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# **The genome of the seagrass** *Zostera marina* **reveals angiosperm adaptation to the sea**

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Seagrasses colonized the sea<sup>1</sup> on at least three independent occasions **to form the basis of one of the most productive and widespread coastal ecosystems on the planet2 . Here we report the genome of**  *Zostera marina* **(L.), the first, to our knowledge, marine angiosperm to be fully sequenced. This reveals unique insights into the genomic losses and gains involved in achieving the structural and physiological adaptations required for its marine lifestyle, arguably the most severe habitat shift ever accomplished by flowering plants. Key angiosperm innovations that were lost include the**  entire repertoire of stomatal genes<sup>3</sup>, genes involved in the synthesis **of terpenoids and ethylene signalling, and genes for ultraviolet protection and phytochromes for far-red sensing. Seagrasses have also regained functions enabling them to adjust to full salinity. Their cell walls contain all of the polysaccharides typical of land plants, but also contain polyanionic, low-methylated pectins and sulfated galactans, a feature shared with the cell walls of all macroalgae4 and that is important for ion homoeostasis, nutrient uptake and O2/CO2 exchange through leaf epidermal cells. The** *Z. marina* **genome resource will markedly advance a wide range of functional ecological studies from adaptation of marine ecosystems under climate warming5,6 , to unravelling the mechanisms of osmoregulation under high salinities that may further inform our understanding of the evolution of salt tolerance in crop plants7 .**

Seagrasses are a polyphyletic assemblage of basal monocots belonging to four families in the Alismatales<sup>1,2</sup> (Supplementary Note 1.1) and Supplementary Fig. 1.1). As a functional group, they provide the foundation of highly productive ecosystems present along the coasts of all continents except Antarctica, where they rival tropical rain forests and coral reefs in ecosystem services<sup>8,9</sup>. In colonizing sedimentary shorelines of the world's ocean, seagrasses found a vast new habitat free of terrestrial competitors and insect pests but had to adapt to cope with new structural and physiological challenges related to full marine conditions.

the Pacific and Atlantic<sup>10</sup>. A clone of *Z. marina* was sequenced from the Archipelago Sea, southwest Finland, using a combination of fosmid-ends and whole-genome shotgun (WGS) approaches (Methods, Supplementary Note 2). The 202.3Mb *Z. marina* genome encodes 20,450 protein-coding genes, 86.6% of which (17,511 genes, Supplementary Note 3.1) are supported by transcriptome data from leaves, roots and flowers (Extended Data Fig. 1, Supplementary Notes 3.2–3.3 and Supplementary Data 1–3). Genes are located in numerous gene-dense islands separated by stretches of repeat elements accounting for 63% of the non-gapped assembly (Extended Data Fig. 2, Supplementary Note 3.1) as compared to only 13% in the only other sequenced alismatid, the freshwater duckweek, *Spirodela polyrhiza* (Alismatales, Araceae)<sup>11</sup>. Gypsy-type (32%) and Copia-type (20%) transposable elements contribute to most of the repetitive DNA. Sequence divergence analysis suggests that the genome retains copies from two distinct periods of invasion by Copia elements, but only one period for Gypsy elements (Extended Data Fig. 3a–c). Genes gained by *Z. marina* ('accessory') are located closer to transposable elements than to conserved ('single copy') genes (Fisher's exact test,  $P < 0.0001$ ) indicating that transposable elements may have played a role in genic adaptation.

We identified 36 conserved microRNAs with high confidence and their predicted targets (Supplementary Note 3.4, Supplementary Data 4 and 5). A novel variant of miR528 (not present in *Spirodela*) was found to be the only member of this miRNA family, and demonstrates that this conserved miRNA is the only one ancestral to the entire monocot lineage. Most likely, *Z. marina* did not take part in the subsequent birth of miRNAs that are common to several other monocots<sup>12</sup>; nor did it experience or retain traces of prominent miRNA duplications.

Analysis of synonymous substitutions per synonymous site  $(K_S)$  age distributions indicates that *Z. marina* carries the remnants of an independent, ancient whole-genome duplication (WGD) event (Fig. 2a, Supplementary Note 4.1)<sup>13</sup>. Duplicated segments account for  $\sim$ 9% of the *Z. marina* genome, probably an underestimate due to the fragmented nature of the assembly. *Zostera* and *Spirodela* diverged somewhere between 135 and 107 million years ago  $(Mya)^{14}$  and

*Zostera marina* (Zosteraceae), or eelgrass (Fig. 1), is the most widespread species throughout the temperate northern hemisphere of

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**Figure 1** | *Zostera marina* **and phylogenetic tree showing gene family expansion/contraction analysis compared with 13 representatives of the Viridiplantae. a**, Gains and losses are indicated along branches and nodes. The number of gene families, orphans (single-copy gene families) and

phylogenomic dating13 of the *Z. marina* WGD suggests that it occurred 72–64 Mya (Fig. 2b), thus independently from the two WGDs reported for *S. polyrhiza*11. This timeframe coincides with the initial diversification of a freshwater clade that includes three of the four families of seagrasses (Supplementary Table 1.1) and with the Cretaceous– Palaeogene (K–Pg) extinction event (Fig. 2c), which provided new ecological opportunities and may have triggered seagrass adaptive radiations.

We mapped signatures of loss and gain of gene families (Supplementary Note 4.2) onto a phylogenetic tree (Fig. 1a). We also mapped losses and gains of Pfam domains (Supplementary Fig. 4.4,



number of predicted genes is indicated next to each species. Background colours (top to bottom) are Alismatales, other monocots, dicots, mosses/ algae **b**, Typical *Zostera marina* meadow, Archipelago Sea, southwest Finland (photo by C.B.).

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Supplementary Data 6). While many genes are shared between *Zostera* and *Spriodela*, clearly some losses and gains are unique to *Zostera* in relation to its marine environment, the alismatid lineage having set the stage for the subsequent freshwater–marine transition. Those unique to *Z. marina* include the absence of all the genes involved in stomatal differentiation (Fig. 3a, Extended Data Table 1 and Supplementary Note 5.1) and the disappearance of genes comprising entire pathways encoding volatiles synthesis and sensing (Supplementary Note 6.1), such as those for ethylene<sup>15</sup> (Fig. 3b, Extended Data Table 2). Terpenoid genes are also drastically reduced to two (Fig. 3c), as compared with four in *Spirodela*, 50 in *Oryza* and>100 in *Eucalyptus*, thus

**Figure 2** | **Ancient whole-genome duplication (WGD). a**, *K*S-based age distribution of the whole *Z. marina* paranome. The *x* axis shows the synonymous distance until a  $K_S$  cut-off of 2, in bins of 0.04, containing the *K*<sub>S</sub> values that were used for mixture modelling (excluding those with a  $K_S \leq 0.1$ ). The component of the Gaussian mixture model plotted in red (as identified by EMMIX) corresponds to a WGD feature based on the SiZer analysis (other components are shown in black). The transition from the blue to the red at a  $K_S$  of  $\sim$ 0.8 in the SiZer panel (below) indicates a change in the distribution and therefore provides evidence for an ancient WGD (Supplementary Table 4.1, Supplementary Fig. 4.1). **b**, Absolute age distribution obtained by phylogenomic dating of *Z. marina* paralogues. The solid black line represents the kernel density estimate (KDE) of the dated paralogues and the vertical dashed black line represents its peak, used as the consensus WGD age estimate, at 67 Mya. Grey lines represent the density estimates from 2,500 bootstrap replicates and the vertical black dotted lines represent the corresponding 90% confidence interval for the WGD age estimate, 64–72 Mya. The original raw distribution of dated paralogues is indicated by the circles. The *y* axis represents the percentage of gene pairs. **c**, Pruned phylogenetic tree with indication of WGD events (boxes)<sup>29</sup>. The Cretaceous–Palaeogene (K–Pg) boundary is indicated by an arrow.



**Figure 3** | **Reconstruction of metabolic (or gene) pathways involved in the production of stomata, ethylene, terpene and pollen in** *Z. marina***. a**, Stomata differentiation from meristemoid mother cells (MMC) to guard mother cell (GMC) to guard cells. **b**, Ethylene synthesis and signalling up to *EIN2* have disappeared; *EIN3* and its downstream targets remain. **c**, Terpenoid biosynthesis in which the pathways producing volatiles are absent but those essential for primary metabolism remain. MVA, mevalonate; MEP, plastidic methylerythritol phosphate; IPP,

precluding synthesis of secondary volatile terpenes (Supplementary Fig. 6.2). Only aromatic acid decarboxylases (AAAD) genes were expanded (Supplementary Fig. 6.3) and these form a clade distinct from *Spirodela.* The loss of volatiles is also consistent with the loss of stomata, through which they are emitted for airborne communication and plant defence. The repertoire of defence-related genes such as the six groups of NBS\_LRR resistance genes (Supplementary Note 6.2) is also reduced to 44 (89 in *Spirodela* and 100–300 in other plants), which may be linked to a lower probability of infection of *Z. marina* due to the absence of stomata, which are a main entry point for pests and pathogens in terrestrial plants.

Land and aquatic floating plants (Embryophyta) are often exposed to intense ultraviolet (UV) radiation and have developed light sensing protein receptors with protective and signalling functions. In contrast, *Z. marina* inhabits a light-attenuated, submarine environment where it must cope with shifted spectral composition, characterized by low penetration of UV-B, red and far-red wavelengths<sup>16</sup>. Accordingly, *Z. marina* has lost ultravioletresistance (UVR8) genes associated with sensing and responding to UV damage (*Spirodela* has not), as well as phytochromes associated with red/far-red receptors (Supplementary Note 7). Whereas photosystems (PSI and PSII) are similar to those of other plants including *Spirodela*, members of the light-harvesting complex B (LHCB) family are expanded in number, possibly in combination



isopentenyl pyrophosphate; DMAPP, dimethylallyl pyrophosphate; FPP, farnesyl pyrophosphate; GPP, geranyl diphosphate; GGPP, geranylgeranyl pyrophosphate; CPP, copalyl pyrophosphate; GA, gibberellic acid; PP, diphosphate; ABA, abscisic acid. **d**, Sporopollenin biosynthesis genes; regulatory genes in the nucleus control downstream processes (arrows) in response to signalling coming from external stimuli through receptors on the plasma membrane. All panels: genes in red are absent; blue are present; the grey line represents the plasma membrane. See Extended Data Tables 1–3.

with non-photochemical quenching (NPQ), thereby enhancing performance at low light (Extended Data Fig. 4).

Seagrasses typically experience full marine seawater  $(35 g kg^{-1})^{17}$ , whereas land plants obtain water with low osmolality (0-2  $g\text{kg}^{-1}$ ) via the rhizosphere and aquatic plants experience fresh  $(0-5g\,kg^{-1})$  to brackish (0.5–20 g kg<sup>−</sup><sup>1</sup> ) conditions. Although *Z. marina* displays a typical repertoire of Na<sup>+</sup> and K<sup>+</sup> antiporters (Supplementary Note 8, Supplementary Table 8.1), one of six  $H^+$ -ATPase (AHA) genes (Supplementary Table 8.2, Supplementary Data 7) is strongly expressed in vegetative tissue and encodes a salt-tolerant  $H^+$ -ATPase. Furthermore, *Z. marina* possesses three AHA genes (along with *Spirodela*) in a cluster unique to alismatids (Supplementary Fig. 8.1).

Uniquely, *Z. marina* has re-evolved new combinations of structural traits related to the cell wall. Synthesis of cutin-cuticular waxes to the outside of the leaf epidermis and suberin–lignin near the plasma membrane (Supplementary Note 9, Supplementary Table 9.1) surround a cell wall matrix of (hemi)celluloses, low-methylated pectin (zosterin) and macroalgal-like sulfated polysaccharides<sup>18</sup> (Supplementary Note 10). The reduction in carbohydrate-related genes that modify the fine structure of cell wall hemicelluloses and pectins in *Z. marina* is not due to loss of pathways, but rather to the large variation within these CAZyme gene families in plants. Available genomes (including *Spirodela*) lack carbohydrate sulfotransferases and sulfatases, suggesting that land

plants have lost these genes as a key adaptation to terrestrial as well as freshwater conditions19,20. In contrast, *Z. marina* has regained the ability to produce sulfated polysaccharides with an expansion of aryl sulfotransferases (12 genes) homologous to aryl sulfotransferases from land plants (Supplementary Note 10). Sulfation facilitates water and ion retention in the cell wall to cope with desiccation and osmotic stress at low tide and, likewise, low methylation of zosterin correlates with the expanded pectin carbohydrate esterase 8 (CE8) family, increasing the polyanionic character of the cell wall matrix. We speculate that several aryl sulfotransferases have evolved because carbohydrate sulfatases have been shown to be active on artificial aryl compounds such as methylumbelliferyl-sulfate<sup>21</sup>. Osmotic equilibrium is further achieved in *Z. marina* by organic osmolytes (mainly sucrose, trehalose and proline) in combination with a small cytoplasm:vacuole volume ratio (10%)<sup>22</sup>. Given that up to 90% of fixed carbon is stored as sucrose in the rhizomes, sucrose synthase (SuSy) and transport (SUT) genes are expanded while those for starch metabolism are greatly reduced, as expected in 'marine sugarcane' (Supplementary Note 7.2, Supplementary Data 8).

The repertoire of redox and other stress-resistance genes (Supplementary Note 8) is typical for angiosperms with the exception of catalase (CAT), which is reduced to a single copy in *Z. marina* (two in *Spirodela*). Late embryogenesis abundant (LEA) and dehydrins are clearly under-represented in both *Zostera* and *Spriodela* relative to other genomes. In contrast, *Zostera* possesses an unusual complement of metallothioneins. Aside from their role as chelators, metallothioneins may be involved in stress resistance; one of these, MT2L, is among the most highly constitutively expressed genes in *Z. marina* (Extended Data Fig. 5, Supplementary Note 8.2).

Sexual reproduction of *Z. marina* takes place underwater, involving completely submerged male and female flowers, and a unique exine-less, filiform pollen that winds around the bifurcate stigmas in a purely abiotic pollination process<sup>23</sup>. Note that freshwater alismatids (and also Sp*irodela*)<sup>24</sup> possess pollen with an exine layer. Exine-less pollen<sup>25</sup> is characteristic of all seagrasses except *Enhalus acoroides* (which is surface pollinated). Ten genes specifically involved in biosynthesis and modification of the pollen exine coat are missing; all other genes involved in the development of viable pollen remain intact (Fig. 3d, Extended Data Table 3, Supplementary Note 11.1). Finally, MADS-box gene transcription factors are also highly reduced to 50 in *Z. marina*, which is most likely related to its highly reduced flowers (also a feature of *Spirodela)* that lack the first two whorls of specialized floral leaves, calyx and corolla (Supplementary Note 11.2, Supplementary Table 11.2).

An increasing proportion of the world population inhabits the coastal zone. This impinges multiple pressures on ecosystems including seagrass beds<sup>26,27</sup>, which in turn compromises the ecosystem services they may provide, including provisioning of harvestable fish and invertebrates, nutrient retention, carbon sequestration and erosion control. In the context of seagrass conservation, elucidating the genomic basis of *Z. marina*'s complex adaptations to ocean waters (Extended Data Fig. 6) will also inform the development of molecular indicators of their physiological status28, as these unique ecosystems rank, unfortunately, among the most threatened on Earth<sup>26,27</sup>.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the [online version of the paper](http://www.nature.com/doifinder/10.1038/nature16548); references unique to these sections appear only in the online paper.

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**Supplementary Information** is available in the [online version of the paper](http://www.nature.com/doifinder/10.1038/nature16548).

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**Author Contributions** J.L.O., T.B.H.R., G.P. and Y.V.d.P. are the lead investigators and contributed equally to the work. J.S., J.W.J., J.G., Y.V.d.P., B.V. and Y.-C.L. coordinated the bioinformatics activities surrounding assembly, quality control, set-up and maintenance of *Z. marina* on the ORCAE site and deposition of the *Z. marina* genome resource. T.B.H.R and T.B. generated and analysed RNA-seq libraries from flowers, rhizome, roots. J.L.O., Y.-C.L. and A.J. generated and analysed RNA-seq libraries from the genome genotype and temperature stress experiments. C.B., W.T.S. and J.L.O. contributed to biological sample collection, preparation and quality control prior to DNA extraction. A.M. performed the HMW DNA extraction and quality control from the genome genotype/ clone. M.A., J.G., H.T. and M.C. contributed to WGS libraries and sequencing, (fosmid)-cloning and quality control. J.G. coordinated the sequencing of FES, quality control projects. Analysis of architectural features of the genome and annotation of specific gene families, including the written contributions to the main paper and Supplementary Information sections, were performed by the following co-authors: J.W.J., the chromosome assembly analysis; B.V. and Y.-C.L., gene family clustering and comparative phylogenomics; A.R.K. and E.B.B., Pfam domains; E.D.P. and P.J.G., miRNA; R.L., K.V. and Y.V.d.P., wholegenome duplication; F.M., Y.-C.L. and Y.V.d.P., transposable elements; B.V., co-linearity and synteny comparisons; M.T., organellar genomes; P.R., stomata gene family; G.M., cell wall polysaccharides and sulfotransferases; T.T., fatty

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**Author Information** Raw reads, the assembled genome sequence and annotation are accessible from NCBI under BioProject number [PRJNA41721](http://www.ncbi.nlm.nih.gov/bioproject/?term=PRJNA41721) with GenBank accession number [LFYR00000000.](http://www.ncbi.nlm.nih.gov/nuccore/LFYR00000000) The accession number for the *Zostera marina* Finnish Clone is BioSample [SAMN00991190.](http://www.ncbi.nlm.nih.gov/biosample/SAMN00991190) Fosmid end sequence: GSS KG963492-[KG999999](http://www.ncbi.nlm.nih.gov/nucgss/KG999999); [KO000001](http://www.ncbi.nlm.nih.gov/nucgss/KO000001)-K0144970, wholegenome shotgun data: [SRA020075](http://www.ncbi.nlm.nih.gov/sra/?term=SRA020075) and RNA-seq: GEO [GSE67579](http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE67579). Further information on the *Zostera marina* project is available via the Online Resource for Community Annotation Eukaryotes (ORCA) at [http://bioinformatics.psb.ugent.](http://bioinformatics.psb.ugent.be/orcae/) [be/orcae/.](http://bioinformatics.psb.ugent.be/orcae/) Reprints and permissions information is available at [www.nature.](http://www.nature.com/reprints) [com/reprints](http://www.nature.com/reprints). The authors declare no competing financial interests. Readers are welcome to comment on the [online version of the paper.](http://www.nature.com/doifinder/10.1038/nature16548) Correspondence and requests for materials should be addressed to J.L.O. [\(j.l.olsen@rug.nl](mailto:j.l.olsen@rug.nl)) or Y.V.D.P [\(yves.vandepeer@psb.vib-ugent.be\)](mailto:yves.vandepeer@psb.vib-ugent.be).

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#### **Methods**

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

**Plant material and DNA preparation.** A single genotype/clone of *Zostera marina* (referred to as the 'Finnish clone') was harvested on 26 August 2010 at 2m depth at Fårö Island (latitude 59° 55.234′ N longitude 21° 47.766′ E) located in the northern Baltic Sea, Finland. Plant material was transported to the lab in seawater, cleaned and further processed. Care was taken to use leaf-meristem tissue harvested from the inner layer of basal shoots to minimize bacterial/diatom contamination. Tissues were immediately frozen in LN<sub>2</sub> and stored at −80 °C for later DNA and RNA extraction. Monoclonality was verified by genotyping 40 ramets of the mega-clone with six highly polymorphic, microsatellite loci<sup>30</sup>. There was no evidence for polyploidy<sup>25,31,32</sup> (*Z. marina* is 2n = 12) or somatic mutations<sup>33</sup> as assessed by multiple peaks in the microsatellite chromatograms. Tissue was subsequently sent on dry ice to Amplicon Express for HMW DNA extraction using a CTAB isolation method modified by R. Meilan (unpublished) but available from him (rmeilan@purdue. edu), based on the original method<sup>34</sup>. Following QC according to JGI guidelines, the DNA was shipped to JGI for library and sequencing preparation.

**Genome sequencing and assembly.** One 35-Kb, fosmid library was generated for end sequencing. The fosmid ends were sequenced with standard Sanger sequencing protocols at the HudsonAlpha Institute for a total of 194,303 Sanger reads (0.29× coverage). Illumina libraries (two fragment libraries (6.62Gb), one 2-Kb JGI mate-pair library (3.57Gb), one 4-Kb JGI mate-pair library (3.41Gb) and two 8-Kb JGI mate-pair libraries (11.94Gb)) were sequenced with Illumina MiSeq/ HiSeq genetic analysers at the Department of Energy's Joint Genome Institute (JGI), using standard protocols. A total of 25.55 Gb of Illumina and 0.14Gb of Sanger sequence was obtained representing 47.7× genomic coverage. Prior to assembly, all reads were screened against mitochondria, chloroplast, and Illumina controls. Reads composed of > 95% simple sequence repeats were removed. For the Illumina, paired-end libraries ( $2 \times 250$ ), reads <75 bp were discarded, for the  $2\times150$  libraries, reads  $<$  50 bp were discarded after trimming for adaptor and quality  $(q<20)$ . An additional deduplication step was performed on the mate pairs that identified and retained only one copy of each PCR duplicate. A total of 212,101,273 reads (Supplementary Table 2.1) was assembled using our modified version of Arachne v. 20071016 (ref. 35). Subsequent directed Arachne modules were applied to collapse adjacent heterozygous contigs. The entire assembly was then run through another Arachne process starting at Stage 6 Rebuilder. This produced 15,747 scaffold sequences (30,723 contigs), with a scaffold L50 of 409.5Kb, 613 scaffolds larger than 100Kb, and a total genome size of 237.5Mb (Supplementary Table 2.2).

Scaffolds were screened against bacterial proteins, organelle sequences, GenBank NR (nr\_prot) and RefSeq protein databases, and removed if found to be a contaminant. Scaffolds consisting of prokaryotes, chloroplast, mitochondria and unanchored rDNA were removed. We also assembled the chloroplast and partial mitochondrial genomes (Supplementary Notes 2.2 and 2.3, Supplementary Fig. 2.1). Additionally, short  $(<1$  Kb) scaffolds or scaffolds containing highly repetitive sequence (>95% 24-mers found more than four times in large scaffolds) or alternative haplotypes were also removed. Following repeat analysis and gene prediction, all scaffolds were subjected to a filtering process (based on NCBI nr\_prot + NCBI taxonomy database) to eliminate remaining bacterial (and other) contaminants (Supplementary Table 2.3).

Assembly validation was performed using a set of 12 fully sequenced fosmid clones. In 4 of the 12 fosmid clones, full-length alignments were not found due to fragmentation in the region of the fosmid clone. In five of the remaining eight fosmid clones, the alignments were of high quality ( $<$  0.05% bp error). The overall base pair error rate (including marked gap bases) in the fosmid clones that aligned to full length was 0.28% (714 discrepant base pairs out of 253,332bp). Supplementary Table 2.4 shows the individual fosmid clones and their contribution to the overall error rate. Note that two fosmid clones (16248, 16249) contributed nearly 81% of the discrepant bases. This probably occurred in polymorphic regions of the genome where the haplotype in the fosmid did not match the haplotype in the reference. There are several indels of various sizes in the clone and assembly, typical of a region of degraded transposons. Further quality analysis indicated that 90% of the set of eukaryotic core genes (CEGMA) were present and 98% were partially represented, suggesting near completeness of the euchromatin component.

**Annotation of repetitive sequences.** Two complementary approaches were used to identify repetitive DNA sequences in the *Z. marina* genome. With respect to masking repeats before gene prediction analysis, a *de novo* repeat identification was carried out with RepeatModeller (v. open-1.0.7; [http://www.RepeatMasker.](http://www.RepeatMasker.org) [org\)](http://www.RepeatMasker.org)<sup>36</sup> to identify repeat boundaries and build consensus models from which potential over represented, non-transposable element, protein-coding genes were removed. RepeatMasker (v. open-4.0.0, WUBlast) was used in combination with this custom repeat library to mask the assembly and prepare it for gene prediction with EuGene.

Furthermore, in order to perform a qualitative and quantitative analysis of repeats with greater resolution<sup>37</sup> the genome assembly was processed for *de novo* repeat detection using the TEdenovo pipeline from the REPET package v. 2.2 (ref. 38); parameters were set to consider repeats with at least five copies. The consensus sequences generated by TEdenovo were then used as probes for whole genome annotation by the TEannot<sup>39</sup> pipeline from the REPET package v. 2.2. The consensus repeat sequences were classified using Pastec<sup>40</sup>. Comparing the genomic positions of transposable elements (TE) to those of exons from the set of predicted genes enabled us to identify that 909 gene predictions most likely represent TEs and these were filtered from the gene set. The REPET package v. 2.2 was also used to annotate repetitive elements in the *Spirodela polyrhiza* genome assembly with the same parameters as for *Z. marina*. See Supplementary Fig. 3.1.

**Transcriptome library preparation, sequencing and assembly.** Leaf, root and flower tissues were separately frozen in liquid nitrogen immediately following harvest from either ambient (field collected) or experimental (mesocosm) conditions (Supplementary Note 3.2). Overall, we obtained between nine and 20 million high-quality reads from each of the flower-leaf-root replicate libraries; and for the Finnish clone library, 148.5 million high quality reads were retrieved (Supplementary Table 3.3).

The *de novo* assembly protocol was adapted from ref. 41. We pooled replicates of each tissue together except for the two leaf tissue libraries, which were kept separate (Supplementary Table 3.4) and performed *de novo* transcriptome assembly for each tissue using  $Trinity<sup>41</sup>(v. 2014-07-17)$  with digital normalization option ON to normalize input read coverage. Frame shift errors and insertion/deletion errors in the assembled transcripts were corrected by FrameDP<sup>42</sup>. Because a *de novo* assembly still generates many spurious transcripts, we used the transcript expression value to remove low quality contigs. We used the RSEM pipeline<sup>43</sup> to obtain the contig expression values and removed contigs with FPKM (fragments per kilobase of transcript per million fragments mapped) value  $<\!1$  and IsoPct (percentage of expression for a given transcript compared with all expression from that Trinity component)<1. In total, we obtained between 39,000 and 53,000 assembled contigs from each library, and 52,000 contigs from the Finnish clone library (Supplementary Table 3.4). Prior to mapping the genome sequence and the predicted genes, we used the CD-HIT<sup>44</sup> program (v. 4.6.1) to collapse redundant contigs, which resulted in 79,134 low redundant transcript contigs.

**Differential gene expression analysis.** High-quality RNA-seq reads were mapped to the genome assembly v.2.1 by TopHat<sup>45</sup>. Differential gene expression analysis was performed by the Cufflink pipeline45 based on the *Z. marina* v.2.1 gene models by converting the number of aligned reads into FPKM values. Genes with significant expression difference ( $\log_2$  > 2) were selected for further investigation by GOstats46 to perform Gene Ontology (GO) term enrichment analysis with *P*≤0.05 (Supplementary Note 3.3, Supplementary Table 3.5)

**MicroRNA analysis.** Genomic precursors of known miRNAs were mapped on the *Z. marina* genome following the procedure described in ref. 47 for the maize genome. miRNA entries from the miRBase database (release 21, 2014) were aligned to the chromosomes of the *Z. marina* genome. Up to three mismatches were allowed in the alignment, using SeqMap<sup>48</sup>. In parallel, novel potential DCL1/ AGO1-dependent miRNAs were enriched by selecting 5′-U 20–22nt small RNAs from three different sequenced libraries from *Z. marina* described in ref. 12. A subset of these small RNAs with abundance >10 TPM (transcripts per million) was retained and aligned to the genome with no mismatches. From every locus, we extracted two  $\sim$ 200-nt regions surrounding each aligned miRNA or candidate (from  $-30$  to  $+160$  and from  $-160$  to  $+30$  nucleotides relative to the putative miRNA start or end coordinate, respectively). Minimum energy RNA secondary structures were predicted for each region using the RNAfold program of the Vienna RNA 1.8.5 package [\(http://www.tbi.univie.ac.at/](http://www.tbi.univie.ac.at/~ivo/RNA/)~ivo/RNA/) using default settings.

In addition, small RNAs from the three sequenced libraries were mapped on these regions, allowing no mismatches, in order to pre-select putative miRNA loci that showed evidence of expression in the three plant tissues analysed. We evaluated RNA structure and small RNA alignment in all the regions based on: (1) dominance of plus-stranded small RNAs; (2) position of the most abundant small RNAs relative to the predicted miRNA coordinates; (3) prevalence of 20–22nt small RNAs in the predicted miRNA locus; (4) position of the putative miRNA with the stem-loop structure; and (5) absence of oversize ( $\geq$ 3 nt) bulges in the miRNA/ miRNA\* alignment. After reduction of overlapping loci to a non-redundant set and removal of stem-loop structures with the wrong orientation compared to miRNAs registered in miRBase, we manually inspected the remaining loci to further evaluate them according to the miRNA annotation criteria proposed by ref. 49. Stringency was relaxed when small RNA expression data strongly indicated the presence of miRNA loci that did not meet the whole set of criteria. Novel miRNA precursors overlapping with TEs or other repetitive elements were filtered out.

Potential miRNA targets were identified *in silico* using the generic small RNAtranscriptome aligner GSTAr from the CleaveLand package  $(v. 4)^{50}$ . Predicted targets were accepted with an Allen score <4 or a MFE (minimum free energy) ratio≥7.5. (Supplementary Note 3.4).

**Gene prediction.** Training of the gene prediction programs started with the collection of high quality ESTs. EST information was used, for example, to train the splice predictor SpliceMachine<sup>51</sup>. Detection of conserved splice sites was further investigated by RNA-seq splice junctions (count  $>$  10) to construct a WAM model in EuGene (v.  $4.1$ )<sup>52</sup>. Coding-potential was modelled with an interpolated Markov Model (IMM) constructed from the BLASTX alignments of proteins from the PLAZA v. 2.5 database<sup>53</sup>. An additional protein 'monocot' Markov Model was built based on the protein sequences from *Brachypodium*, maize and sorghum. Starting from EST and protein alignments, a set of 215 gene models was manually constructed and curated using the genome browser GenomeView<sup>54</sup>. The 215 models were then used as a training set for EuGene in order to optimize the different splice site and coding-potential models, as well as the weights for the extrinsic EST and homology evidence. An overall fitness score of 80.1% was achieved, which is high enough to obtain reliable results without overfitting. GeneMark<sup>55</sup> and Augustus<sup>56</sup> were separately trained (using the same input data as EuGene) and their predictions were integrated with EuGene using a custom script to evaluate the best gene structure at each locus. All gene models were automatically screened to highlight possible erroneous structures (for example, in-frame stop codons, deviating splice junctions) and manually curated. Transfer-RNA gene models were predicted by tRNAscan-SE (v. 1.31)<sup>57</sup> and their structures were verified with Infernal (v. 1.1rc1, rfam11 covariant model database)<sup>58</sup>. For each gene, UTRs were assigned by identifying a set of ESTs and RNA-seq assemblies that uniquely overlapped with it. We subsequently selected the longest mapped transcript on either end of the predicted coding sequence and designated the section outside the coding sequence as the UTR. Finally, all genes were uploaded to the ORCAE platform ([http://bioinfor](http://bioinformatics.psb.ugent.be/orcae) [matics.psb.ugent.be/orcae\)](http://bioinformatics.psb.ugent.be/orcae) 59, enabling all members of the consortium to refine and curate the gene model and assign gene function. A list of protein domains, as well as the derived Gene Ontology (GO) terms and KEGG pathway identifiers were generated using an InterProScan (v. 5.2.45)<sup>60</sup> analysis and available in ORCAE. More specifically, gene functional descriptions were added either manually by consortium expert scientists or automatically through sequence homology searches. The automated method relies on the EC (Enzyme Commission) number reported by InterProScan to retrieve the enzyme name with BLASTP search against UniProtKB/Swiss-Prot<sup>61</sup> to filter out hits that are below 60% identity and 70% query/hit coverage. Although such high stringency on per cent identity and sequence coverage reduced the available number of functional descriptions, it reduced the false-positive prediction rate, as desired here.

**Construction of age distributions and WGD analyses.** *K*<sub>S</sub>-based age distributions were constructed as previously described<sup>62</sup>. In brief, the  $K_S$  values between genes were obtained through maximum likelihood estimation using the CODEML program<sup>63</sup> of the PAML package (v. 4.4c)<sup>64</sup>. Gene families for which *K*<sub>S</sub> estimates between members did not exceed a value of 5 were subdivided into subfamilies. For each duplicated gene in the resulting phylogenetic gene tree, obtained by PhyML<sup>65</sup>, all  $m$  $K_S$  estimates between the two child clades were added to the  $K_S$  distribution with a weight  $1/m$  (where  $m$  is the number of  $K_S$  estimates for a duplication event), so that the weights of all  $K_S$  estimates for a single duplication event summed to one. Mixture modelling was used to confirm a WGD signature in the  $K_S$  distribution (Fig. 2 and Supplementary Fig. 4.1), for which all duplicates with  $K_S$  values  $\leq 0.1$  were excluded to avoid the incorporation of allelic and/or splice variants, while all duplicates with  $K_S$  values  $>$  2.0 were removed because  $K_S$  saturation and stochasticity can mislead mixture modelling above this range<sup>62</sup>. For further details see Supplementary Note 4.1.

Absolute dating of the identified WGD event was performed as described previously13,29. In brief, paralogueous gene pairs located in duplicated segments (anchors) and duplicated pairs lying under the WGD peak (peak-based duplicates) were collected for phylogenetic dating. Anchors, assumed to be corresponding to the most recent WGD, were detected using i-ADHoRe 3.0 (refs 66,67). Only a low number of duplicated segments and hence anchors could be identified, most likely because of the fragmented assembly of *Z. marina.* However, the identified anchors did confirm the presence of a broad WGD peak between a  $K_S$  of 0.8 and 1.6 (data not shown). For each WGD paralogueous pair, an orthogroup was created that included the two paralogues plus several orthologues from other plant species as identified by InParanoid (v.  $4.1$ )<sup>68</sup> using a broad taxonomic sampling: one representative orthologue from the order Cucurbitales, two from the Rosales, two from the Fabales, two from the Malpighiales, two from the Brassicales, one from the Malvales, one from the Solanales, two from the Poales, one orthologue from *Musa acuminata*69 (Zingiberales), and one orthologue from *Spirodela polyrhiza*11 (Alismatales). In total, about 180 orthogroups from anchor pair duplicates and peak-based duplicates were collected. The node joining the two *Z. marina* WGD paralogues was then dated using the BEAST v.  $1.7$  package<sup>70</sup> under an uncorrelated

relaxed clock model and a LG+G (four rate categories) evolutionary model. A starting tree with branch lengths satisfying all fossil prior constraints was created according to the consensus APGIII phylogeny<sup>71</sup>. Fossil calibrations were implemented using log-normal calibration priors on the following nodes: the node uniting the Malvidae based on the fossil *Dressiantha bicarpellata*72 with prior offset = 82.8, mean = 3.8528, and s.d. =  $0.5$ )(ref. 73), the node uniting the Fabidae based on the fossil *Paleoclusia chevalieri*74 with prior offset=82.8, mean=3.9314, and s.d.=0.5(ref. 75), the node uniting the Alismatales (including *Z. marina* and *Spirodela polyrhiza*) with the other monocots based on the oldest fossil mono- $\cot$  pollen, *Liliacidites*<sup>76,77</sup> from the Trent's Reach locality, with prior offset = 125, mean = 2.0418, and s.d. = 0.5 (refs 14,78) and the root with prior offset = 124, mean =  $4.0786$ , and s.d. = 0.5 (ref. 79). The offsets of these calibrations represent hard minimum boundaries, while their means represent locations for their respective peak mass probabilities in accordance with some of the most recent and taxonomically complete dating studies available for these specific clades $14,80$ . A run without data was performed to ensure proper placement of the marginal calibration prior distributions<sup>81</sup>. The Markov chain Monte Carlo (MCMC) for each orthogroup was run for  $10^6$  generations, sampling every 1,000 generations resulting in a sample size of  $10<sup>4</sup>$ . The resulting trace files of all orthogroups were evaluated manually using Tracer v. 1.5<sup>70</sup> with a burn-in of 1,000 samples to ensure proper convergence (minimum ESS for all statistics at least 200). In total, 169 orthogroups were accepted and all age estimates for the node uniting the WGD paralogous pairs were then grouped into one absolute age distribution (Fig. 2, too few anchors were available to evaluate them separately from the peak-based duplicates), for which kernel density estimation (KDE) and a bootstrapping procedure were used to find the peak consensus WGD age estimate and its 90% confidence interval boundaries, respectively.

Intra- and inter-genomic co-linearity was investigated (Supplementary Tables 4.2 and 4.3) using MCScanX82 based on a BLASTP search of all genomic protein coding genes with an E-value cut-off of  $e^{-10}$ . Only one large duplicated segment was detected, which was most likely due to the fragmented assembly of *Z. marina*; only 27 scaffolds had a size larger than 1 Mb, accounting for only 23.4% of all protein-coding genes. We therefore additionally used i-ADHoRe (v. 3.0)<sup>66</sup> to investigate genomic co-linearity by including all possible scaffolds.

**Gene family comparisons.** Protein sets were collected for 14 species: *Z. marina* (ORCAE v. 2.1), *Arabidopsis thaliana* (TAIR10), *Thellungiella parvula* ([http://thel](http://thellungiella.org) [lungiella.org](http://thellungiella.org)) *Populus trichocarpa* (Phytozome v. 9.0), *Vitis vinifera* (Phytozome v. 9.0), *Amborella trichopoda* (<http://amborella.huck.psu.edu>), *Oryza sativa* japonica (Phytozome v. 9.0), *Zea mays* (Phytozome v. 9.0), *Brachypodium distachyon* (Phytozome v. 9.0), *Spirodela polyrhiza* (<http://mocklerlab.org>), *Selaginella moellendorffii* (Phytozome v. 9.0), *Physcomitrella patens* (Phytozome v. 9.0), *Chlamydomonas reinhardtii* (Phytozome v. 9.0), and *Ostreococcus lucimarinus* (ORCAE v. 6/3/2013). These species were selected in order to provide a phylogenetic representation traversing green algae, basal plants, monocots, and dicots. Following an 'all-vresus-all' TimeLogic Decypher Tera-BLASTP (Active Motif Inc.; e-value threshold  $1 \times e^{-3}$ , max hits 500) comparison, OrthoMCL (v. 2.0; mcl inflation factor  $3.0)^{83}$  was used to delineate gene families. Confidence in establishing gene losses in *Zostera* was enhanced by using a combination of reciprocal blast, TblastN, re-annotation of *Spirodela* (and other monocot genes), and careful phylogenetic analysis. OrthoMCL results and related protein resources are available in the ORCAE download section.

To further understand gene family expansion or contraction in *Z. marina* in comparison with other sequenced genomes, gene family sizes were calculated for all gene families (excluding orphans and species-specific families) (Supplementary Note 4.2). The number of genes per species for each family was transformed into a matrix of *z*-scores in order to centre and normalize the data. The first 100 families with the largest gene family size in *Z. marina* were selected. The *z*-score profile was hierarchically clustered (complete linkage clustering) using Pearson correlation as a distance measure. The functional annotation of each family was predicted based on sequence similarity to entries in the InterProScan and Pfam protein domain database where more than 30% of proteins in the family share the same protein domain. The phylogenetic profile and phylogenetic tree topology provided at PLAZA<sup>84</sup> were used to reconstruct the most parsimonious series of gene gain and loss events. The Dollop program from the PHYLIP package<sup>85</sup> was used to determine the minimum gene set at ancestral nodes of the phylogenetic tree. The Dollop program is based on the Dollo parsimony principle, which assumes that novel gene families arise exactly once during evolution but can be lost independently in different phylogenetic lineages.

**Search for presence/absence of orthologues for specific genes and families.** A dedicated search for orthologues/homologues was performed for genes and proteins involved in stomata differentiation (Supplementary Note 5.1), volatile biosynthesis and sensing with focus on ethylene and terpenes (Supplementary Note 6.1), as well as genes involved in male flower specification and pollen

differentiation (Supplementary Note 11.1). To this end, queries were chosen from documented genes involved in these pathways (usually from *Arabidopsis* but occasionally from *Oryza*, *Zea* and tomato). Next, the search for homologues in *Zostera marina*, *Spirodela polyrhiza*, *Oryza sativa japonica* and *Arabidopsis thaliana* (when not used as a query) was performed using BLASTP. To avoid missing or poorly annotated genes a TBLASTN search was conducted using the above queries against the *Zostera marina* and *Spirodela polyrhiza* genomes. Putative orthologues were identified based on reciprocal BLASTP searches with *Arabidopsis* (or the other queries). Owing to speciesspecific duplications, this may produce some paralogous genes to appear orthologous to the query, or vice versa (see Extended Data Tables 1–3). To further confirm correct orthology assignments, phylogenetic trees were built using a broader sampling of protein sequences from both the query species and the three target species. Ambiguously aligned sequences (especially due to indels) were checked manually and corrected or removed.

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**Extended Data Figure 1** | **Number of genes expressed in five tissues of** *Z. marina***. a**, Venn diagram of genes with expression values (FPKM) higher than 1 are considered as expressed in the tissue. **b**, Pairwise differential gene expression analysis between tissues. The male flower shows the highest number of differentially expressed genes.



**Extended Data Figure 2** | **Circos plot of the ten largest scaffolds of** *Z. marina***.** Tracks from outside to inside. GC percentage, gene density, and transposable element (TE) density (density measured in 20-Kb sliding windows and gene expression profiles from five tissues (root, leaf, male flower, female flower early and female flower late) presented as  $\log_2$  FPKM values.



**Extended Data Figure 3** | **Potential impact of transposable elements (TEs) on** *Z. marina* **evolution. a**, Frequency distribution of pairwise sequence identity values between copies of Copia- and Gypsy-type LTR retrotransposons and DNA transposons, and their cognate consensus sequences (younger repeats share higher sequence similarity). Two peaks are detectable for Copia-type elements. **b**, Distance to the closest TE for

the set of *Z. marina* single-copy genes and the set of *Z. marina* accessory genes. TE-proximal accessory genes are more frequent than TE-proximal single-copy genes. **c**, Frequency of pairwise sequence identity between accessory gene-proximal Ty3-Gypsy elements and their cognate consensus sequences. A number of high-identity copies (that is, putatively young duplicate genes) is observed.



**Extended Data Figure 4** | **Unrooted maximum likelihood tree of genes encoding light-harvesting complex A (LHCA) and LHCB proteins of** *Z.marina***,**  *Spirodela polyrhiza* **and** *Arabidopsis thaliana***.** The analysis was carried out on protein sequences using PhyML 3 with LG substitution model and 100 bootstrap replicates. Supplementary Note 7.1, Supplementary Table 7.3.

### LETTER RESEARCH



**Extended Data Figure 5** | **Alignment of metallothionein (MT) and half-metallothionein (HMT) genes in** *Z. marina* **as compared with other plants.** Alignments were performed in ClustalW on the Lyon PBIL web server and edited manually. The upper alignments are for type 1–3 MTs and HMTs; the lower alignment is for type 4 EcMTs where there is no *Zostera* homologue. Conserved residues are shown in red and residues in the same amino acid group in blue. Cys and His residues, putatively involved in binding metals, are highlighted in green and yellow, respectively. Aromatic amino acids absent in canonical animal MTs are highlighted in grey. MTs and MT-like proteins were obtained from: *Arabidopsis thaliana* (ARATH), Japanese rice (ORYSJ), *Cicer arietinum* (CICAR), banana (MUSAC), wheat (WHEAT), potato (SOLTU), *Setaria Italica* (SETIT), *Vitis vinifera* (VITVI) and the alismatids: *Posidonia oceanica* (POSOC) highlighted in grey, *Spirodela polyrhiza* (SPIPO) highlighted in blue, and *Zostera marina* (ZOSMA) highlighted in yellow. See Supplementary Note 8.2.





**Extended Data Figure 6** | **Conceptual summary of physiological and structural adaptations made by** *Z. marina* **in its return to the sea.** Ecosystem services shown in blue. Physical processes related to salinity, light and CO<sub>2</sub> availability shown in white within light-green boxes. Gene losses and gains associated with morphological and physiological processes shown in white within the dark-green box on the right.

## LETTER RESEARCH

**Extended Data Table 1** | **Genes involved in stomata development in** *Z. marina* **compared to other angiosperms**



The genes documented to be involved in stomatal development in *Arabidopsis*86 were used as queries to find orthologues in rice and *Siprodela polyrhiza* (duckweed). See Supplementary Note 5.1, Supplementary Fig. 5.1 for sequence alignment and phylogenetic tree. NF-1, not found, supported by phylogeny; NF-2, not found, unambiguous reciprocal BlastP; NC, not conserved.

\*BASL is not evolutionarily conserved, precluding the finding of its homologue in monocots, if it would exist.

†PAN genes have been searched for using the documented PAN1 and PAN2 genes from maize as baits.

#### **Extended Data Table 2** | **Ethylene-responsive transcription factor genes (ERF) in** *Zostera marina*



MF, male flowers; FFE, female flowers early; FFL, female flowers late; R, roots; L, leaves; NF-1, not found as supported by reciprocal Blast and phylogeny. See Supplementary Note 6.1, Supplementary Fig. 6.1 for sequence alignment and phylogenetic tree. Grey indicates genes not involved in ethylene biosynthesis and signal pathways but strongly co-expressed, indicative of multiple functions.

#### **Extended Data Table 3** | **Genes involved in pollen development of** *Z. marina* **compared to other angiosperms**



The five genes encoding proteins associated on the ER-located sporopollenin metabolon in *Arabidopsis*87 are highlighted in grey. The genes documented to be involved in pollen development in *Arabidopsis* or in rice were used as queries to find orthologues. MF, male flowers; FFE, female flowers early; FFL, female flowers late; R, roots; L, leaves; NF-1, not found, supported by reciprocal Blast and phylogeny; NF-2, not found, single copy gene; amb, ambiguous with homologues too similar to point to a specific orthologue. See Supplementary Note 11.1, Supplementary Fig. 11.1 for sequence<br>alignment and phylogenetic t