



Potential impacts of climate change on *Welwitschia mirabilis* populations in the Namib Desert, southern Africa

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Abstract: Climate change is threatening natural ecosystems in the Earth, and arid regions of southern Africa are particularly exposed to further drying. *Welwitschia mirabilis* Hook. (Welwitschiaceae) is an unusual gymnosperm tree that is recognized as an icon of the Namib Desert, southern Africa. Many aspects of its biology were investigated in the past, with a special emphasis for its physiology and adaptations, but nothing is known about its potential sensitivity to current climate changes. In this study, we adopted an approach based on distribution data for *W. mirabilis* and ecological niche models for clarifying the species-climate interactions and for predicting the potential impacts of climate change on *W. mirabilis* populations in three well-separated sub-ranges (northern, southern and central) in northwestern Namibia, southern Africa. We evidenced that the populations occurring in the northern sub-range have peculiar climatic exigencies compared with those in the central and southern sub-ranges and are particularly exposed to the impact of climate change, which will consist of a substantial increase in temperature across the region. These impacts could be represented by demographic changes that should be detected and monitored detailedly to plan efficient measures for managing populations of this important species on the long-term scale.

Keywords: global warming; range fragmentation; climatic suitability; *Welwitschia mirabilis*; ecological niche model; Namib Desert

Citation: Pierluigi BOMBI. 2018. Potential impacts of climate change on *Welwitschia mirabilis* populations in the Namib Desert, southern Africa. Journal of Arid Land, 10(5): 663–672. <https://doi.org/10.1007/s40333-018-0067-1>

1 Introduction

Global climate change is recognized as one of the major threats to biodiversity and natural ecosystems. Extinction events, range shifts, and density variations at population and species levels have been widely documented in cold, temperate, and tropical ecosystems (Walther et al., 2002; Parmesan and Yohe, 2003; Cristofari et al., 2018). In addition, further pressures on natural populations, driving to changes in species diversity and ecosystem functioning and service provision are expected for the years to come (Hole et al., 2009; Moritz and Agudo, 2013; Ge et al., 2015). Afro-tropical species and ecosystems are expected to be particularly sensitive to climate change and to face substantial alterations (Thuiller et al., 2006a; Peterson et al., 2014; Blach-Overgaard et al., 2015; Midgley and Bond, 2015). For this reason, demographic studies

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Received 2017-08-30; revised 2018-03-09; accepted 2018-04-05

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have been carried out on some species in arid regions of southern Africa for detecting climate change impacts (Foden et al., 2007; Guo et al., 2011).

Ecological niche models have been extensively used to project the likely impacts of future climate change for different time slices. In very general terms, the ecological niche models are based on the definition of the species-environment interactions and their spatial explicit projection in the future climate (Guisan and Zimmermann, 2000). In recent years, ecological niche models underwent deep improvements both in the theory and methodology (Guisan and Thuiller, 2005). A large spectrum of modeling approaches and techniques have been developed for dealing with different data types and aims (Elith and Leathwick, 2009; Guisan et al., 2013). These approaches have been used for predicting species distribution in a wide range of biota and for targeting initiatives, allowing to improve conservation strategies (Bakkenes et al., 2002; Iverson and Prasad, 2002; Thuiller et al., 2004, 2006b).

Arid regions of southern Africa host a high percentage of endemic species and a variety of remarkable taxa (Maggs et al., 1998). *Welwitschia mirabilis* Hook. is an endemic, unique species of the taxonomic family Welwitschiaceae, and also an unusual member of gymnosperms in the Namib Desert, southern Africa. The distribution of *W. mirabilis* is restricted to the central and northern Namib Desert, extending from the Kuiseb River in Namibia to the Nicolau River, north of Namibe, in Angola (Kers, 1967; Giess, 1969). It occurs in isolated populations, with highly variable density (Henschel and Seely, 2000). Although a great number of studies have dealt with the morphology, development, ecophysiology, ecology, and distribution of *W. mirabilis*, many important aspects of the biology of this iconic species still need further understanding (Henschel and Seely, 2000; Valverde et al., 2016; Krüger et al., 2017). To the best of our knowledge, no assessment of the potential impacts of climate change on this important species is available to forecast the variation in environmental suitability in the future.

The aim of this study is to fill this gap, predicting how the climatic conditions in the Namib Desert will satisfy environmental exigencies of *W. mirabilis* populations in the future. We adopted an approach based on ecological niche models for clarifying the climatic components of the ecological niche of *W. mirabilis* populations and for evaluating whether and where such climatic exigencies will be satisfied across its current geographic range. More specifically, we combined distribution data with current climate surfaces for exploring the species-climate relationship and for modeling the climate suitability of *W. mirabilis* populations in the Namib Desert under the current and future conditions. Furthermore, we projected a defined relationship into a wide spectrum of climate change scenarios for estimating the extent to which the climatic suitability will change Namib Desert in the future. Our results are important to understand more about the ecology of this unusual plant (i.e., *W. mirabilis*), for identifying which populations could suffer the worst impacts of climate change, and for improving the management strategies of *W. mirabilis* fields.

2 Study area and data collection

Distribution data for *W. mirabilis* were downloaded from the 'Digital Atlas of Namibia' (Bubenzer et al., 2004), which provides maps compiled by the 'National Herbarium of Namibia' based on the databases of the 'National Botanical Research Institute of Namibia' and published in the 'Atlas of Namibia' (Mendelsohn et al., 2002) for the 'Ministry of Environment and Tourism' (Fig. 1). These maps are provided as polygon shape files for a selection of six plant species that are important at the country scale. Polygons describe the species distributions in terms of their extent of occurrence (IUCN, 2012), encompassing all known localities of presence. We defined our study area as a bounding box, which is 2/3 larger than the latitudinal and longitudinal extent of the data polygons.

The available data could describe *W. mirabilis* as spread in three well-separated sub-ranges in the Namib Desert, southern Africa (Fig. 1). The northern sub-range is a circular area in central Kaokoland at the border between Namib Desert, Nama Karoo, and savanna biomes. The central sub-range is a large area in Damaraland and Swakopmund districts laying over Namib Desert and Nama Karoo. The southern sub-range is an elongated area in the central Namib Desert,

Swakopmund district, southwestern Africa. The three areas are roughly 100 km away from each other and are separated by unoccupied zones.

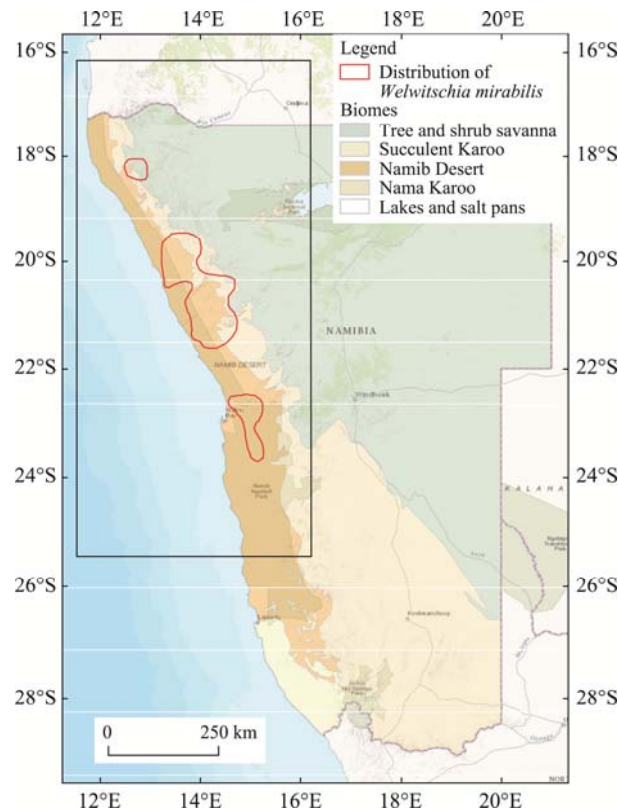


Fig. 1 Distribution of *Welwitschia mirabilis* in the Namib Desert, southern Africa

3 Materials and methods

Climate data for the study area were downloaded from the 'WorldClim databank' (version 1.4; www.worldclim.org) (Hijmans et al., 2005a), which consists of 19 bioclimatic variables with global coverage. These variables were derived from the monthly temperature and precipitation values, using measures for the period 1950–2000 from a great number of weather stations (>15,000) around the globe. Data are available at different resolutions (i.e., 10.0, 5.0, 2.5, and 0.5 min of arc) as grid (raster) layers. In particular, we used climate surfaces with pixels of 2.5 arc-min, corresponding to a resolution of approximately 5 km (Hijmans et al., 2005b).

In order to describe the climatic niche of *W. mirabilis* populations in a way that is easy-to-read, we represented its geographic distribution in a two-dimensional environmental space. For doing this, we extracted values of annual mean temperature and mean annual precipitation for all the pixels contained in our study area and projected the extracted values in a scatterplot. Values corresponding to pixels contained in the extent of occurrence of *W. mirabilis* were evidenced by coloring the relative points according to their sub-range. This approach produced a visual representation of the climatic niche of the species in its individual sub-ranges in relation with the climatic availability of the study area. In addition, we utilized the same approach for visualizing the climatic shift expected in the three different sub-ranges according to the available scenarios.

We adopted an approach based on ecological niche models to estimate the climatic suitability for *W. mirabilis* populations across the study area (Guisan and Zimmermann, 2000). Since *W. mirabilis* has different climatic niches in its different sub-ranges (see Results for details), we modeled the species distribution in its different sub-ranges separately. We generated 1000 random points across the study area and considered them as pseudo-presence points when they fell into

the considered sub-range and as pseudo-absences otherwise. Climate variables were assigned to the presence/absence points and selected for reducing multicollinearity by discarding those with variance inflation factor higher than 5 (Belsley et al., 1981). After this selection, seven variables were retained for modeling (i.e., bio3 (isothermality), bio4 (temperature seasonality), bio9 (mean temperature of driest quarter), bio14 (precipitation of driest month), bio15 (precipitation seasonality), bio17 (precipitation of driest quarter), and bio18 (precipitation of warmest quarter)). The individual projections for the different sub-ranges were merged by summing up the suitability values in each pixel.

Because the choice of algorithm can influence the performance of ecological niche models, we adopted an ensemble forecasting approach in this study (Araújo and New, 2007). We used eight different techniques and then chose the most common and performing methods, for modeling climatic suitability in the R package (R Development Core Team, 2016) Biomod2 (Thuiller et al., 2016). In particular, we fitted Generalized Linear Model (GLM) (Dobson, 1990), Generalized Boosting Model (GBM) (Friedman et al., 2000), Classification Tree Analysis (CTA) (Breiman et al., 1984), Artificial Neural Network (ANN) (Ripley, 1996), Surface Range Envelop (SRE) (Busby, 1991), Flexible Discriminant Analysis (FDA) (Hastie et al., 1994), Multiple Adaptive Regression Splines (MARS) (Friedman, 1991), and Random Forest Model (RFM) (Breiman, 2001) on presence/absence data for each sub-range. Each model was validated by the ROC (receiver operating characteristic) value (Fielding and Bell, 1997), and a final consensus model was generated by calculating the weighted average of the individual models using the relative ROC values as weights.

To predict the variation of climatic suitability of *W. mirabilis* populations under future conditions, we projected the fitted models into the available Global Climate Models (GCMs). These GCMs were elaborated by 19 research centres for the four RCPs (Representative Concentration Pathway) for 2050 (average for the period 2041–2060) and downscaled at the same spatial resolution as the current climatic surfaces (i.e., 2.5 arc-min) using WorldClim as baseline climate (Hijmans et al., 2005a). Overall, we considered 59 different GCMs and used the same model parameterizations as in the current conditions for projecting the climatic suitability in the future and for creating the final consensus model from the ensemble of the individual models. As a result, we obtained one single spatial prediction of climatic suitability for *W. mirabilis* populations under future conditions.

4 Results

4.1 Climatic exigences

The evident separation of the three sub-ranges has implications for the climatic niches of *W. mirabilis* populations. Overall, this species occurs in the areas with very low annual precipitation respect to the entire study area (Fig. 2). In particular, populations in the southern sub-range (blue in Fig. 2) occupy the driest zones, with a mean annual precipitation <50 mm and an annual mean temperature of 15°C–21°C. Populations in the central sub-range (green in Fig. 2) have a relatively large spectrum of conditions for mean annual precipitation, from very dry (close to 0 mm) to semi-arid (250–300 mm), and a preference for hot conditions (annual mean temperature between 19°C and 22°C) with few exceptions. Populations in the northern sub-range (red in Fig. 2) encounter conditions of mean annual precipitation in the spectrum of the central sub-range (roughly between 100 and 200 mm) but evidently colder (annual mean temperature between 17°C and 20°C).

4.2 Climatic suitability

Species distribution models obtained very high validation scores, i.e., mean ROC values of 0.84 (± 0.20), 0.95 (± 0.03), and 0.85 (± 0.14) for the northern, central, and southern sub-ranges, respectively. The prediction map under current climatic conditions (Fig. 3a) evidenced that the northern and central sub-ranges are divided by a wide unsuitable zone. Differently, the central and southern sub-ranges are connected in terms of climatic suitability. The prediction map under

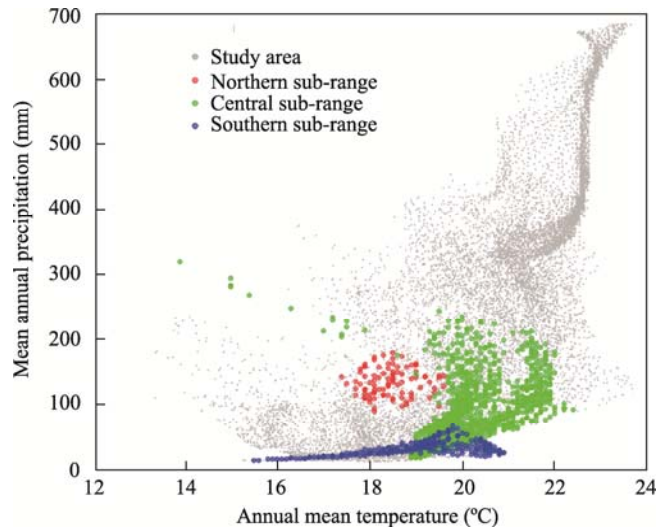


Fig. 2 Visualization of the climatic niche of *W. mirabilis* populations in the three main sub-ranges as well as the whole study area

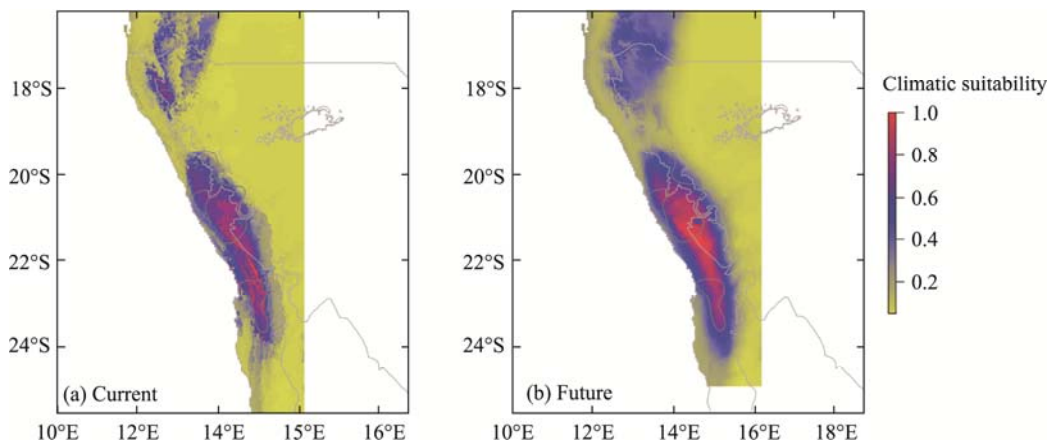


Fig. 3 Climatic suitability for *W. mirabilis* populations under current (a) and future (b) conditions

future conditions (Fig. 3b) shows a similar pattern, with central and southern sub-ranges connected and northern sub-range isolated. In addition, lower levels of climatic suitability are forecasted for the northern sub-range in the future.

The variation of climatic suitability between current and future conditions is expected to occur especially along the borders of species distribution range (Fig. 4). This is particularly evident for the central and southern sub-ranges, where the forecasted suitability variation has positive values (green in Fig. 4) along the borders facing the centre of the continent and negative values (red in Fig. 4) along the borders facing the ocean. The northern sub-range makes an exception to this pattern. Indeed, the entire northern sub-range is expected to suffer a very strong reduction of climatic suitability for *W. mirabilis* populations.

4.3 Climatic variation

The variation of climatic suitability evidenced above corresponds to a substantial increase of temperature expected across the species distribution. According to the 59 available GCMs, the temperature is expected to become higher in the future than in the current in the three sub-ranges (Fig. 5). In particular, the available scenarios of future climate predict that the annual mean temperature will become almost completely different from current conditions in the northern

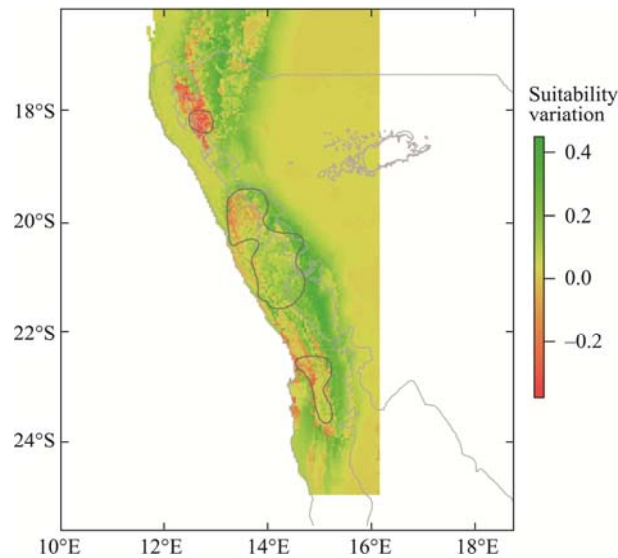


Fig. 4 Variation of climatic suitability for *W. mirabilis* populations between current and future conditions. Note that green shades represent suitability gains and red shades represent suitability losses.

sub-range (Fig. 5a). Differently, in the central and southern sub-ranges (Figs. 5b and c, respectively), the climatic niche of *W. mirabilis* populations will be still satisfied, although the temperature will change. These evidences are in accordance with results from ecological niche models.

5 Discussion

As evidenced by our results, the *W. mirabilis* populations in the northern sub-range have a climatic niche substantially different from those in the central and southern sub-ranges. This is evident both by the visualization in the two-dimensional environmental space and by the outputs of the ecological niche models. Indeed, the northern sub-range is colder than the central sub-range and wetter than the southern sub-range at the same time (Fig. 2). In addition, a wide unsuitable area is present between northern and central sub-ranges (Fig. 3). On the contrary, the central and southern sub-ranges appear relatively similar, being represented by a unique, continuous cloud of points in Figure 2 and being geographically connected in terms of climatic suitability (Fig. 3). This ecological difference between northern and central-southern populations could suggest some level of isolation between these two units, somehow in agreement with the proposal of two distinct taxa (Leuenberger, 2001). Reasons behind Leuenberger's (Leuenberger, 2001) proposal (i.e., different cone morphology) were confuted by a recent study of Jacobson et al. (2014). But, unfortunately, populations from the Namibian northern sub-range were not considered in this study. Similarly, a preliminary study of *Welwitschia* genetic variability also did not include populations from the Namibian northern sub-range or from Angola (Jacobson and Lester, 2003), leaving the issue of the intraspecific taxonomic arrangement open.

W. mirabilis populations in the northern sub-range are more exposed to impacts of climate change than those in the central and southern sub-ranges. Our results show that climatic suitability under future conditions will remain roughly the same for the central and southern sub-ranges. Differently, the northern sub-range will become rather unsuitable in terms of temperature (Fig. 5), losing most of the current climatic suitability in the future (Fig. 3). This evidences the importance of including *W. mirabilis* populations in the northern sub-range into future studies, monitoring plans, and assessing campaigns. Unfortunately, the northern sub-range lays in a remote region, where good roads are rare and traveling time is expanded. As a consequence, this area was seldom included in activities of systematic data collection and

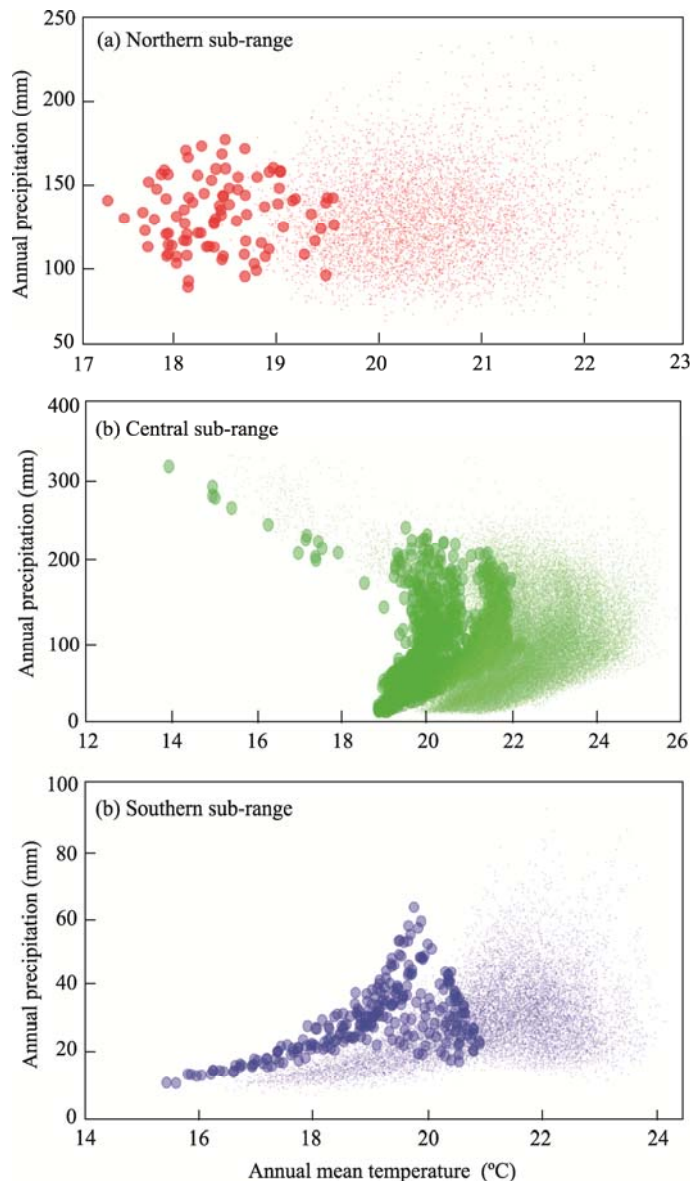


Fig. 5 Variation of climate conditions expected in the three sub-ranges of *W. mirabilis*. In the figure, the big circles are the current values of temperature and precipitation in the northern, central, and southern sub-ranges, while the small circles are the future values in the same areas according to the 59 available Global Climate Models.

available information mainly regards species occurrence or abundance. In the light of our outcomes, demographic studies aimed to understand whether climate change is influencing the structures and dynamics of *W. mirabilis* populations should be carried out, especially in the northern sub-range.

The loss of climatic suitability we evidenced does not mean that of *W. mirabilis* populations will go extinct in the northern sub-range. Individuals of this species are among the oldest living plants, with ages estimated up to 3000 years (Bornman et al., 1972). In addition, it seems that this taxon faced substantial climate changes in the past. Indeed, some traits of *W. mirabilis* are not typically associated with desert plants (e.g., C3 metabolism, high water demands), justifying the hypothesis that earlier habitats of *W. mirabilis* were more mesic than the current habitat in the Namib Desert (Jacobson and Lester, 2003). In addition, paleontologic evidences suggest that *W.*

mirabilis populations existed in the areas corresponding to southern Africa and western South America more than 105 million years ago, before the separation of the continents and the upheaval of the Great Escarpment (Jacobson and Lester, 2003). Since the upheaval of the Great Escarpment was associated to high rainfall in the region, it is plausible that the current fragmented distribution of *W. mirabilis* was due to aridification during the Tertiary and Quaternary (Jacobson and Lester, 2003). In this light, living *W. mirabilis* trees are probably perfectly able to cope with the reduction of climatic suitability. Differently, recruitment could be affected by climate change. Indeed, a certain amount of rain, in specific rainfall pattern, is necessary for seed germination and seedling development (Bornman et al., 1972). For this reason, females annually produce many seeds but recruitment events are rare (many years apart) driving to coetaneous groups of individuals (Henschel and Seely, 2000). Therefore, climate change could prevent the germination of new plants, influencing the population structure.

Obviously, some aspects of our study could be improved with additional data and information. First, distribution data we used are limited to the Namibian range and exclude the northernmost area of presence in Angola. Unfortunately, information about the species distribution in Angola is very general and, to the best of our knowledge, neither precise localities of species presence nor polygons indicating the extent of occurrence of *W. mirabilis* has been published. Moreover, the Angolan portion of *W. mirabilis* range is relatively small, being restricted to the coastal area south of Namibe (Kers, 1967). Second, distribution data we used are not precise localities of presence but general extents of occurrence. Unfortunately, available information about precise sites occupied by *W. mirabilis* is not abundant enough for niche modeling. In addition, density and precision of such information are heterogeneous among sub-ranges. This would have introduced into ecological niche models an unpredictable amount of uncertainty. Furthermore, techniques for cross-scale modeling are available (Bombi and D'Amen, 2012), which allow to produce detailed suitability models from general distribution data (Bombi et al., 2012). Third, ecological niche models assume the focal species is in a status of static equilibrium with environment. This assumption introduces a certain level of uncertainty into the outputs when dealing with non-equilibrium situations (Pearman et al., 2008). Modelers are developing new methods for incorporating adaptation in the process (Bush et al., 2016) to improve the predictive performances in dynamic conditions. Nevertheless, these methods require additional information about the species adaptive potential that is unavailable for *W. mirabilis*. Fourth, we used the ensemble of all the available GCMs for projecting the species niche into the future conditions. Recent studies evidenced that this approach can underestimate the impacts of climate change on wild species (Bombi et al., 2017), including unrealistic models into the possible scenarios. Unfortunately, unavailable information on species responses would be necessary for reducing the uncertainty related to future climate predictions. Overall, further detailed data about species distribution and biology would be needed to overcome these issues, but the state of the art of our knowledge already allowed us to evidence an effect.

6 Conclusions

Welwitschia mirabilis is recognized as a crucial element of the Namib Desert ecosystems. This species has a complex role in the ecosystems where it occurs, representing an element of attraction for many species of vertebrates and arthropods. In addition, this plant species is considered as a symbol of the Namib Desert and contributes to the wellbeing of local human populations drawing tourists in the area. Therefore, the development of a management plan for *W. mirabilis* populations, especially for those in the northern sub-range, would be fundamental and this plan should take into account the fact that climate change could have potential impacts on this species. In this regard, further efforts for collecting additional data on the distribution, demography, and adaptive potential of *W. mirabilis* populations are required.

Acknowledgements

The study was supported by the LifeWatch-ITA European Research Infrastructure on Biodiversity and the Project LIFE+ ManFor C.BD. (LIFE09 ENV/IT/000078). The author wishes to thank Michael THOMPSON for the improvement of the English style, and two anonymous reviewers for their helpful comments.

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