Wiegand, Pütz, Bartolini, and Paruelo (2016)                                      |
| Brute force—with super-individuals | To aggregate individuals of similar age or size into super-individuals that are used as entities in large-scale model | Het   |               | Xavier, Grose, and Whetton (2007)                                                            |
| Scaling method                  | Purpose                                                                 | Input | Visualization | Examples                                                                                       |
|--------------------------------|-------------------------------------------------------------------------|-------|----------------|-----------------------------------------------------------------------------------------------|
| Representative cells           | To run the simulation only once for each cell representing many cells with same conditions | Het   |                | Lehsten, Mischurow, Lindström, Lehsten, and Lischke (2019) and Nabel (2015)                   |
| Scale transition theory        | To mathematically or heuristically derive larger-scale processes from small-scale processes, interactions and heterogeneity, for example, to use nonlinear averaging to scale-up nonlinear population growth | Het   |                | Bolker and Pacala (1997) and Melbourne and Chesson (2005)                                      |
| Meta-model—classic            | To fit a meta-model (dynamic, statistical, or neural network) to the results of running a small-scale model at a range of relevant conditions and only applying the simpler meta-model to obtain large-scale results; in contrast to analytical approximations (see post-model scaling), the small-scale model is only of interest as basis for scaling-up | Het   |                | Acevedo, Ablan, Urban, and Pamarti (2001), Acevedo, Urban, and Ablan (1995), Acevedo, Urban, and Shugart (1996), Meier, Lischke, Schmatz, and Zimmermann (2012), Rammer and Seidl (2019) and Urban, Acevedo, and Garman (1999) |
| Meta-model—with interactions  | To add an interaction model to the meta-model                              | Het   |                | ZelStage model (Urban et al., 1999)                                                            |
| Post-model scaling             |                                                                         |       |                |                                                                                               |
| Approximation by analytical model in general | To approximate small-scale model results by an analytical model and apply only analytical model at large extent; similar to meta-models, but applied to models whose original aim did not include scaling-up | Non-het |                | Moorcroft, Hurtt, and Pacala (2001)                                                            |
| Approximation by analytical model for individual-based models | To approximate large-scale results of individual-based models with analytical model and only apply analytical model in other large-scale scenarios | Het   |                | Fahse, Wissel, and Grimm (1998) and Johst et al. (2013)                                        |
| Spatial statistics             | To combine scale-dependent pair correlation functions and species abundance distributions to predict species numbers across scales via species–area curves | Non-het |                | Azaele et al. (2015)                                                                          |

*a Type of model input: Het—spatially heterogeneous, Non-het—no spatially heterogeneous input possible.
*b Visualizations show only examples. For simplicity, visualizations refer to spatial scaling only. This can be transferred analogously to temporal and hierarchical scaling.
*c This is only theoretical, because in practice, according to our claim in this paper, all modelling implies scaling.
recommendations to guide the application of scaling approaches (see also Figure 3) and point out theoretical avenues towards minimizing some related problems in scaling.

2 | SCALING APPROACHES ALONG THE MODEL CHAIN

2.1 | Pre-model scaling

Pre-model scaling refers to aggregation or disaggregation of values to make them usable as model inputs. This (dis)aggregation is not necessarily related to a scaling procedure across space or time (note that if parameters were aggregated across spatial or temporal scales, the result would be called an effective parameter and the procedure would fall into the category in-model scaling). Pre-model scaling often involves statistical modelling if parameter values are determined by parameterization from detailed empirical data instead of by calibration with coarse empirical data (e.g. via pattern-oriented modelling; Grimm et al., 2005). Parameters of a model usually have a standard or reference value that needs to be extracted from empirical data. These data can be obtained from literature sources, own measurements, expert knowledge or their combination. Even if only one source is used, the parameter value is in almost all cases the result of some statistical calculation to aggregate over a sample. This ranges from simple averages and estimates of linear regression models or generalized linear mixed effects models to results of more advanced techniques such as Bayesian analysis. All these approaches do some kind of aggregation, that is, scaling-up. This is more obvious where values from several sources are combined into one model parameter value, again using statistics and thus scaling-up. Fortunately, pre-model scaling problems such as nonlinear
relationships, heterogeneity in the form of non-normal distributions, and interactions can be addressed with a broad array of statistical methods including nonlinear regression models, generalized linear models or mixed models (Crawley, 2012).

Model inputs such as spatial maps or time series can also be subject to pre-model scaling. The spatial resolution of input maps can significantly and differentially affect model behaviour. For instance, scaling-up is always associated with a loss of information or aggregation error, for example with respect to landscape homogenization, when animal dispersal and animal population dynamics are simulated (Bocedi, Pe'er, Heikkinnen, Matsinos, & Travis, 2012). This can lead to a bias depending on the aim of the model. Thus, the spatial resolution of input maps needs to be adapted to the requirements of the model by scaling-up and scaling-down. Common coarsening methods for aggregation of information in gridded maps apply the majority rule (use the most common state of fine cells for the aggregated cell), simple averaging (use the average of the fine-scale information), area-weighted mean (weigh the average by the area a state is covering inside the aggregated cell) and kriging (fit a spatial function of the information; Chiles & Delfiner, 1999). Scaling-down in contrast increases the information content by generating finer-scale patterns based on interpolating large-scale patterns, sometimes using additional information and models. One example is climate regionalization, where climatic variables are scaled down (Ekstroem, Grose, & Whetton, 2015). This includes that temperatures are interpolated and refined by fine-scale elevation data, and precipitation patterns by additional data about prevailing wind fields resulting in luv and lee effects of mountain ranges (Karger et al., 2017). Similarly, temporal scaling-down adds values between given values, by more or less explicit assumptions about the underlying process. For example, daily temperature data are often created by interpolation based on linear or sine-wave functions between monthly values (Lischke, Loffler, & Fischlin, 1997).

2.2 In-model scaling

Modelling is a robust way to bridge scale discrepancies and to include multiple spatial and temporal interactions when analysing complex ecological phenomena (Urban, 2005). This is reflected in the broad range of model-based scaling-up methods. Reviews of ecological model-based scaling methods are available for general ecology (Rastetter, Aber, Peters, Ojima, & Burke, 2003; Urban, 2005; Urban et al., 1999; Wu et al., 2006), species distribution models (Miller et al., 2004), landscape research (Lischke et al., 2007), forest ecosystem management (Seidl et al., 2013) and dynamic vegetation models (Snell et al., 2014); thus, we only give a short overview here (see also Table 1). Scaling-down refers to the refinement of processes within the model, sometimes by coupling a submodel for some processes to the original model. For example, the dynamic global vegetation model LPJ, which was based on representative (average) individuals of plant functional types, was merged with an individual-based model of tree species, significantly improving the simulated vegetation dynamics (Smith et al., 2008).

The purposes of in-model scaling-up are deriving simpler models for (a) computationally more efficient simulations (in terms of memory and time), (b) mathematical tractability and (c) detecting and formalizing emergent dynamics. A few in-model scaling-up methods will be reviewed in the following (see also Table 1).

Averaged or ‘effective’ parameters (Wu et al., 2006) can sometimes be derived from the small scale for the large scale and directly used as input to the original small-scale model to generate large-scale output (Cipriotti et al., 2016). This only works in the absence of scale-dependent processes or interactions at the large scale.

Brute force approaches work with copies of a small-scale model that are run in each cell of a large-scale grid using the local conditions of the respective cell as inputs (Cipriotti et al., 2016; Peters et al., 2004). To work trustworthily, they also require the absence of additional large-scale processes and large-scale interactions. A brute force approach can be combined with interactions at the large scale to broaden its applicability. However, this comes at the cost of huge computing power requirements and is often not feasible. In individual-based modelling, computing power constraints of brute force approaches can be alleviated using super-individuals or representative individuals. In individual-based models (and some complex physiological models), population- or community-level dynamics are not imposed, but emerge from individual-level interactions (Grimm & Railsback, 2005). Aggregating such individuals into super-individuals that share similar properties such as age or size is very useful to save computing time (Scheffer, Baveco, DeAngelis, Rose, & van Nes, 1995). For example, Xavier et al. (2007) successfully implemented a multiscale individual-based model of microbial and bioconversion dynamics with bacterial groups clustered into super-individuals called granules. Representative individuals are also typical of forest gap models (Bugmann, 2001). Similarly, grid cells can be clustered into similar representative cells. For example, the forest landscape models LANDIS II (Scheller et al., 2007) and iLand (Seidl, Rammer, Scheller, & Spies, 2012) work with ecoregions that are considered homogeneous over the entire simulation time. In their dynamic global vegetation model, Lehsten et al. (2019) simulate local forest dynamics only along transects, interpolate between them, but simulate dispersal in all fine-scale grid cells. Nabel (2015) dynamically assign cells of similar species composition to representative cells in which local forest dynamics are simulated and then redistributed into space. However, the aggregation bias associated to a super-individual or representative cell approach arising from the clustering of similar but not necessarily identical entities can advocate against this approach and in favour of non-aggregated approaches (Parry & Evans, 2008).

A very elegant but also very difficult way of scaling-up is to derive the broad-scale process functions of the model given the scaling-up of the state variables, ideally mathematically or partly heuristically (see Figure 1 in Lischke et al., 2007). For example, Lischke et al. (1998) aggregated tree populations of a forest gap model into height classes. The main heuristic assumption was that a crucial element of the gap models—namely the between-gap heterogeneity...
resulting from the small-scale demographic stochasticity in each gap—can be mimicked by a random spatial distribution of the trees. Based on this assumption, it was possible to mathematically derive the main process functions.

Scale transition theory also provides analytical methods for such a mathematical scaling-up of population dynamics based on nonlinear population growth and spatial or temporal variation in population densities as well as abiotic factors (Chesson, Donahue, Melbourne, & Sears, 2005). These methods are based on a mathematical manipulation, the Laplace transform, and draw on the phenomenon of nonlinear averaging (Chesson, 1978, 1981). It has, for example, been applied to scale-up the population dynamics of caddisflies from riffles to whole streams (Melbourne & Chesson, 2005). Scale transition theory has been elaborated and widely extended over the past decades (Benedetti-Cecchi et al., 2012; Chesson, 1998, 2009, 2012; Chesson et al., 2005; Melbourne & Chesson, 2006).

Meta-models are doubly simplified, because they are models of models (Urban, 2005; reviewed in Pietzsch et al., 2020). To obtain a meta-model, first a small-scale simulation model is run under all sets of conditions that are relevant for large-scale model analysis. Then, the meta-model is fitted to the results of these simulations given the respective conditions. The meta-model is then used to make predictions at the large scale that are either directly used or entered into a simplified large-scale simulation model. In contrast to the method of analytical approximations (see Section 2.3), in meta-modelling, the small-scale model is not of interest in itself, but only as a basis for scaling-up with the meta-model. Typically, a meta-model is a statistical model such as a multiple regression, mixed effects or generalized linear model (Meier et al., 2012). More recent meta-modelling techniques include neural networks. Rammer and Seidl (2019) fitted a neural network to the IBM iLand, run in many combinations of environmental conditions. The meta-model then gave the trajectories of the forest dynamics. In forest gap modelling, Markov models have also been used as meta-models (Acevedo et al., 1995, 1996) and can under certain conditions be generated automatically (Acevedo et al., 2001). Urban et al. (1999) presented a semi-Markov meta-model based on a cellular automaton. These meta-models interpolate over the whole parameter space given by the simulated conditions. In other cases, meta-models consist of libraries or lookup tables that contain results of small-scale simulations only for the simulated sets of conditions without interpolation (Cipriotti et al., 2016; Seidl et al., 2012). These libraries or lookup tables are then used in large-scale simulations to save ad hoc simulation time. If meta-models are used to avoid running many costly simulations of the original model, then they can be ‘black boxes’, as in the case of the neural networks. If the meta-model structure is based on ecological theory, such as the Lotka-Volterra competition model, meta-modelling could additionally give ecological insights by relating the processes and parameters of the original complex model to that of a mathematically well studied simpler theoretical model.

2.3 | Post-model scaling

Post-model scaling addresses all scaling procedures that are applied to the output of models. Post-model scaling procedures comprise analytical approximations as well as statistical methods such as those described in the section on pre-model scaling but also advanced spatial statistical approaches such as the pair correlation function. Usually, simulation results are presented at different degrees of aggregation, for example, as total biomass versus environmental gradients or as biodiversity in time and space, all of them representing a simple post-model scaling-up. In contrast, if representative-cell approaches have been used, the results are then redistributed in space, that is, scaled-down again.

Analytical approximations are elegant tools to simplify scaling-up approaches by fitting an analytical model to the large-scale output of a small-scale simulation model (Grimm & Railsback, 2005, chapter 11). If a suitable analytical expression can be found, it is a huge advantage, because analytical models are computationally much simpler than corresponding simulation models (Tietjen & Huth, 2006). Hence, it is not a scaling method per se, but a method to improve computing times during scaling. In contrast to meta-modelling (see Section 2.2), the analytical approximation is not necessary to fulfil the aim of the original model, but contributes an additional post-hoc scaling-up component. For instance, Moorcroft et al. (2001) used partial differential equations to approximate the results of a stochastic individual-based simulation model of ecosystem dynamics. Their small-scale simulation model operates at the scale of 15 m × 15 m, representing a forest gap of a few individuals. With their analytical approximation of time-dependent model behaviour, they were able to scale up to one-degree grid cells. Analytical approximations can also help to gain insight from scaling-up between individuals and populations. Fahse et al. (1998) were able to show that the results of their individual-based simulation model on bird flocking corresponded well with the analytical model of logistic growth. This was perceived as very beneficial during model analysis. However, applications of analytical approximations are not always unequivocal. In their analytical approximations of the population-level results of an individual-based competition model, Johst et al. (2013) found that the same type of resource competition was best described by different functional forms depending on the type of exogenous fluctuation, that is, food availability or weather. Fitting analytical models should thus not only follow statistical criteria, but be tailored to the ecological case study.

Advanced spatial statistics can also be used for scaling-up simulation results. Azaele et al. (2015) applied the pair correlation function to scale spatial species abundance distributions from a 1 m radius to a 400 m radius. The pair correlation function describes spatial point patterns (e.g. of study subject locations) across scales by determining how the density of points varies depending on the distance to reference points. However, predicting species abundance distributions from pair correlation functions relies on restrictive assumptions that exclude scale dependence in the functional form of the species abundance distribution, require the random environmental
variability in the region to be moderate and prohibit spatial heterogeneity. This strongly restricts the applicability of the pair correlation function for scaling-up.

3 | CONCLUSIONS AND FUTURE DIRECTIONS

This review takes the perspective that scaling is inherent to modelling and elaborates how the scaling approaches that are available can be classified into pre-model, in-model and post-model scaling methods. This implies that scaling and the associated problems are probably more widespread than previously thought, since they cover so many different areas of modelling. Thus, we recommend that ecologists be aware of scaling problems especially where models do not explicitly aim at scaling.

In terms of practical recommendations, we have derived a rough identification key for scaling approaches based on the approaches reviewed here (Figure 3). Given the aims and conditions of an ecological modelling study, the key provides decision support which scaling approach may be most suitable in which cases. However, the small number of aims and conditions covered by the key also reveals that the available scaling approaches do not often provide generally applicable solutions. Often, scaling approaches will have to be tailored to the specific case or new methods will have to be developed.

Future directions for theoretical and methodical advancement include four steps: First, the potential of existing scaling approaches should be more fully exploited. For example, in meta-modelling, many more statistical techniques may be explored. Second, underexplored scaling techniques such as Bayesian approaches (Clark, 2003; Ellison, 2004; Wikle, Berliner, & Cressie, 1998) and spatial statistics (Azaele et al., 2015; Illian, Penttinen, Stoyan, & Stoyan, 2008) should be further developed. For example, a Bayesian approach has already been successfully applied to scale-up from the individual level to the population level in an animal movement model (Hooten, Buderman, Brost, Hanks, & Ivan, 2016). Third, gaps in the coverage of current approaches (Figure 3), for example, dealing with scale dependence, should be filled by developing entirely new scaling approaches. Finally and importantly, scaling-up results need to be evaluated (Moorcroft et al., 2001; Schindler, 1998). Such an evaluation requires knowledge of the true pattern for comparison, at least for parts of the target scale, and should ideally control for environmental variability (Azaele et al., 2015), which is difficult to achieve. One avenue to address this evaluation challenge, for example, in vegetation modelling is to take advantage of data from mega-plots such as the ForestGEO plots with extents of more than 25 ha or of new methods in remote sensing that provide data at almost any desired grain and extent and compare them with scaling results. Another avenue is to design virtual test systems where the true patterns are known at any scale so that the relative performance of different scaling methods can be evaluated (e.g. case study 2 in Scaiini, Fritsch, Scherer, & Simpkins, 2018). Following these directions may help to improve current scaling solutions and diversify the portfolio of available scaling methods in the future.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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