Local adaptation of a dominant coastal tree to freshwater availability and solar radiation suggested by genomic and ecophysiological approaches

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Local adaptation is often a product of environmental variations in geographical space and has implications for biodiversity conservation. We investigated the role of latitudinal heterogeneity in climate on the organization of genetic and phenotypic variation in the dominant coastal tree Avicennia schaueriana. In a common garden experiment, samples from an equatorial region, with pronounced seasonality in precipitation, accumulated less biomass, and showed lower stomatal conductance and transpiration, narrower xylem vessels, smaller leaves and higher reflectance of long wavelengths by the stem epidermis than samples from a subtropical region, with seasonality in temperature and no dry season. Transcriptomic differences identified between trees sampled under field conditions at equatorial and subtropical sites, were enriched in functional categories such as responses to temperature, solar radiation, water deficit, photosynthesis and cell wall biosynthesis. Remarkably, the diversity based on genome-wide SNPs revealed a north-south genetic structure and signatures of selection were identified for loci associated with photosynthesis, anthocyanin accumulation and the responses to osmotic and hypoxia stresses. Our results suggest the existence of divergence in key resource-use characteristics, likely driven by seasonality in water deficit and solar radiation. These findings provide a basis for conservation plans and for predicting coastal plants responses to climate change.

Adaptation is frequently a consequence of spatial variations in selective environmental forces acting on phenotypic diversity1,2. As selective forces operate, they may reduce the heritable variation within a population, leading to the specialization of individuals2. Conversely, in highly stochastic environments, selection can increase the potential of species for phenotypic plasticity3. Therefore, the investigation of adaptation is fundamental for understanding the ability of a species to respond to environmental changes4,5.

Alarmingly high rates of environmental changes have been observed over the last decades, particularly affecting coastal ecosystems6–9, for instance, bleaching of coral reefs worldwide10, large-scale dieback of mangrove communities11 and increased mortality of seagrass beds12. Further environmental changes and coastal biodiversity loss threaten critical socioeconomic and ecological services provided by coastal ecosystems, including their pivotal role as blue carbon stocks13. In this context, studies on the organization of adaptive variation of coastal habitat-forming species are necessary to improve predictions on the impacts of future climate conditions on marine ecosystems14 and to support efficient conservation practices15.

Coastal habitat-forming species are frequently water-dispersed over long distances16,17 and often occur across wide latitudinal ranges. The connectivity among populations of these species may be influenced, not only by their

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dispersion capabilities but also by means of natural selection, caused by varying establishment success over the broad environmental heterogeneity across latitudes18. For instance, distinct coastal species in the North Atlantic show north–south organization of diversity with evidence of selection for different thermal regimes19–21. Similarly, in the Southwest Atlantic, an overlapping north–south structure of genetic diversity, with reduced gene flow between northern and southern populations, has been observed in phylogenetically distant habitat-forming species22–24. In this context, one could expect northerly and southerly populations of these species to adapt differently to contrasting environments over their latitudinal ranges, especially due to their large population sizes2,25. However, the neutrality of the molecular markers used in previous studies has precluded inferences regarding adaptive variation. The existence of local adaptation remains virtually unknown in species that play a central role in sustaining the coastal biodiversity in the South Atlantic. This limited knowledge about the organization of non-neutral variation compromises accurate predictions and suitable conservation efforts for sustainable management.

Here, we tested the hypothesis that latitudinal heterogeneity in climate variables shapes the variation in genotypes and phenotypes involved in the optimization of resource use in widely distributed coastal species. To reduce the potential for incorrect conclusions about selection26, we integrated three independent but complementary approaches, making predictions as follows: (1) using a common garden experiment, individuals from contrasting latitudes would show genetically based phenotypic divergence in ecophysiological traits; (2) under contrasting latitudes, transcriptomic changes would be detected in genes involved in responses to environmental variation; and (3) signatures of selection along the species distribution would be detected in genes involved in responses to latitudinal variation in air temperature, solar radiation and freshwater availability determined by air vapor pressure deficit (VPD), rainfall and tidal regime. We considered the new-world coastal tree species *Avicennia schaueriana* Stapf & Leechman ex Moldenke, found in the Lower Lesser Antilles and widespread from Venezuela to the southernmost temperate mangroves on the Atlantic coast of South America (~28 °S)27, as a model to test this hypothesis. The broad latitudinal range of *A. schaueriana*, spanning wide environmental variation (Fig. 1), and the previously detected north–south structure of neutral variation22, facilitate ecological predictions and the accumulation of divergence, motivating this choice. Our results indicated that fluctuations in VPD, rainfall and solar radiation may be associated with the observed phenotypic and genotypic variation as well as the regulation of gene expression in this dominant coastal tree. We discuss the implications of our results for the persistence of coastal biodiversity in the context of climate change.

**Materials and Methods**

**Propagule sampling.** Mature propagules were collected from 15 *Avicennia schaueriana* Stapf & Leechman ex Moldenke mother trees that were at least 100 m apart from one another in the following two genetically contrasting populations22: (1) the southernmost range limit of American mangroves, in the subtropical region, and (2) an equatorial site in one of the world’s largest macrotidal mangrove forests28,29, near the northernmost limit of the species range (Fig. 1). We refer to samples collected in the former site as “subtropical” and those in the latter as “equatorial” throughout this work. Sampling in more localities was impracticable due to the absence of mature propagules during fieldwork. A detailed characterization of each of these contrasting sites can be found in Table 1, in Supplementary Methods and in the Supplementary Information Fig. S1.

**Comparative ecophysiology using a common garden experiment.** Propagules were germinated as described for *Avicennia germinans*30. We grew propagules in trays with local mangrove soil for two months. After this period, 44 similar-sized seedlings from 30 distinct mother trees, 15 from equatorial and 15 from subtropical...
sites — with an average height of 18 cm, most with three leaf pairs and senesced cotyledons — were transplanted to 6 L pots filled with topsoil and sand (1:1). Seedlings were cultivated for seven months under homogenous conditions in a glasshouse at the University of Campinas, São Paulo, Brazil (22°49′S 47°04′W), where automatic sensors coupled with a data logger (Onset Computer Corp.) measured the atmospheric humidity and temperature every 15 minutes. Seedlings were irrigated daily at 10 a.m. and 5 p.m. with a 3-minute freshwater spray. Twice a week, nutrients were added to the soil using 600 mL of 0.4X Hoagland solution with 15.0 g L$^{-1}$ NaCl per pot. Pots were rotated weekly to reduce the effects of environmental heterogeneity. The environmental conditions in the glasshouse were different from those at each sampling site, as is shown in the Supplementary Information Fig. S2.

The light reflectance of stems was measured in ten plants from each sampling site using a USB4000 spectrophotometer (OceanOptics, Inc.) coupled to a deuterium-halogen light source (DH-2000; OceanOptics) using a light emission range from 200–900 nm. Photosynthesis, stomatal conductance and transpiration rates were measured every 2.0–2.5 hours in five six-month-old individuals from each sampling site on two different days using a Li-Cor 6400 XT (Li-Cor Corp.).

After harvest, three plants without flowers or flower buds from each sampling site were split into leaves, stems and roots, washed with distilled water, dried for 7 days at 70 °C and weighed. The individual leaf area, total leaf area and leaf lamina angle per plant were measured through photographic analyses using ImageJ

| Köppen-Geiger climate characterization | Subtropical | Equatorial |
|--------------------------------------|------------|------------|
| Latitude (°)                        | 285        | 0          |
| Annual Average tidal amplitude (m)$^1$ | 0.45 (Microtidal) | 4.00 (Macrotidal) |
| Annual mean temperature (°C)$^2$     | 20.1       | 26.4       |
| Minimum temperature of the coldest month (°C)$^2$ | 11.8       | 22.0       |
| Maximum temperature of the warmest month (°C)$^2$ | 28.7       | 31.1       |
| Annual precipitation (mm)$^3$        | 1,435      | 2,216      |

Precipitation of the driest month (mm)$^4$ 88 4
Precipitation of the wettest month (mm)$^4$ 162 452
Mean air vapor pressure deficit (kPa)$^7$ $-0.436$ $-0.625$
Mean irradiance (KJ m$^{-2}$ day$^{-1}$)$^3$ 14.270 17.414
Mean sea surface salinity (g/kg)$^6$ 35.50 34.96
Sea surface salinity in the saltiest month (g/kg)$^6$ 36.24 36.87
Sea surface salinity in the freshest month (g/kg)$^6$ 33.73 32.54
Average day length (hours (±SD)) 12.103 (±1.251) 12.115 (±0.033)

Table 1. Characterization of subtropical and equatorial sampling sites of propagules used in the common garden experiment and of plant material used for RNA-sequencing. SD: standard deviation.$^1$ Source: Vestbo et al. (2018)$^{112}$

| True mangrove species in the area | Avicennia schaueriana | Laguncularia racemosa |
|----------------------------------|----------------------|----------------------|

Plant material used for RNA extraction and RNA-sequencing. Plant material used for RNA-sequencing (RNA-Seq) was collected from the equatorial and subtropical sites described in the “Propagule sampling” section and immediately stored in RNAlater (Ambion, Inc.). Because precise species identification requires the analysis...
of both vegetative branches and flower morphology, opportunities for sampling were limited by the reproductive phenology at each visited site. Leaves, stems and flowers from three adult trees at least 100 m apart were collected from July-August of 2014, which corresponds to the end of winter at the subtropical site and the beginning of the dry season at the equatorial site. To minimize the detection of differential transcript expression due to circadian changes, sampling was conducted during low tide and from 10:30 AM to 4:00 PM. A detailed description of the environmental conditions at the time of sampling is available in Supplementary Information Table S1.

We extracted RNA according to Oliveira et al., and evaluated its integrity and purity using agarose gel electrophoresis and a NanoVue spectrophotometer (GE Healthcare Life Sciences). Illumina TruSeq RNA Sample Preparation kits (Illumina, Inc.) were used to construct cDNA libraries. cDNA quality was assessed using the Agilent 2100 Bioanalyzer (Agilent Technologies) and concentrations were quantified by qPCR using the Sequencing Library qPCR Quantification kit (Illumina, Inc.). Sequencing was performed using two 36-cycle TruSeq SBS paired-end kits (Illumina, Inc.) and the Genome Analyzer IIx platform (Illumina, Inc.).

**Assembly and characterization of the *A. schaueriana* transcriptome.** Adapter sequences were trimmed, and 72-bp paired-end reads were filtered by quality (Phred score ≥ 20 for at least 70% of the read length) using the NGS QC Toolkit v.2.3.4. High-quality reads were used for transcriptome assembly in the CLC Genomics Workbench (https://www.qiagenbioinformatics.com/). We used the default settings, except for the distance between pairs (300–500 bp) and k-mer size (45 bp).

Reads were mapped to transcripts using bowtie2 in single-read mode using the default parameters. Transcripts without read-mapping support were removed. Functional annotation was performed using blastx v.2.2.31 with an e-value < 1e−10, NCBI RefSeq v.77. The Arabidopsis Information Resource (TAIR) v.9 and NCBI non-redundant (nr) databases were used as reference databases. We excluded transcripts that were exclusively similar to non-plant sequences. Protein family domains were identified using HMMER3, which iteratively searched transcripts against the Pfam database. To assign Gene Ontology (GO) terms to transcripts, we used the Arabidopsis thaliana gene association file from the GO Consortium and retrieved the information for transcripts showing similar coding sequences in the *A. thaliana* genome. Redundant transcripts were clustered with Cd-hit-est v.4.6.14 using local alignment mode with 95% identity and 70% coverage of the shortest sequence thresholds. Open reading frames (ORFs) were identified using Transdecoder (http://transdecoder.sf.net). We reduced the redundancy of transcripts in the final assembly by retaining for each CD-HIT-EST cluster either the sequence with the longest ORF or, in the absence of sequences containing ORF, the longest sequence. The completeness of the final transcriptome was assessed using BUSCO. Additionally, a reciprocal blastn alignment using an e-value threshold of 10−10 and a minimum alignment length of 100 nucleotides with at least 70% identity was used to compare the *A. schaueriana* transcriptome with other publicly available transcriptomes of congeneric species.

**Comparative transcriptomics using RNA-sequencing.** Tissue-specific count data were obtained from the number of reads uniquely mapped to each transcript of the non-redundant transcriptome using bowtie2 and normalized using edgeR. Differentially expressed transcripts (DETs) between equatorial and subtropical tree tissue-specific samples were detected using the exact test for negative binomial distributions with an FDR < 0.05 (Benjamini-Hochberg). GO term enrichment analyses of the DETs were performed using GOseq with the Wallenius approximation method and P-value < 0.05. Differential expression results were verified using reverse transcription real-time PCR (qRT-PCR) (Supplementary Methods).

**Detection of candidate adaptive loci.** We sampled leaves from 79 adult plants at ten locations spanning most of the distribution of *A. schaueriana* Stapf & Leechman ex Moldenke (Fig. 1, Supplementary Information Table S2). We isolated DNA using the DNeasy Plant Mini Kit (QIAGEN) and NucleoSpin Plant II (Macherey Nagel) following the manufacturers’ instructions. DNA quality and quantity were assessed using 1% agarose gel electrophoresis and the QuantiFluor dsDNA System with a Quantus fluorometer (Promega). Nextera-tagmented DNA was prepared using the Agilente Bioanalyzer (Agilent Technologies) and concentrations were quantified by qPCR using the Illumina TruSeq SBS paired-end kits (Illumina, Inc.) and the Genome Analyzer IIx platform (Illumina, Inc.).

After excluding plants morphologically identified as *A. schaueriana* with genomic signs of hybridization with *A. germinans* (L.) L., we assessed the genetic structure considering all SNPs, using the discriminant analysis of principal components (DAPC) and ADMIXTURE v.1.3.0. For DAPC analyses, we considered the number of groups (K) varying from 1–50 and the Bayesian information criteria for inferring K. Additionally, we used the optim.a.score function to avoid overfitting during the discrimination steps. For the ADMIXTURE analyses, we performed three separate runs for K varying from 1–15 using the block-relaxation method for point estimation; computing was terminated when estimates increased by <0.0001, and the most likely K was determined by cross-validation.
We used two programs to minimize false-positive signs of natural selection: LOSITAN\(^{49}\), assuming an infinite allele model of mutation, using a confidence interval of 0.99, a false-discovery rate (FDR) of 0.1, the neutral mean FST and forcing of the mean FST options; and pcadapt 3.0.4\(^{50}\), which simultaneously identifies the population structure and the loci excessively related to this structure, using an FDR < 0.1.

Putative evidence of selection was considered only for SNP loci that were conservatively identified by pcadapt and five independent runs of LOSITAN to avoid false-positives\(^{51}\). As selection is presumably stronger in coding regions of the genome and there is no reference genome for the species, we used the \textit{de novo} assembled transcriptome, characterized herein, as a reference to identify candidate loci within putative coding regions. We performed a reciprocal alignment between nextRAD sequences (75 bp) and longer expressed sequences (≈300–11600 bp) using blastn v.2.2.31\(^{52}\), with a threshold of at least 50 aligned nucleotides, a maximum of one mismatch and no gaps.

Results

Comparative physiology using a common garden experiment. Seedlings from equatorial and subtropical sites diverged in key ecophysiological traits in the common garden experiment (Figs. 2–4). The inclination angle of the leaf lamina and the average size of individual leaves were smaller in equatorial than in subtropical plants, but total leaf area and specific leaf area did not differ between the groups (Fig. 2, Supplementary Information Table S3). Additionally, equatorial plants showed lower stomatal conductance and transpiration rates than did subtropical plants (Fig. 3). Subtropical plants accumulated more biomass in leaves and roots than did equatorial plants. However, the stem dry mass ratio (MR) (stem dry biomass/plant dry biomass) was greater in equatorial plants, whereas the leaf MR (leaf dry biomass/plant dry biomass) was greater in subtropical plants (Fig. 2). The dry biomass area in the stems and the root MR (root dry biomass/plant dry biomass) were indistinguishable between the groups (Supplementary Information Table S3). Unexpectedly, 63% of the equatorial plants flowered after six months of growth (Supplementary Information Fig. S3g). Since this was not observed in any subtropical plants, flowering plants were not used in the biomass allocation analyses.

Mean stem vessel diameter was smaller in the equatorial seedlings than in the subtropical seedlings, indicating enhanced hydraulic safety (Fig. 2, Supplementary Information Fig. S3h,i). However, the vessel density, vessel grouping index, vessel lumen area in sapwood and total hydraulic conductivity of the stems were not significantly different between the groups (P-value > 0.05) (Supplementary Information Table S3). Plants from contrasting latitudes exhibited different stem epidermis pigmentation, with equatorial seedlings reflecting more long-wavelength of red light (635–700 nm) and less medium wavelength of green light (520–560 nm) than subtropical seedlings (Fig. 4).

Characterization of the \textit{A. schaueriana} transcriptome. In the absence of a reference genome, we obtained the first reference transcriptome for \textit{A. schaueriana} from leaves, stems and flowers of adult individuals under field conditions (Supplementary Information Fig. S4, Table S1). Over 209 million paired-end 72-bp reads showing high quality were \textit{de novo} assembled into a reference, non-redundant transcriptome containing 49,490 sequences, of which 30,227 (61%) were putative protein-coding sequences. Over 91.9% of these reads were mapped to a single transcript, indicating minimum redundancy and a wide representation of sequenced data (Supplementary Information Table S4). Moreover, 91.8% of universal plant orthologous genes were present in the reference transcriptome (Supplementary Information Table S4). Sequences with complete ORFs represented approximately 42% (12,796) of all putative protein-coding transcripts (Supplementary Information Table S5, Fig. S5). Most of these putative protein-coding sequences (94.33%) showed significant similarity (e-value < 1e-10) to proteins in the Plant RefSeq and TAIR databases (Supplementary Information Fig. S5c). More than 80% of protein-coding sequences matched proteins from \textit{Sesamum indicum} or \textit{Erythranthe guttata}, which, as \textit{A. schaueriana}, belong to the order Lamiales (Supplementary Information Fig. S6d). We identified 27,658, 18,325 and 13,273 putative orthologs between the \textit{A. schaueriana} reference transcriptome and transcriptomes derived from \textit{A. marina} leaves\(^{52}\) and \textit{A. officinalis} leaves\(^{53}\) and roots\(^{54}\), respectively (Supplementary Information Table S6).

Comparative transcriptomics between trees from contrasting latitudes. To identify the effects of environmental variation on gene expression and, thus, on phenotypes at contrasting source sites in the field, we compared expression levels in trees under field conditions as comparable as possible, considering tidal conditions and seasonal climates (Supplementary Information Table S1). As expected, we observed a consistent source-site pattern in overall transcript expression levels from leaves and stems (Supplementary Information Fig. S6a,b). However, floral transcript expression levels did not show a clear pattern among samples of the same origin (Supplementary Information Fig. S6c), leading to the identification of only one DET. Thus, we did not include flowers in the subsequent analyses (Supplementary Information Fig. S6f). Conversely, 1,837 and 904 transcripts showed significantly different (FDR < 0.05) relative abundances between equatorial and subtropical samples in leaves and stems, respectively (Supplementary Information Fig. S6d,e). Among the total 2,741 DETs, 1,150 (41.91%) were putative protein-coding transcripts.

The assignment of transcripts to GO terms was possible for 25,184 (83.31%) of the 30,227 putative protein-coding sequences, allowing GO enrichment analyses. Analyses were conducted separately for leaves and stems and for each of the two sets of DETs: one showing higher expression levels in equatorial than in subtropical samples (which we refer to as “DET-Eq”) and the other showing higher abundance in subtropical than in equatorial samples (which are refer to as “DET-St”). We focused on the biological processes associated with key aspects of the responses to contrasting climate conditions between the equatorial and subtropical sites (Table 1, Supplementary Information Fig. S1). The enriched processes among the DET sets included photosynthesis;
Figure 2. Pairwise comparison of morphological traits between *Avicennia schaueriana* seedlings from equatorial and subtropical sites grown using a common garden experiment. Violin plots represent the distribution of observations for plants from equatorial (red) and subtropical (blue) sampling sites. Box plots represent the mean, standard error, and maximum and minimum values. (a) Leaf inclination angle (n = 15 leaves per group, 5 plants per group); (b) individual leaf area (n = 250 leaves per group, 3 plants per group); (c) vessel lumen diameter (n = 180 vessels per group, 3 plants per group); (d–f) leaf, stem and root dry weight (n = 15 plants per group); and (g–i) leaf, stem and root dry mass ratio (organ-specific dry biomass/plant dry biomass) (n = 15 plants per group). For all variables represented in the figure, the absence of a difference between groups was rejected using unpaired Student's T-tests or the Mann-Whitney-Wilcoxon U-tests (b,c), at a significance threshold of 0.05.
cell wall biosynthesis; and plant responses to ultraviolet radiation (UV), temperature stimulus and water stress (Supplementary Information Tables S7–S11, Supplementary Information Fig. S6i–l).

**Photosynthesis.** Among the DET-St set, we observed various putative genes participating in the biosynthesis of the photosynthetic apparatus, chlorophyll and photoreceptors; the function of electron transporters and chloroplast movement coordination. Contrastingly, the DET-Eq set showed enrichment of transcripts similar to Muslim.
proteins required for disassembling the light-harvesting complex of photosystem II in thylakoid membranes\(^5\) (Supplementary Information Table S11).

**Cell wall biosynthesis.** Transcripts similar to those encoding 33 distinct proteins and transcription factors that play central roles in the biosynthesis and deposition of cell wall components, such as cellulose, hemicellulose, lignin and pectin, were identified in DET-Eq (Supplementary Information Table S10).

**Response to UV.** Both the DET-St and DET-Eq sets showed enrichment of functions related to responses to UV radiation; however, the transcript annotations differed between these sets. The DET-St set included putative UV-B protectants and genes involved in UV-B-induced antioxidant biosynthesis, such as plastid copper/zinc superoxide dismutases, photosystem II repair proteins, and L-ascorbic acid. In contrast, the DET-Eq set showed enrichment of transcripts associated with photo-oxidative damage reduction and the positive regulation of anthocyanin biosynthesis in response to UV. Antioxidants induced by UV irradiation\(^8\), such as putative iron superoxide dismutases and pyridoxine biosynthesis genes, were among the DET-Eq set (Supplementary Information Table S11).

**Response to temperature.** In the DET-St set, we observed putative genes presenting critical roles in chloroplast protein translation during cold acclimation and that provide tolerance to chilling temperatures\(^5\)\(^5\)\(^8\). There included transcripts similar to the **GLYCINE-RICH RNA-BINDING (RZ1A)**, which has chaperone activity during cold acclimation\(^6\)\(^8\), and to the cold-inducible ATP-dependent **DNA HELICASE ATSGS1**, required for DNA damage-repair\(^8\). Interestingly, DET-St included a putative **AGAMOUS-LIKE 24 (AGL24)** transcription factor that is involved in the vernalization-induced floral transition\(^6\). Contrastingly, various transcripts similar to heat-shock-inducible chaperones and to **ADENINE NUCLEOTIDE ALPHA HYDROLASE-LIKE (AT3G53990)**, involved in chaperone-mediated protein folding\(^8\), were among the DET-Eq set, potentially enhancing tolerance to heat in equatorial plants. Additionally, a transcript similar to the **ETHYLENE-RESPONSIVE ELEMENT BINDING (RAP2.3)**, which confers resistance to heat and hydrogen peroxide\(^8\), was observed in this group (Supplementary Information Table S11).

**Response to water stress.** Transcripts associated with the response and tolerance to water deficits and with cellular ion homeostasis and osmotic adjustment were enriched among DET-Eq; for instance, a transcript similar to the **ETHYLENE-RESPONSIVE TRANSCRIPTION FACTOR (RAP2.4)**, which regulates the expression of several drought-responsive genes, including aquaporins\(^6\)\(^8\). Accordingly, the putative aquaporin **PLASMA MEMBRANE INTRINSIC (PIP1;4)**\(^6\) was found in this set. In DET-Eq, we observed putative genes participating in the synthesis of raffinose, an osmoprotective soluble trisaccharide\(^7\), as well as transcripts similar to osmosensitive ion channels belonging to the **EARLY-RESPONSIVE TO DEHYDRATION STRESS FAMILY**. Correspondingly, we observed an ion channel, **SLAC1 HOMOLOGUE 3 (SLAH3)**, required for stomatal closure induced by drought stress\(^6\), and the putative **NINE-CIS-EPOXYCAROTENOID DIOXYGENASE 3 (NCED3)**, which increases plant resistance to water deficits through the accumulation of abscisic acid (ABA), leading to stomatal closure. Possibly as a consequence of decreased stomatal conductance, a transcript similar to the photosensitive gene, **ALANINE-GLYCOSYLAMINO TRANSFERASE 2 HOMOLOG 3 (AT3G08860)**, was also observed in DET-Eq (Supplementary Information Table S11).

We confirmed the results obtained from RNA-Seq data computational analyses by qRT-PCR using ten DETs from leaf samples (Supplementary Information Fig. S7, Table S12, Supplementary Note).

**Detection of SNPs with signs of selection.** RNA-Seq enabled the assembly of a reference transcriptome and the identification of biological processes influenced by the environmental divergence over the South American coastline. To complement these analyses, we evaluated the effects of plasticity and adaptive selection in the differential expression\(^29\)\(^71\), we searched for gene sequence variations among trees sampled along the Atlantic coast of South America (Fig. 1, Supplementary Information Table S2). After quality filtering of the sequenced data, we selected 77 individuals without evidence of interspecific hybridization with *A. germinans* for downstream analyses. We identified a set of 6,170 high-quality unlinked biallelic SNP loci with a minor allele frequency \(\geq 0.05\) and \(\geq 8x\) coverage. The overall genetic structure of genome-wide SNPs corroborated the results of a previous study based on putatively neutral loci\(^2\), dividing the species into two populations: north and south of the northeast extremity of South America (Fig. 5).

We observed 122 loci showing considerable departures from neutral expectations of interpopulational divergence, as conservatively detected\(^2\) by pcadapt and LOSITAN. Fifteen of these loci aligned to *A. schaueriana* transcripts that were similar to gene models in *A. thaliana* and *S. indicum* (Supplementary Information Table S13), enabling screening for their potential functional roles. However, five reference proteins did not have informative functions described for the model plant, hindering functional inferences. Conversely, among the remaining annotated candidates, we found five putative genes involved in processes related to contrasting equatorial and subtropical environmental variables (Fig. 6). One candidate locus was detected in the putative transcription factor **RAP2.4**, which is induced in response to water and salt stress\(^8\) and regulates the expression of aquaporins\(^8\). Two other candidates showed similarity to the transcription factors **ZINC-FINGER PROTEIN 1 (ZFN1)**, involved in the regulation of the expression of several water stress genes\(^8\) and the **HYPOXIA RESPONSE ATTENUATOR (HRA1)**, which is strongly induced in response to low oxygen levels\(^7\). A putative **UDP GLUCOSYL TRANSFERASE**, an enzyme that catalyzes the final step of anthocyanin biosynthesis wherein pigmentation is produced\(^7\), also showed evidence of positive selection. Additionally, one candidate locus was in a transcript similar to that of the **TETRATRICOPEPTIDE REPEAT** gene (AT2G20710, TPR), which might play a role in the biogenesis of the photosynthetic apparatus\(^7\).
Discussion

In this study, we integrated genomic and ecophysiological approaches to investigate the foundations of adaptive variations in a dominant tree from the Atlantic coast of South America. We tested the hypothesis that latitudinal variations in climate regimens shape genetic and phenotypic variation involved in resource-use strategies in widespread coastal trees, using the mangrove *Avicennia schaueriana* as a model. Overall, our results supported this hypothesis. Using a common garden experiment, we detected differences between plants from contrasting latitudes in key ecophysiological traits involved in carbon and water balances, indicating the heritability of trait divergence. Accordingly, transcriptomic changes between plants sampled in contrasting latitudes showed enrichment in processes associated with central aspects of the environmental divergence across latitudes, such as responses to temperature, solar radiation, water deficit, photosynthesis and cell wall biosynthesis. The relevance of these biological processes in the field was corroborated by the identification of SNP loci with signs of selection present within putative coding transcripts similar to genes involved in responses to osmotic stress, accumulation of anthocyanins and photosynthesis, unveiling a genetic basis for environmental adaptation. Figure 6 summarizes the integration of three independent approaches that converge to suggest population adaptation to contrasting environments over the species range.

Evidence of a conservative resource-use strategy in the equatorial population of *A. schaueriana*.

Traits exhibited by equatorial plants compared to subtropical plants during the common garden experiment, such as a smaller leaf size and angle, higher levels of red light-reflecting pigments, narrower vessels and lower rates of stomatal conductance, limit carbon acquisition and may have imposed constraints on carbon gain in equatorial plants. Accordingly, these plants also accumulated less biomass than subtropical plants. Despite causing limitations in growth, these characteristics allow plants to maintain suitable leaf temperatures for photosynthesis while reducing the risk of cavitation, UV exposure and water loss through the minimization of evaporative cooling. Given the relevance of these traits to water and carbon balances, especially in high-salinity environments, the prevalence of more conservative resource-use traits among equatorial samples in the common garden experiment suggested selection favoring drought tolerance, likely being advantageous during hot and dry conditions.

Figure 5. Population structure of *A. schaueriana* inferred from genome-wide genotyping of single nucleotide polymorphic (SNP) loci. (a) Density of individuals on the first retained discriminant function, calculated using a multivariate method, the discriminant analysis of principal components (DAPC); different colors represent inferred populations; (b) map showing the distribution of the species (green-shaded area), the geographical location of sampling sites (points) and barplots in which each color denotes an ancestral cluster and each bar represents an individual as inferred by the model-based method implemented in Admixture 1.3.0.
seasons in equatorial wetlands (August-December). These seasons frequently present a combination of stressful conditions, including high UV exposure, highly fluctuating soil salinity resulting from a wide daily tidal range, and a high vapor pressure deficit (VPD) resulting from high temperature (>30 °C) and an air humidity below 60%. Accordingly, equatorial plants also showed lower transpiration rates than subtropical plants in the common garden. Additionally, 63% of the six-month-old equatorial plants flowered from July-August in the common garden, whereas subtropical plants did not flower. This period is consistent with phenological observations reported for A. schaueriana in both equatorial and southern subtropical forests, and could indicate a genetic basis for the observed variation. Early flowering is a phenotype with complex genetic components, rarely studied in non-model organisms but renowned as an adaptive mechanism for maximizing the chances of reproduction under stressful environments.

In their native environment, equatorial plants showed increased expression levels of putative heat-shock proteins, drought-induced ion transporters, aquaporins and genes that play central roles in stomatal closure and photorespiration compared to subtropical plants, likely contributing to heat and drought tolerance. These findings provided multiple lines of evidence of heat and water stress responses. Higher expression of aquaporins and genes involved in the accumulation of organic solutes may contribute to lowering the cellular water potential while improving drought tolerance in equatorial plants compared to subtropical plants. Additionally, higher expression of several transcripts associated with secondary cell wall biosynthesis and thickening may enhance the rigidity of cells and reduce the risk of collapse during dehydration-rehydration cycles in equatorial trees. Moreover, equatorial plants showed lower expression of several putative photosynthesis genes than subtropical plants. In response to drought, high-light and heat stress plants minimize the photooxidative damage by reducing photosynthetic activity via repression of photosynthesis genes. Remarkably, evidence of selection was detected in two putative transcription factors, RAP2.4 and ZFN1, which play key roles in the regulation of osmotic stress-response and to hypoxia (HRA1). Our results suggest that the establishment success of propagules over the latitudinal gradient of the Atlantic coast of South America may play a role in shaping the diversity of genotypes and phenotypes in the widespread mangrove Avicennia schaueriana. Evidence of an acquisitive resource-use strategy in the subtropical population of A. schaueriana. Subtropical plants showed higher stomatal conductance and transpiration rates, higher levels of green light-reflecting pigments, larger leaf area, wider leaf lamina angle and larger xylem vessel diameter than equatorial plants in the common garden experiment. These characteristics enhance light energy absorbance and carbon use efficiency.
recommend that populations of coastal trees occurring north and south from the northeast extremity of South America was detected in a transcription factor highly induced by oxygen deprivation (HRA1)74. Additionally, the HRA1 differentially affect the sensitivity of northerly and southerly populations to a rapidly changing climate. It has been suggested that the diversification of the dominant coastal tree *A. schaueriana*9. The functional divergence described herein might differ from that occurring in equatorial latitudes. Plants optimize the use of light energy and adjust photosynthetic activity through the regulation of light-harvesting and photosystem-component genes83. Thus, the higher expression levels of transcripts associated with photosynthesis in subtropical than in equatorial plants may have facilitated the absorption of light energy in subtropical plants during winter. Although solar irradiance levels were indistinguishable between sampling sites at the time of sampling, transcriptomic changes in putative UV-inducible antioxidant and photodamage repair genes suggest the use of distinct strategies to respond to differential seasonality in photoperiod and solar radiation levels between subtropical and equatorial latitudes. Notably, we observed signatures of selection in two transcripts, one showing similarity to a UDP-GLUCOSYL TRANSFERASE, a key enzyme in the anthocyanin biosynthesis pathway, and the other in a transcript similar to a TPR gene required for chlorophyll biosynthesis and repair. These results imply that solar radiation, in addition to VPD, may act as an environmental factor driving selection in *A. schaueriana*.

Although soil oxygen availability affects plant growth in intertidal areas, we did not focus our experiments on the relevance of hypoxia in shaping adaptive divergence in coastal wetlands84. However, evidence of selection was detected in a transcription factor highly induced by oxygen deprivation (HRA1)74. Additionally, the HRA1 putative homolog also showed 1.75-fold higher expression in subtropical leaves relative to that in equatorial leaves, even though sampling was conducted at low tide at both sites. As tidal amplitude decreases with increasing latitude along the Atlantic coast of South America85 (Fig. 6), trees are exposed to increasing anoxia conditions southwards. These findings suggest that subtropical populations may have better stress preparedness for hypoxia than equatorial populations.

**Climate change and conservation implications.** Our results provide compelling evidence that adaptations to contrasting seasonality in freshwater availability and solar radiation are involved in the organization of diversity of the dominant coastal tree *A. schaueriana*. The functional divergence described herein might differentially affect the sensitivity of northerly and southerly populations to a rapidly changing climate. It has been suggested that the southernmost mangrove forests in the Americas could expand poleward in the near future86. We expect that the observed acquisitive resource-use characteristics may indeed favor subtropical tree growth under increased CO2 concentrations, temperatures and rainfall, as predicted for this region by 210087. However, great landward relocation of subtropical populations will be necessary for their persistence in the face of a rising sea level88 due to the narrowing of intertidal zones towards mid-latitude areas on the Atlantic coast of South America. Even though subtropical populations apparently show better preparedness for hypoxia than equatorial populations, this scenario is aggravated by the dense coastal occupation by urban and agricultural areas, which may preclude the landward migration of subtropical mangroves in South America. Contrastingly, equatorial populations frequently have wider plains that are potentially available for expansion and lower human occupation, but might be threatened by reduced rainfall and increased VPD during more intense El Niño-Southern Oscillation events87. Increased temperature will stimulate both respiration99 and photosynthesis100 and may eventually offset benefits in carbon acquisition caused by increased CO2 concentrations101. The critical temperature threshold for photosynthesis may be overcome more frequently, possibly reducing carbon assimilation and productivity.

With a conservative resource-use strategy, further limitations in net carbon assimilation could, in extreme events, lead to biomass loss or tree mortality triggered by carbon starvation101. This scenario could be especially severe to the semiarid northeast South American coast, particularly in the face of intense human use of adjacent plains and decreasing trends in terrestrial freshwater availability and coastal freshwater input101-103. Our results corroborate previous studies of marine species from the Northern Atlantic that associated limited north-south population connectivity with adaptation to latitudinal environmental dissimilarity10-21. As widespread trees distributed along the Atlantic coast of South America show an overlapping north-south organization of genetic variation20-24, we hypothesize that they may also show divergent adaptations to heterogeneous resource availability over their distribution ranges. Conservation efforts of coastal ecosystems should focus on resilient, habitat-forming species that give shelter and act as climate rescuers for several stress-sensitive species106. We recommend that populations of coastal trees occurring north and south from the northeast extremity of South America should warrant attention as distinct conservation management units107 for the long-term persistence of coastal biodiversity and the ecosystem services they provide.

**Conclusions**

Studies on local adaptations in tropical trees are frequently limited by difficulties in implementing traditional approaches, such as reciprocal transplants, due to their long generation time and the lack of basic biological information, such as knowledge of their evolutionary history, reproductive mode and phenological patterns. Despite limitations, we provided integrated ecophysiological and genomic evidence that supports the hypothesis acquisition, at the expense of a higher cavitation risk88,89. Conversely, this apparent riskier strategy may compensate for seasonal declines in growth resulting from decreasing temperature, photoperiod and light quality at higher latitudes89. Although low temperatures reduce enzymatic activity99 and, thus, plant growth, the severity of low-temperature stress in the southernmost subtropical mangrove forests of the Americas is likely insufficient to favor the selection of freezing-tolerant adaptations, in contrast to results reported for mangroves at their northernmost edge on the Atlantic coast of the Americas92. At the southernmost limit of American mangrove forests, the minimum air temperature does not drop below 0°C and remains within the expected mangrove physiological thresholds93. Additionally, the small population size of *A. schaueriana* at this location27 and the arrival of maladapted propagules from northerly populations likely reduce the potential strength of selection favoring cold-adaptive traits.

Corroborating the observed differences in morphophysiological traits, we found divergence at the molecular level that may also be related to the increasing amplitude of variation in photoperiod and light quality towards high latitudes. Plants optimize the use of light energy and adjust photosynthetic activity through the regulation of high-latitude stress in the southernmost subtropical mangrove forests of the Americas is likely insufficient to favor the selection of freezing-tolerant adaptations, in contrast to results reported for mangroves at their northernmost edge on the Atlantic coast of the Americas92. At the southernmost limit of American mangrove forests, the minimum air temperature does not drop below 0°C and remains within the expected mangrove physiological thresholds93. Additionally, the small population size of *A. schaueriana* at this location27 and the arrival of maladapted propagules from northerly populations likely reduce the potential strength of selection favoring cold-adaptive traits.
that seasonal environmental heterogeneity in solar radiation and freshwater availability may shape the variation of traits associated with resource-use in *A. schaueriana*, a widespread tree along the Atlantic coastline of South America. The non-neutral trait diversity revealed by the interdisciplinary approach used here provides a perspective into the molecular and phenotypic scales at which environmental selection may shape functional variation of this dominant species, on a continental scale. Integrating high-throughput sequencing and ecophysiological approaches, has shown to be a promising strategy in the study of adaptation in non-model, long-lived species. To provide more realistic predictions of how dominant coastal trees may respond to current global climate changes we encourage the development of further studies accounting for phenotypically and genetically informed distribution modeling. Since freshwater availability has been decreasing in coastal areas worldwide, strongly compromising the productivity of coastal plants, such studies should focus on biological variation involved in the balance between carbon gain and water loss. This knowledge can improve predictions on the future of ecosystems they form and generate key information for forest conservation and management efforts. Without the realization and dissemination of such studies, the success of conservation plans for tropical forests and their potential to mitigate climate change will likely be seriously compromised.

**Data availability**

Expression data and sequences that support the findings have been deposited in GenBank with the primary accession code GSE116060. A Variant Call Format file and its complementary file, both required for all of the genome-wide SNP diversity analyses are provided as Supplementary Files (Supplementary Datasets 1, 2).

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**References**

1. Hereford, J. A Quantitative Survey of Local Adaptation and Fitness Trade-Offs. *Am. Nat.* 173, 579–588 (2009).
2. Kawecki, T. J. & Ebert, D. Conceptual issues in local adaptation. *Ecol. Lett.* 7, 1225–1241 (2004).
3. Albert, C. H., Grassein, F., Schneidereit, C. & H. Vasse, C. Schneidereit, C. & H. Vasse, C. When and how should intraspecific variability be considered in trait-based plant ecology? *Perspect. Plant Ecol. Evol. Syst.* 13, 217–225 (2011).
4. Aitken, S. N., Yeaman, S., Holliday, J. A., Wang, T. & Curtis-McLane, S. Adaptation, migration or extirpation: climate change outcomes for tree populations. *Evol. Appl.* 1, 95–111 (2008).
5. Jump, A. S. & Peñuelas, J. Running to stand still: adaptation and the response of plants to rapid climate change. *Ecol. Lett.* 8, 1010–1020 (2005).
6. Hoegh-Guldberg, O. & Bruno, J. F. The Impact of Climate Change on the World’s Marine Ecosystems. *Science* (80-.). 328, 1523–1528 (2010).
7. Loarie, S. R. *et al.* The velocity of climate change. *Nature* 462, 1052–1055 (2009).
8. Duke, N. C. A World Without Mangroves? *Science* (80-.). 317, 41 (2007).
9. Poldoro, B. A. *et al.* The loss of species: Mangrove extinction risk and geographic areas of global concern. *PLoS One* 5, e10095 (2010).
10. Normile, D. El Niño’s warmth devastating reefs worldwide. *Science* (80-.). 352, 15–16 (2016).
11. Duke, N. *et al.* Large-scale dieback of mangroves in Australia. *Mar. Freshw. Res.* 68, 1816–1829 (2017).
12. Marba, N. & Duarte, C. M. Mediterranean warming triggers seagrass (Posidonia oceanica) shoot mortality. * Glob. Chang. Biol.* 16, 2366–2375 (2010).
13. Kaufman, J. R. *et al.* Carbon stocks of mangroves and salt marshes of the Amazon region. *Brazil. Biol. Lett.* 14, 2018028 (2018).
14. Moran, E. V., Hartig, F. & Bell, D. M. Intraspecific trait variation across scales: Implications for understanding global change responses. *Glob. Chang. Biol.* 22, 137–150 (2016).
15. Wee, A. K. S. *et al.* The integration and application of genomic information in mangrove conservation. *Conserv. Biol.* 33, 206–209 (2018).
16. Takayama, K., Tamura, M., Tateishi, Y., Webb, E. L. & Kajita, T. Strong genetic structure over the American continents and transoceanic dispersal in the mangrove genus *Rhizophora* (Rhizophoraceae) revealed by broad-scale nuclear and chloroplast DNA analysis. *Am. J. Bot.* 100, 1191–1201 (2013).
17. Mori, G. M., Zucchi, M. I. Sampaio, I. & Souza, A. P. Species distribution and introgressive hybridization of two *Avicennia* species from the Western Hemisphere unveiled by phylogenetic patterns. *BMC Evol. Biol.* 15, 1–15 (2015).
18. Marshall, D. J., Monro, K., Bode, M., Kroug, M. J. & Sweare, S. Phenotype-environment mismatches reduce connectivity in the sea. *Ecol. Lett.* 13, 128–140 (2010).
19. Jeffery, N. W. *et al.* Range-wide parallel climate-associated genomic clines in Atlantic salmon. *R. Soc. Open Sci.* 4, 171394 (2017).
20. Bradbury, I. R. *et al.* Parallel adaptive evolution of Atlantic cod on both sides of the Atlantic Ocean in response to temperature. *Proc. R. Soc. B Biol. Sci.* 277, 3725–3734 (2010).
21. Chai, N. D., Kaliuziecki, S. T., Trussell, G. C. & Vollmer, S. V. Phylogenomic analyses reveal latitudinal population structure and polymorphisms in heat stress genes in the North Atlantic snail *Nucella lapillus*. *Mol. Ecol.* 23, 1863–1873 (2014).
22. Mori, G. M., Zucchi, M. I. & Souza, A. P. Multiple-geographic-scale genetic structure of two mangrove tree species: the roles of mating system, hybridization, limited dispersal and extrinsic factors. *PLoS One* 10, e0118710 (2015).
23. Francisco, P. M., Mori, G. M., Alves, F. M., Tambarussi, E. & Souza, A. P. De. Population genetic structure, introgression, and five hybridization in the genus *Rhizophora* along the Brazilian coast. *Ecol. Evol.* 8, 3491–3504 (2018).
24. Takayama, K., Tateishi, Y., Murata, J. & Kajita, T. Gene flow and population subdivision in a pantropical plant with sea-drifted seeds *Hibiscus tiliaceus* shoot mortality. *Evol. Appl.* 5, 137–150 (2016).
25. Savolainen, O., Pyhäjärvi, T. & Knürr, T. Gene Flow and Local Adaptation in Trees.
26. Barrett, R. D. H. & Hoekstra, H. E. Molecular sandspurs: tests of adaptation at the genetic level. *Nat. Rev. Genet.* 12, 767–780 (2011).
27. Soares, M. I. G., Estrada, G. C. D. & Tognella, M. P. M. Southern limit of the Western South Atlantic mangroves: Assessment of the potential effects of global warming from a biogeographical perspective. *Estuar. Coast. Shelf Sci.* 101, 44–53 (2012).
28. Souza-Filho, P. W. M. *et al.* Holocene coastal evolution and facies model of the Bragança macrotidal flat on the Amazon Mangrove Coast, northern Brazil. *J. Coast. Res.* 1, 306–310 (2006).
29. Kjerfve, B. *et al.* Morphodynamics of muddy environments along the Atlantic coasts of North and South America. In Muddy coasts of the World: Processes, Deposits and Function (eds Healy, T., Wang, Y. & Healy, J.-A.) 4, 479–532 (Elsevier Science B. V., 2002).
30. Reef, R. *et al.* The effect of atmospheric carbon dioxide concentrations on the performance of the mangrove *Avicennia germinans* over a range of salinities. *Physiol. Plant.* 154, 358–368 (2015).
31. Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH Image to ImageJ: 25 years of image analysis. *Nat. Methods* 9, 671–675 (2012).
32. Scholz, A., Klepsch, M., Karimi, Z. & Jansen, S. How to quantify conduits in wood? Front. Plant Sci. 4, 1–11 (2013).
33. Oliveira, R. R., Viana, A. J. C., Rejtegui, A. C. E. & Vincentz, M. G. A. An efficient method for simultaneous extraction of high-quality RNA and DNA from various plant tissues. Genet. Mol. Res. 14, 18828–18838 (2015).
34. Patel, R. K. & Jain, M. NGS QC toolkit: A toolkit for quality control of next generation sequencing data. PLoS One 7, e30619 (2012).
35. Langmead, B., Trapnell, C., Pop, M. & Salzberg, S. Ultrafast and memory-efficient alignment of short DNA sequences to the human genome. Genome Biol. 10, R25 (2009).
36. Camacho, C. et al. BLAST ++: architecture and applications. BMC Bioinformatics 10, 421 (2009).
37. O’Leary, N. A. et al. Reference sequence (RefSeq) database at NCBI: Current status, taxonomic expansion, and functional annotation. Nucleic Acids Res. 44, D733–D745 (2016).
38. Berardini, T. Z. et al. The Arabidopsis Information Resource: Making and Mining the “Gold Standard” Annotated Reference Plant Genome. Genesis 53, 474–485 (2015).
39. Finn, R. D. et al. Pfam: The protein families database. Nucleic Acids Res. 42, 222–230 (2014).
40. Blake, J. A. et al. Gene ontology annotation: Going forward. Nucleic Acids Res. 43, D1049–D1056 (2015).
41. Li, W. & Godzik, A. Cx-hit: A fast program for clustering and comparing large sets of protein or nucleotide sequences. Bioinformatics 22, 1658–1659 (2006).
42. Simão, F. A., Waterhouse, R. M., Ioannidis, P., Kriventseva, E. V. & Zdobnov, E. M. BUSCO: Assessing genome assembly and annotation completeness with single-copy orthologs. Bioinformatics 31, 3210–3212 (2015).
43. Robinson, M. D., McCarthy, D. I. & Smyth, G. K. edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26, 139–140 (2010).
44. Young, M. D., Wakefield, M. J., Smyth, G. K. & Oshlack, A. Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biol. 11, R14 (2010).
45. Robinson, M. D., McCarthy, D. J. & Smyth, G. K. edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26, 139–140 (2010).
46. Young, M. D., Wakefield, M. J., Smyth, G. K. & Oshlack, A. Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biol. 11, R14 (2010).
47. Rusello, M. A., Waterhouse, M. D., Etterson, D. P. F. & Johnson, E. A. From promise to practice: pairing non-invasive sampling with genomics in conservation. Peer. 3, e1106 (2015).
48. Danecek, P. et al. The variant call format and VCFtools. Bioinformatics 27, 2156–2158 (2011).
49. Jombart, T., Devillard, S. & Balloux, F. Discriminant analysis of principal components: a new method for the analysis of genetically structured populations. BMC Genet. 11, 94 (2010).
50. Alexander, D. H., Novembre, J. & Lange, K. Fast model-based estimation of ancestry in unrelated individuals. Genem 19, 1655–1664 (2009).
51. Antao, T., Lopes, A., Lopes, R. J., Beja-Pereira, A. & Luikart, G. LOSITAN: A workbench to detect molecular adaptation based on a Fst-outlier method. BMC Bioinformatics 9, 1–5 (2008).
52. Lui, K., Bazin, E. & Blum, M. G. B. pcadapt: An R package to perform genome scans for selection based on principal component analysis. Mol. Ecol. Resour. 33, 67–77 (2016).
53. Lotterhos, K. E. & Whitlock, M. C. The relative power of genome scans to detect local adaptation depends on sampling design and statistical method. Mol. Ecol. 24, 1031–1046 (2015).
54. Huang, J. et al. Transcriptome Sequencing and Analysis of Leaf Tissue of Avicennia marina Using the Illumina Platform. PLoS One 9, e108785 (2014).
55. Lys, H., Li, X., Guo, Z., He, Z. & Shi, S. De novo assembly and annotation of the Avicennia officinalis L. transcriptome. Mar. Genomics 38, 17–20 (2017).
56. Krishnamurthy, P. et al. Transcription analysis of salt stress tolerance in the roots of the mangrove Avicennia officinalis. Nat. Sci. Reports 7, 1–19 (2017).
57. Park, S.-Y. et al. The Senescence-Induced Staygreen Protein Regulates Chlorophyll Degradation. Plant Cell 19, 1649–1664 (2007).
58. Myouga, F. et al. A Heterocomplex of Iron Superoxide Dismutases Defends Chloroplast Nucleoids against Oxidative Stress and Is Essential for Chloroplast Development in Arabidopsis. Plant Cell Online 20, 3148–3162 (2008).
59. Wang, S. et al. Chloroplast RNA-Binding Protein RBD1 Promotes Chilling Tolerance through 23S rRNA Processing in Arabidopsis. PLoS Genet. 12, 1–21 (2016).
60. Goulas, E. et al. The chloroplast lumen and stromal proteomes of Arabidopsis thaliana show differential sensitivity to short- and long-term exposure to low temperature. Plant J. 47, 720–734 (2006).
61. Kim, Y. O., Kim, J. S. & Kang, H. Cold-inducible zinc finger-containing glycin-rich RNA-binding protein contributes to the enhancement of freezing tolerance in Arabidopsis thaliana. Plant J. 42, 890–900 (2005).
62. Hartung, F., Suer, S. & Puchta, H. Two closely related RecQ helicases have antagonistic roles in homologous recombination and DNA repair in Arabidopsis thaliana. Proc. Natl. Acad. Sci. 104, 18836–18841 (2007).
63. Michaels, S. D. et al. AGL24 acts as a promoter of flowering in Arabidopsis and is positively regulated by vernalization. Plant J. 33, 867–874 (2003).
64. Jung, Y. J. et al. Universal Stress Protein Exhibits a Redox-Dependent Chaperone Function in Arabidopsis and Enhances Plant Tolerance to Heat Shock and Oxidative Stress. Front. Plant Sci. 6, 1–11 (2015).
65. Ogawa, T. et al. Functional Analysis of Arabidopsis Ethylene-Responsive Element Binding Protein Confering Resistance to Bax. Plant Physiol. 138, 1436–1445 (2005).
66. Lin, R. C., Park, H. J. & Wang, H. Y. Role of Arabidopsis RAP2.4 in regulating light- and ethylene-mediated developmental processes and drought stress tolerance. Mol. Plant 1, 42–57 (2008).
67. Rae, L. E., Lao, N. T. & Kavanagh, T. A. Regulation of multiple aquaporin genes in Arabidopsis by a pair of recently duplicated DREB transcription factors. Plant Cell 234, 429–444 (2011).
68. Alexandersson, E. et al. Whole gene family expression and drought stress regulation of aquaporins. Plant Mol. Biol. 59, 469–484 (2005).
69. T. Z. B. A. and Shigeoka, S. Galactinol and Raffinose Constitute a Novel Function to Protect Plants from Oxidative Stress to Heat Shock and Oxidative Stress. Plant Cell Online 20, 3148–3162 (2008).
70. Wolf, J. B. W., Lindell, J. & Backstrom, N. Speciation genetics: current status and evolving approaches. Philos. Trans. R. Soc. B Biol. Sci. 365, 1717–1733 (2010).
71. Liu, W. C., Chen, Y. & Lowry, D. B. Gene Regulatory Divergence Between Locally Adapted Ecotypes in Their Native Habitats. Mol. Ecol. 27, 4174–4188 (2018).
72. A. C. H. & Iwabuchi, M. Expression of a subset of the RAP2.4 in regulating light- and ethylene-mediated developmental processes and drought stress tolerance. Mol. Plant 1, 42–57 (2008).
73. Ahrens, C. W. et al. The search for loci under selection: trends, biases and progress. Mol. Ecol. 27, 1342–1356 (2018).
74. Sakamoto, H., Araki, T., Meshi, T. & Ivabuchi, M. Expression of a subset of the Arabidopsis Cys2/His2-type zinc-finger protein gene family under water stress. Gene 248, 23–32 (2000).
75. Guiontoli, B. et al. A trihelix DNA binding protein counterbalances hypoxia-responsive transcriptional activation in Arabidopsis. PLoS Biol. 12 (2014).
76. Töhö, T. et al. Functional genomics by integrated analysis of metabolome and transcriptome of Arabidopsis plants over-expressing an MYB transcription factor. Plant J. 42, 218–235 (2005).
76. Bohne, A. V., Schwenkert, S., Grimm, B. & Nickelsen, J. Roles of Tetratricopeptide Repeat Proteins in Biogenesis of the Photosynthetic Apparatus. *Int. Rev. Cell Mol. Biol.* **324**, 187–227 (2016).
77. Reef, R. & Lovelock, C. E. Regulation of water balance in Mangroves. *Ann. Bot.* **115**, 385–395 (2015).
78. Menezes, M. P. M., de, Berger, U. & Mehlig, U. Mangrove vegetation in Amazonia: a review of studies from the coast of Pará and Maranhão States, north Brazil. *Acta Amaz.* **38**, 403–420 (2008).
79. Alkire, M. A. M., S. B., Botasso, P. C. & Soffiati, P. Stem growth and phenology of three subtropical mangrove tree species. *Brazilian J. Bot.* **40**, 907–914 (2017).
80. Kazan, K. & Lyons, R. The link between flowering time and stress tolerance. *J. Exp. Bot.* **67**, 47–60 (2016).
81. Lovelock, C. E. et al. The vulnerability of Indo-Pacific mangrove forests to sea-level rise. *Nature* **526**, 559–563 (2015).
82. Gall, H. L. et al. Cell Wall Metabolism in Response to Abiotic Stress. *Plants* **4**, 112–166 (2015).
83. Kimura, M. et al. Identification of Arabidopsis Genes Regulated by High Light – Stress Using cDNA Microarray. *Photochem. Photobiol.* **77**, 226–233 (2003).
84. Wang, D. et al. Genome-wide temporal-spatial gene expression profiling of drought responsiveness in rice. *BMC Genomics* **12**, 149 (2011).
85. Moumeni, A. et al. Comparative analysis of root transcriptome profiles of two pairs of drought-tolerant and susceptible rice near-isogenic lines under different drought stress. *BMC Plant Biol.* **11**, 174 (2011).
86. Wei, W. et al. Melatonin enhances plant growth and abiotic stress tolerance in soybean plants. *J. Exp. Bot.* **66**, 695–707 (2015).
87. McRae, G. J. A Simple Procedure for Calculating Atmospheric Water Vapor Concentration. *J. Air Pollut. Control Assoc.* **30**, 394–394 (1980).
88. Stuart, S. A., Choot, B., Martin, K. C., Holbrook, N. M. & Ball, M. C. The role of freezing in setting the latitudinal limits of mangrove forests. *New Phytol.* **173**, 576–583 (2007).
89. Carlson, J. E., Holsinger, K. E. & Prunier, R. Plant responses to climate in the cape floristic region of South Africa: Evidence for adaptive differentiation in the proteaceae. *Evolution (N. Y.)* **65**, 108–124 (2011).
90. Pecot, S. D., Horsley, S. B., Battaglia, M. A. & Mitchell, R. J. The influence of canopy, sky condition, and solar angle on light quality in a longleaf pine woodland. *Can. J. For. Res.* **35**, 1356–1366 (2005).
91. Arcus, V. L. et al. On the Temperature Dependence of Enzyme–Catalyzed Rates. *Biochemistry* **55**, 1681–1688 (2016).
92. Cavanaugh, K. C. et al. Poleward expansion of mangroves is a threshold response to decreased frequency of extreme cold events. *Proc. Natl. Acad. Sci. USA* **111**, 723–723 (2014).
93. Osland, M. J. et al. Climatic controls on the global distribution, abundance, and species richness of mangrove forests. *Ecol. Monogr.* **87**, 341–359 (2017).
94. Colmer, T. D. & Flowers, T. J. Flooding tolerance in halophytes. *New Phytol.* **179**, 964–974 (2008).
95. Schaeffer-Novelli, Y., Cintrón-Molero, G., Adaime, R. R. & de Camargo, T. M. Variability of mangrove ecosystems along the Brazilian coast. *Estuaries* **13**, 204–218 (1990).
96. Godoy, M. D. P. & De Lacerda, L. D. Mangroves Response to Climate Change: A Review of Recent Findings on Mangrove Extension and Distribution. *Ann. Brazilian Acad. Sci.* **87**, 651–667 (2015).
97. IPCC. *Climate Change 2014: Synthesis Report. Contribution of Working Groups I, II and III to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change*. Cambridge University Press (2014).
98. Elliott, J. C. Vulnerability assessment of mangroves to climate change and sea-level rise impacts. *Wetl. Ecol. Manag.* **23**, 115–137 (2015).
99. Heské, M. A. et al. Convergence in the temperature response of leaf respiration across biomes and plant functional types. *Proc. Natl. Acad. Sci.* **113**, 3832–3837 (2016).
100. Jordan, D. R. & Ogren, W. The CO2/O2 specificity of ribulose 1,5-bisphosphate carboxylase/oxygenase. *Planta* **161**, 308–313 (1984).
101. Drake, B. G., González-Meler, M. A. & Long, S. P. More efficient plants: a consequence of rising atmospheric CO22? *Annu. Rev. Plant Physiol. Plant Mol. Biol.* **48**, 609–639 (1997).
102. Saenger, P. & Moerly, J. Vegetative phenology along the Queensland coastline. *Proc. Ecol. Soc. Aust.* **13**, 257–265 (1985).
103. Doughty, C. E. et al. Drought impact on forest carbon dynamics and fluxes in Amazonia. *Nature* **519**, 78–82 (2015).
104. Rodell, M. et al. Emerging trends in global freshwater availability. *Nature* **557**, 651–659 (2018).
105. Osland, M. J. et al. Climate and plant control on soil organic matter in coastal wetlands. *Glob. Chang. Biol.* https://doi.org/10.1111/gcb.14376 (2018).
106. Bulleri, F. et al. Harnessing positive species interactions as a tool against climate-driven loss of coastal biodiversity. *Plos Biol.* **16** (2018).
107. Mortitz, C. Defining ‘Evolutionarily Significant Units’ for conservation. *Trends Ecol. Evol.* **9**, 373–375 (1994).
108. Ikeda, D. H. et al. Genetically informed ecological niche models improve climate change predictions. *Glob. Chang. Biol.* **23**, 164–176 (2017).
109. Holliday, I. A. et al. Advances in genetic information in forest trees and applications to genetic resources conservation and breeding. *Mol. Ecol.* **25**, 706–717, https://doi.org/10.1111/mec.13963 (2017).
110. Hjornes, R. J., Cameron, S. E., Parra, J. L., Jones, P. G. & Jarvis, A. Very high resolution interpolated climate surfaces for global land areas. *Int. J. Climatol.* **25**, 1965–1978 (2005).
111. Fick, S. E. & Hijmans, R. J. WorldClim 2: new 1-km spatial resolution climate surfaces for global land areas. *Int. J. Climatol.* **43**, 4032–4035 (2017).
112. Vestbo, S., Obst, M., Quevedo Fernandez, F. J., Intanai, I. & Funch, P. Present and Potential Future Distributions of Asian Horseshoe Crabs Determine Areas for Conservation. *Front. Mar. Sci.* **5** (2018).
113. Silva, E. J. & Barber, P. H. MARSPEC: ocean climate layers for marine spatial ecology. *Brazilian J. Bot.* **38**, 1079–1095 (2015).
114. Forsythe, W. C., Rijkie, E. J., Stahl, R. S., Wu, H. I. & Schoolfield, R. M. A model comparison for daylength as a function of latitude and day of year. *Ecol. Modell.* **80**, 87–95 (1995).

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Author contributions
A.P.S., R.S.O., G.M.M. and M.V.C. designed the study. M.V.C. and G.M.M. conducted fieldwork. M.V.C. and C.S.-M. cultivated seedlings and performed analyses of morphophysiological data. M.V.C. prepared samples and performed RNA-sequencing. M.V.C., M.D., D.-H.O. and G.M.M. analyzed RNA-Seq data. C.C.S. and M.V.C. verified RNA-Seq data through qRT-PCR. G.M.M. prepared samples and performed genome-wide SNP genotyping. G.M.M. and M.V.C. analyzed nextRAD results. A.P.S., M.I.Z., G.M.M. and R.S.O. contributed material/reagents/analysis tools. M.V.C. and G.M.M. wrote the manuscript. All authors discussed the results and contributed to the manuscript.

Competing interests
The authors declare no competing interests.

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