ETOILE Regulates Developmental Patterning in the Filamentous Brown Alga Ectocarpus siliculosus $^{\text{m}}$

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Brown algae are multicellular marine organisms evolutionarily distant from both metazoans and land plants. The molecular or cellular mechanisms that govern the developmental patterning in brown algae are poorly characterized. Here, we report the first morphogenetic mutant, etoile (etl), produced in the brown algal model Ectocarpus siliculosus. Genetic, cellular, and morphometric analyses showed that a single recessive locus, ETL, regulates cell differentiation: etl cells display thickening of the extracellular matrix (ECM), and the elongated, apical, and actively dividing E cells are underrepresented. As a result of this defect, the overrepresentation of round, branch-initiating R cells in the etl mutant leads to the rapid induction of the branching process at the expense of the uniaxial growth in the primary filament. Computational modeling allowed the simulation of the etl mutant phenotype by including a modified response to the neighborhood information in the division rules used to specify wild-type development. Microarray experiments supported the hypothesis of a defect in cell–cell communication, as primarily Lin-Notch-domain transmembrane proteins, which share similarities with metazoan Notch proteins involved in binary cell differentiation were repressed in etl. Thus, our study highlights the role of the ECM and of novel transmembrane proteins in cell–cell communication during the establishment of the developmental pattern in this brown alga.

INTRODUCTION

Brown algae are multicellular organisms that live along the coasts all over the world under tropical, temperate, and arctic climates. Their sizes are diverse, ranging from microscopic filaments (*Laminaria* gametophytes) to up to 50-meter-long giant kelps (*Macrocystis* sp). They also display a wide range of morphologies, spanning from uniseriate filaments to blades with conic, plain, or digitated flat shapes.

Brown algae belong to the heterokont phylum, also comprising oomycetes and diatoms (Baldauf, 2008). It is estimated that this phylum emerged more than one billion years ago from a unicellular eukaryotic ancestor common to all the eukaryotic lineages known today (Yoon et al., 2004). Therefore, the very distant phylogenetic relationship between brown algae and other multicellular organisms, namely, the archaeplastida (land plants and red and green algae) and the opisthokonta (metazoan and fungi), raises the possibility that these plants developed distinct cellular mechanisms to achieve multicellular development.

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The brown algae *Fucus* and *Pelvetia* were previously used to study the establishment of plant zygote polarity during the first divisions of the embryo (Hable and Kropf, 2005; reviewed in Kropf, 1992). During the first divisions, the cytoskeleton and associated proteins are highly implicated in the establishment of cell polarity, in addition to a Rho family GTPase (Fowler et al., 2004). At later embryonic stages, cell wall–mediated intercellular communication processes seem to participate in the determination and maintenance of cell fate in the fucoid embryo (Berger et al., 1994; Bouget et al., 1998), but the nature of the underlying molecular components remains unknown.

Despite their use as models for embryo polarization, mature thalli of fucoid algae cannot be cultured in vitro and are therefore not appropriate models to study postembryonic development. Furthermore, genomic and genetic tools are not available. By contrast, the emerging brown alga model *Ectocarpus siliculosus* (Peters et al., 2004; Charrier et al., 2008) displays both a body and a reproductive cycle amenable to morphogenetic and genetic studies (Peters et al., 2008). In addition, its genome is sequenced (Cock et al., 2010) and proteomic (Ritter et al., 2010) and transcriptomic tools such as microarrays (Dittami et al., 2009) and real-time PCR normalization genes (Le Bail et al., 2008b) are available.

The early developmental pattern of wild-type *E. siliculosus* sporophytes was previously described (Le Bail et al., 2008a). Despite a high level of morphological plasticity, several constants were observed. After zygote germination, *E. siliculosus*

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sporophytes grow a uniaxial prostrate filament composed of two cell types, E (for elongated) and R (for round). E cells are always located at apical and subapical positions, and R cells are grouped in the center of the filament. Filament growth is ensured essentially by apical E cell division and elongation, while R cells are produced by progressive subapical E cell differentiation. Subsequent axillary branching occurs mainly in the central part of the filament. This overall prostrate filamentous body then differentiates erect filaments, which will later carry the sporangia. Previous computer modeling showed that the developmental patterning that takes place at the early filament stages could be accounted for by local positional information based on the recognition of neighboring cells (Billoud et al., 2008). The progression of the developmental program (branching and shift to the reproductive phase) depends on the phytohormone auxin, which suggests an overall regulation of the developmental pattern based on auxin-mediated positional information (Le Bail et al., 2010). Analyses of morphologically affected mutants were then initiated to further decipher the mechanisms involved in the regulation of the morphogenesis in brown algae. Here, we report a morphogenetic mutant of a brown alga, *e´toile* (*etl*), in *E. siliculosus*. We provide evidence that cell differentiation is altered in this mutant, resulting in an overall modification of its early developmental pattern, and point to a probable defect in the interplay between the extracellular matrix (ECM) and novel transmembrane proteins containing a Notch domain, whose presence and role previously have only been identified in animals.

RESULTS

Morphogenesis in the Brown Alga E. siliculosus Is Regulated by the Recessive Single Locus ETL

The life cycle of *E. siliculosus* is biphasic, heteromorphic, and dioecious (Figure 1). Both male and female unfertilized gametes are able to generate haploid sporophytes called parthenosporophytes (reviewed in Charrier et al., 2008), which are particularly amenable to mutant selection. Gametes from the male strain Ec32 were irradiated by UV-B, and after 2 weeks, the mutant *etl* was isolated. It displayed a hyperbranched phenotype, which remained stable for at least six parthenogenetic generations. *etl* gametophytes displayed no morphological variation compared with wild-type gametophytes.

The mutant *etl* was then crossed with a female wild-type strain (Ec568). The heterozygous state of the F1 was tested by PCR amplification of a microsatellite marker, which exhibits length polymorphism between the two strains (Figure 2A). The 10 zygotes obtained from this cross all showed a wild-type phenotype, indicating that the *etl* mutation is recessive (Figure 2B).

To analyze the segregation of the *etl* mutation in the subsequent generations, 12 unilocular sporangia, each of them the site of a single meiotic event, were isolated from the heterozygote diploid sporophyte (Figure 1F). For each of them, 26 to 74 gametophytes were subisolated, and the morphology of the corresponding parthenosporophytes grown from the germination of unfertilized gametes (parthenogenesis) was analyzed. Altogether, 263 gametophytes produced parthenosporophytes with a mutant phenotype, and 275 gametophytes produced parthenosporophytes with a wild-type phenotype (Table 1). The results obtained were consistent with a 1:1 segregation ratio and a Mendelian inheritance of a single locus mutation $(x^2 \text{ test}, df = 90, x^2 = 96, P \text{ value} = 0.313).$

Additional crosses were performed to further analyze the inheritance of the *etl* mutation. The sex of nine mutant gametophytes derived from the F1 was determined by attempting to cross them with male Ec32 and female Ec568 gametophytes. Four mutant gametophytes could be crossed only with the male Ec32 strain and five only with the female Ec568 strain, demonstrating that the *etl* mutation segregated independently of the sex locus.

Higher Branching Activity Increases etl Overall Growth Rate

Wild-type filaments of young *E. siliculosus* sporophytes (prostrate filaments) first grow as a linear primary filament (PF; Figure 3A) before lateral branching occurs mainly in the center of the filament giving rise to secondary filaments (SFs; Figure 3B) that grow parallel to the surface at the early stages (Figure 3C). By contrast, *etl* growth was characterized by an early arrest in apical growth and an earlier emergence of numerous branches all along the PF (Figures 3D and 3E), ultimately producing a bushy organism growing in all spatial dimensions (Figure 3F). An overall growth curve showing the number of cells as a function of time demonstrated that *etl* kept on growing faster than the wildtype filament 3 d after germination of the mitospores (Figure 3G). Detailed analyses of both the position and the time of emergence of the SFs can account for this increase in the growth rate. Indeed, while in wild-type filaments branching occurs preferentially in the central part of the filament, in *etl*, branches emerge with similar probabilities all along the filament (Figure 3H). In addition, observations of the developmental stage at which SF emergence occurs showed that branches develop earlier in *etl* than in the wild type. First and second SFs emerge at the 11- and 14-cell stages, respectively, in the wild type, while they emerge at the 5- and 8-cell stages in *etl* (Figures 3I and 3J). Finally, the emergence of branches was also more frequent in the *etl* mutant than in the wild type, with the PF growing on average 2.2 cells between the first and the second emergences, compared with 3.6 cells in the wild type.

Auxin, which was shown to modulate wild-type *E. siliculosus* branching pattern (Le Bail et al., 2010), had no noticeable effect on *etl* development.

etl Regulates Cell Differentiation

At the early stages of development, two cell types can be differentiated in filaments of wild-type sporophytes: R cells with a length/width ratio \leq 1.5 and E cells with a ratio > 1.5 (Figure 3A; Le Bail et al., 2008a). In *etl*, cell size and shape were significantly altered compared with the wild type, with E cells being chunkier and R cells being bigger (Figure 4A; P < 0.01). While in the wild type the two cell types were present in the same proportion within the filament (50% each; Le Bail et al., 2008a), in *etl* 90% of the cells were R (P < 0.01 for each stage;

Figure 1. Mutagenesis and Genetic Characterization of the Mutant *etl*.

The life cycle of *E. siliculosus* can be both sexual (left) and asexual (right).

(A) Unfertilized gametes can generate haploid sporophytes called parthenosporophytes.

(B) Gametes from the strain Ec32 were irradiated with UV-B, and the *etl* mutant was isolated after 2 weeks of growth.

(C) The phenotype was stable for at least six parthenogenetic generations produced via germination of mitospores released from plurilocular sporangia. (D) *etl* spores were isolated from unilocular sporangia to generate gametophytes, which displayed no morphological variation compared with wild-type gametophytes.

(E) Male *etl* gametes were crossed with female gametes of the strain Ec568 and the F1 generation displayed a wild-type phenotype.

(F) Unilocular sporangia were isolated from the F1, and the genotype of the spores was identified from the phenotype of the corresponding parthenosporophyte.

Figure 4B). Interestingly, in spite of this difference, the evolution in the cell type proportion was similar in the wild type and in *etl* during the entire experiment (up to the 20-cell stage). In both cases, the cell type proportion increased progressively from the 3-cell stage to the 12-cell stage, where it reached a plateau. These data indicate that *etl* alters the cell type proportion, without altering its evolution and its stabilization as the filament grows.

Furthermore, *etl* displayed a modification of several structural features of the cell. Transmission electron microscopy (TEM) showed that, in addition to the general diameter increase observed in *etl* cells, the ECM looked thicker compared with the wild type (Figures 4C and 4D; details in Figures 4E and 4F). Measurements on TEM sections (*n* > 80) confirmed that the ECM was significantly thicker in *etl* than in the wild type (*t* test, P value < 0.001; Figure 4G). Moreover, when observed by scanning electron microscopy, *etl* cells displayed a less regular and a rougher surface than wild-type cells (Figures 4H and 4I). However, the organization of the ECM did not seem to be modified in *etl* cells, as cellulose fibers were still observed parallel to the cell surface, and wild-type and *etl* ECM materials had a similar density (Figures 4E and 4F). In addition to a thicker ECM, *etl* cells displayed a slightly modified Golgi apparatus: it appeared larger and often encircled the cell nucleus (4% of *etl* cells displayed a discrete Golgi compared with 24% in the wild type, $n = 30$; Figures 4J and 4K). Furthermore, a higher proportion of *etl* cells displayed a low number of cisternae (25% of *etl* cells contained less than three cisternae per Golgi, compared with 10% in the wild type; Figures 4J and 4K). At their extremities, cisternae displayed large white vesicles at a higher proportion in *etl* cells (56%) than in wild-type cells (40%; Figure 4K), and this was accompanied by an abundant *cis*-Golgi. Therefore, the higher abundance of Golgi stacks with a reduced number of cisternae bearing big white budding vesicles indicate that the Golgi apparatus is more active in *etl* cells than in the wild type. This probably accounts for the thicker ECM, as the main components of the brown algal ECM (i.e., alginate and fucoidan polysaccharides) are transported to the ECM via Golgi-derived vesicles (Nagasato and Motomura, 2009).

In addition to changes in the proportion of R and E cells, the position of these cell types along the filament was altered. Figure 5 shows their distribution in >100 wild-type or *etl* primary filaments grown in similar conditions and observed from the 2-cell stage to the 10-cell stage. Because of the high level of morphological plasticity in this alga, wild-type filaments displayed a large range of combinations. However, a preference for filaments with R cells located in the center and always adjacent to each other was observed in the wild type (Le Bail et al., 2008a). In *etl*, the filament population was different. While some combinations were shared by the wild-type and *etl* populations, many others were specific to *etl*. In contrast with the wild type, *etl* R cells were distributed along the entire filament, and in rare but significant cases, R cells were separated by E cells. Altogether, these observations show that, in addition to the higher proportion of R cells and the modification of some of their subcellular features, tissue patterning in *etl* was impaired, as both the position of R cells along the filament and their clustering were affected.

Figure 2. The *etl* Locus Is Recessive.

The male mutant *etl* (genetic background Ec32) was crossed with the female strain Ec568.

(A) Ten F1 individuals were tested by PCR for the presence of the polymorphic microsatellite marker 427 (40 bp difference; Heesch et al., 2010) between the two strains (only five shown).

(B) The 10 F1 individuals displayed a phenotype similar to that of the progeny of the control cross Ec568 \times Ec32, showing the recessive character of the *etl* mutation. Bars = 100 µm.

Table 1. Segregation Analysis of Gametophytes Derived from

Computational Simulation of etl Development Suggests Impairment in Local Positional Information

To generate a hypothesis of the mechanisms perturbed in *etl*, we modeled its development attempting to reproduce the celltype distribution observed at each stage of filament growth (from the 2- to the 10-cell stages). Both the types of distribution and their frequency of occurrence were taken into account (Figure 5). The early developmental pattern of *E. siliculosus* wildtype sporophyte filaments was previously modeled using the cellular automaton ECTOMAT (Billoud et al., 2008). ECTOMAT uses five transition rules, based on the cellular processes taking place during the development of the alga in vivo (taking into account both cell division and cell differentiation rules: I, division of an R cell with at least one free side; E, E cell differentiation from an R cell; A, apical division of an E cell; R, R cell differentiation from an E cell; and D, simultaneous division and differentiation of an E cell into two R cells; Billoud et al., 2008; Figures 6A and 6B). These processes were modeled as stochastic transitions between cell states, which make use of information about the cellular neighborhood of the cell undergoing the transition. For example, rules R and D occur only if the considered cell is an E cell and if it has one R cell on one side and another cell (of any type) on the other side. Likewise, rule A depicts the apical division and is modeled as a transformation of an E cell into two E cells, provided that the initial E cell is bordered by one cell and an empty space. As these cellular transitions were also observed in the *etl* mutant, we investigated whether *etl* development patterning could be accounted for by changing only the probability of occurrence of these transition rules. For both wild-type and *etl* modeling, the probabilities were optimized for the distribution of R cells counted at each filament stage and were then used to simulate the generation of the population observed in Figure 5 (both filament pattern and frequency of occurrence at each filament stage). Figure 6B displays the best probabilities obtained for both wild-type and *etl* filament populations. It shows that, while the optimization heuristic method managed to simulate the wild-type population (low indicator of discrepancy $=$ 40.6), it failed to obtain a set of probabilities able to simulate the observed *etl* population (high indicator of discrepancy = 250.4). This means that the transition rules used to simulate the development of wild-type *E. siliculosus* were not adequate and that the automaton rules themselves needed to be changed.

In vivo observations revealed that, compared with the wildtype organism, *etl* undergoes additional cellular transitions during its early development. Figure 6A shows that the three additional rules (dashed lines in red) all contribute to the generation of more R cells, in agreement with the previous description of *etl* filaments. These rules were incorporated in the ECTOMAT model, and the probabilities of the now eight transition rules were optimized as described above (Figure 6B). This approach allowed the successful simulation of the in vivo *etl* population at all the developmental stages (low indicator of discrepancy = 3.5). As an example, Figure 6C provides a comparison of the observed

Individual Unilocular Sporangia from One F1 (Ec568 3 *etl*) Sporophyte

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Unilocular Sporangium	No. of et/ Gametophytes	No. of Wild-Type Gametophytes
1	$13*$	$13*$
$\overline{2}$	$32*$	$42*$
3	$14*$	$22*$
4	$14*$	$23*$
5	$20*$	$14*$
6	30	16
7	$23*$	$13*$
8	$34*$	$28*$
9	$14*$	$18*$
10	19	37
11	40	20
12	$22*$	$17*$
Total	263*	$275*$

Segregation analysis was performed from 12 meiotic events (one event per unilocular sporangium). Twenty-six to 60 *etl* or wild-type gametophytes were isolated from each sporangium and grown to produce parthenosporophytes, on which the phenotype was observed. x^2 tests show that the population does not significantly deviate from a 50% (wild type)/50% (*etl*) segregation, indicating that a single mutated locus is responsible for the *etl* phenotype. Asterisk indicates that the value does not deviate significantly from 1:1 (χ^2 test, P > 0.05).

(A) to (F) Morphology of wild-type (A, B, C) and *etl* (D, E, F) sporophytes. Sporophytic wild-type PFs are composed of two cell types, namely, the elongated E cells located at the apical parts and the round R cells located in the center (A). Ten days after germination, SFs differentiate, mainly in the center and on R cells (Le Bail et al., 2008a) (B), thereby generating a loosely filamentous organism after 20 d (C). In *etl* sporophytes, cell types are less distinct (D), and the arrest of apical division and the more prolific branching (E) leads to a bushy, compact organism (F).

(G) Changes in cell number in the whole organism over time. For each observed individual, the time series was adjusted so that the first division occurred at $t = 0$. The number of observed wild-type (WT) and *etl* filaments was 54 and 35, respectively. Mean values \pm sD.

(H) Position of the first nonaxial division, giving rise to the first SF (SF I). The *y* axis represents the relative deviation from the values expected under the assumption of a uniform distribution of SF.

(I) and (J) Developmental stage (number of cells in the primary filament) when the first (top) and second (bottom) nonaxial division occurs, giving rise to the first and second SFs (SF I [I] and SF II [J]), respectively. Means (represented as a vertical bar) and SD are indicated under the *x* axis. *n* = 50 individuals for both (I) and (J).

Figure 4. Modification in Cell Size, Proportion, and Ultrastructure in *etl*.

(A) Cell size. In *etl*, E cells are shorter (31 μ m versus 37 μ m in the wild type [WT]) and wider (11 μ m versus 8 μ m in the wild type; *t* test, P < 0.01) and R cells are larger (17 μ m diameter) than in the wild type (14 μ m; P < 0.01). Means and SD are indicated (error bars) (*n* > 100).

(B) Proportion of R cells at each stage. Means and SD are indicated (error bars). *n* > 100 for each stage in both the wild type and *etl*.

(C) and (D) General morphology of the wild-type (C) and *etl* (D) cells observed by TEM.

(E) and (F) Details of the ECM in the wild-type (E) and *etl* (F) cells showing a similar organization with two layers: a thin outer black layer and a thick inner gray layer displaying an alignment of cellulose fibers.

(G) Thickness of the ECM. Several measurements were performed on >80 wild-type and *etl* cells observed by TEM. The mean \pm SD of the smallest thickness per cell are presented (*t* test, P < 0.01).

and the simulated populations at the 10-cell stage (correlation coefficient = 0.89). When these eight transition rules were used to similarly simulate the development of the wild-type population (correlation coefficient $= 0.85$), the optimal probabilities of the three additional rules S, P, and V remained close to 0 (Figure 6B). This shows that the gain in adequacy for *etl* modeling brought by these new rules did not result from the increase in the number of free parameters. Instead, it supports the conclusion that these cellular transitions are specific to *etl* and do not participate in the development patterning of wild-type filaments, in agreement with our in vivo observations.

These results allowed us to speculate on the cellular mechanisms altered in *etl*. In addition to generating more R cells, the additional rules implemented a modification in the type of immediate neighboring cells required to promote a given transition in a cell (Figure 6A). Rule P imposed a different and weaker constraint on the nature of the neighboring cells than rule D, as only an empty space bordering the differentiating cell was required. Rule S, whose probability of occurrence was among the highest (Figure 6B), was unique as it did not impose any type of constraint on the neighboring cells. Altogether, computational simulations demonstrated that *etl* could be modeled using both additional transitions occurring during the initial developmental patterns and modified local constraints, which were weaker (rule P and V), different (rule P), or nonexistent (rule S). This result indicates that *etl* may be impaired in the recognition of neighboring cells, a process which is likely to participate in the establishment of local positional information required for proper filament patterning in the wild type.

etl Underexpresses Lin-Notch Domain Transmembrane Proteins

To identify the target genes of this mutation, we analyzed the transcriptome of *etl* using a large-scale microarray approach (Dittami et al., 2009). In addition to the wild-type organism, we included as a technical reference another *E. siliculosus* morphological mutant, called mutant A. After normalization of the data (see Supplemental Data Set 1 online), a statistical analysis was performed using significance analysis of microarrays (false discovery rate = 0.05). In *etl*, 24 ESTs displayed a modified level. This low number was not due to technical limitation of sensitivity or too stringent analysis conditions, as 943 ESTs were shown to be up- or downregulated in mutant A compared with the wild type (see Supplemental Data Set 2 online). Confirmation by quantitative RT-PCR (qRT-qPCR) allowed the identification of only two upregulated ESTs and five downregulated ESTs. The upregulated ESTs belong to a genomic region corresponding to a putative transposable element (Cock et al., 2010) related to a

⁽H) and (I) Cell surface viewed with a scanning electron microscope in wild-type (H) and *etl* (I) cells.

⁽J) and (K) Details of wild-type (J) and *etl* (K) cell ultrastructure by TEM. Nu, nucleus; Chl, chloroplast. The arrows indicate Golgi bodies. Note that this illustrates the most extreme and opposite phenotypes observed in the two organisms.

Different filament organizations, according to the position of R cells (circle) and E cells (rectangle), were observed at each cell stage, from the 2-cell (left) to the 10-cell (right) stage. Some combinations were observed in both wild-type (WT) and *etl* populations (purple), while others occurred exclusively in wild-type (blue) or *etl* (red) populations. The respective frequency (%) of each combination in either of the two populations is indicated on the left side of the corresponding drawing with the same color code. *n* > 100 individuals at each stage.

retrotransposon-like family member from zebra fish (*Danio rerio*; XP001339559.3 BLAST E value = 10^{-34}). This upregulation was not correlated to an increase in the transposition activity of this element, as shown by the qPCR measurement of its genome copy number (see Supplemental Figure 1 online). The five downregulated genes, named *ERG #1-5* (for etl-regulated gene), whose sequences were not modified in *etl* compared with those in the wild-type organisms, contained specific protein domains. ERG5 contained a short-chain dehydrogenase domain. The four other proteins featured a signal peptide and one or several predicted transmembrane domains (Figure 7). In addition, Lin-Notch domains (Interpro IPR000800, also named Notch domain or LNR) were present in up to three copies in three of the ERG sequences (ERG2, ERG3, and ERG4). In *E. siliculosus*, at least 14 predicted proteins with Lin-Notch domains (E value $< 10^{-5}$) are present (see Supplemental Data Set 3 online), while at the same significance level, none have been found in the genomes of the monocellular diatoms *Thalassiosira pseudonana* and *Phaeodactylum tricornutum*, which constitute the phylogenetically most closely related organisms with available genome sequences.

The low number of ESTs showing a modified expression level in *etl* remains surprising, considering the pleiotropic phenotype of this mutant. Therefore, it seems likely that the phenotypic alterations reflect additional molecular phenomena, which cannot be detected by transcriptome analysis. Indeed, in addition to transcriptional regulation of gene expression, various processes are able to account for phenotypic variation. Hundreds of plant genes are regulated by microRNAs (Zhang et al., 2010), which can cleave the mRNA at the recognition site or block mRNA translation (reviewed in Voinnet, 2009), thereby reducing protein production, while maintaining the corresponding transcripts at nearly normal levels (Brodersen et al., 2008). Therefore, we

Figure 6. Computational Modeling of the Early Developmental Pattern in the Wild Type and *etl*.

(A) Transition rules. Filaments are schematized by circles (R cells) and rectangles (E cells). The different events of cell differentiation are indicated by arrows. A letter, corresponding to the transition rule, is given above the arrow. The symbols accompanying the letter designate any required cells adjacent to that undergoing the transition: * for an empty neighborhood, $^{\circ}$ for an R cell, and $^{\Delta}$ for either an R or an E cell. No symbol means that the transition occurs with any type of neighboring cell. Black arrows and letters represent the rules used to model the development of the wild-type filaments; red dashed arrows and red letters are those added to model the development of *etl* filaments.

(B) Optimized probabilities of occurrence used to simulate the developmental pattern of wild-type (WT) and *etl* filaments. Each starting configuration (left) is defined by a cell type for the central cell (in brown) and its immediate neighbors (open gray shapes). The possible states are E (elongated, a rectangle), R (round, a circle), and 0 (empty, a star). Superimposed symbols mean that either state is allowed. The transitions (arrows) affect only the central cell; they are called I for initial, E for elongate, A for apical, P for partition, V for variegate, R for round, D for double, and S for split. All transitions are stochastic (i.e., they occur with a given probability). For each transition rule, the optimized probabilities enabling the best simulation at each developmental stage are shown on the right part, for both the wild type and the *etl* mutant. In each case, the simulation was optimized with either five or eight rules. An indicator of discrepancy between the simulated population and the observed population is given (see Methods for details).

cannot exclude the possibility that additional genes are misexpressed in *etl*, which cannot be detected by the microarray approach.

DISCUSSION

Most of the previous studies on brown algal development were performed on fucoid zygote polarization (*Fucus* sp or *Silvetia* sp). They allowed the identification of molecular actors involved in this initial process of development (for an update, see Hable and Kropf, 2005). Interestingly, Berger et al. (1994) showed that, later in the embryonic development, a component of the cell wall was determining the cell fate. Unfortunately, because of the experimental limitations of fucoid models, this component remains unidentified more than 15 years later. This study used another model, *E. siliculosus*, which is more amenable to the investigation of thallus development. Previous studies on its wild-type sporophyte filament have shown that, at the early stages, the apical E cells both elongate and divide away from the center of the filament, ensuring most of its growth, while the resulting subapical E cells progressively differentiate into R cells (Le Bail et al., 2008a). The balance between these two activities (i.e., cell elongation and division on one hand and cell differentiation on the other hand) might be at least partially regulated by auxin, as this phytohormone is accumulated in the E cells where it inhibits both E-to-R cell differentiation and branching (Le Bail et al., 2010).

In this study, we characterized a morphogenetic mutant of a brown alga. The single locus mutation *etl* is responsible for a hyperbranching activity with an earlier and more synchronized emergence of branches in no preferential position along the filament. The partial loss of the E cell type in *etl*, resulting in a high abundance of R cells (90% compared with 50% in the wild type) can easily account for this phenotype, as in the wild type mainly R cells ensure branching (Le Bail et al., 2008a). This also accounts for the arrest in mitotic activity observed early in the apical parts of the *etl* filaments. Furthermore, *etl* cells displayed modifications of their structural features: in addition to slight modifications in their dimensions, *etl* cells showed a thicker ECM with a rougher surface, which was correlated with a higher activity of the Golgi apparatus. Moreover, several observations supported an alteration of positional information: (1) cell transition events were modified, and modeling showed that this could result from a partial loss in cell–cell communication, and (2) primarily Notch domain transmembrane proteins were underexpressed. Altogether, these results show that *ETL* regulates cell differentiation and developmental patterning in the brown alga *E. siliculosus*

⁽C) Simulation of a virtual population of *etl* filaments at the 10-cell stage. The simulation was conducted with the optimized probabilities given in (B). All patterns observed in vivo (Figure 5) were generated by the simulation. The frequency of each combination, either generated by the simulation (black) or observed in vivo (gray), is shown. Patterns with a simulated frequency of <1% are grouped in others. The linear correlation coefficient between the frequencies (r) and the P value of the Pearson test (p) is indicated.

Figure 7. ERG Proteins Are Targets of the *ETL* Locus.

Microarray experiments were conducted on four independent biological replicates corresponding to either wild-type or *etl* samples. Significance analysis of microarrays (Tusher et al., 2001) identified 19 significantly downregulated genes in *etl*, of which only five were confirmed by qRT-PCR. (A) Map of the five ERG proteins. The genome sequences of the *ERG* genes are available at http://bioinformatics.psb.ugent.be/webtools/bogas/ with the accession numbers indicated in parentheses. Protein domains are indicated with a color code. The number of amino acids of the corresponding protein is indicated at the right-hand side of the gene map.

(B) Expression level. Transcript levels were measured by qRT-PCR and normalized according to Le Bail et al. (2008b). Data are expressed as the log2 of the ratio with the wild-type (WT) organism. $n = 4$ biological replicates; error bars are SD.

and point to a combined role of ECM and novel transmembrane proteins in cell–cell communication involved in these processes.

The understanding of the functional relationship between the ECM and ERG1-4 transmembrane proteins will require detailed analyses focused on these two cellular and molecular actors. Indeed, both *E. siliculosus* ECM and ERGs display peculiar features, which make interpretation of the data challenging. The brown alga ECM is composed mainly of alginate and sulfated fucan (fucoidan) matrix polysaccharides, which embrace a low proportion of rigid cellulose fibers (Kloareg and Quatrano, 1988). Interestingly, Michel et al. (2010) recently showed that the *E. siliculosus* ECM was synthesized by a patchwork of enzymes inherited from various ancestors through independent evolutionary events. While cellulose is synthesized by enzymes closely related to land plant cellulose synthases (CESAs), fucoidans are synthesized by an ancestral pathway shared with animals. In the latter organisms, this pathway is responsible for the biosynthesis of sulfated ECM polysaccharides (e.g., heparan sulfate). Even more surprisingly, a horizontal gene transfer from an Actinobacterium is thought to account for the acquisition of alginates as well as hemicellulose. Altogether, these polysaccharides provide brown algae with a unique ECM adapted to the marine environment (Michel et al., 2010).

The unique nature of the brown algal ECM sharing features with land plants and metazoans makes comparisons with these organisms both risky and tempting. In animals, numerous examples of the combined role of the ECM and transmembrane proteins in cell development have been reported (Friedl and Zallen, 2010). The ECM regulates cell polarization by activating mechanosensitive transmembrane proteins, so-called integrins, and by stimulating the assembly of focal adhesion complexes. Reciprocally, the transmembrane and intracellular Wnt signaling pathway impacts the organization of the ECM. The ECM also regulates cell differentiation. Sulfated polysaccharides present within the ECM play a determining role in cell fate by regulating ligand binding to transmembrane receptors. For example, heparan sulfate, a sulfated ECM glycosaminoglycan, regulates the diffusion of the ligand of the epidermal growth factor (EGF) receptor through the ECM by mediating its sequestration by proteoglycans (Prince et al., 2010). This sequestration engages the cell in growth inhibition instead of cell proliferation. Interestingly, brown algal fucoidans also inhibit the binding of various ligands to animal transmembrane receptors (Heinzelmann et al., 1998). Hence, in brown algae, fucoidans could similarly restrain ligand diffusion to the Notch domain receptors, a process that could be enhanced in *etl* because of its thicker ECM. The reported positive feedback loop exercised by Notch proteins on Notch genes (Liu et al., 2009) illustrates a mechanism of autoregulation that could also apply to *ERG* genes and could possibly account for the decrease in their transcript level observed in *etl*.

In land plants, the cell wall has long been viewed as a static compartment, through which signal molecules must diffuse (reviewed in Brownlee, 2002). However, it was recently recognized as a dynamic and responsive compartment with an important role in relaying information from external stimuli during development (Kohorn, 2000; Humphrey et al., 2007). Several cell wall proteins are thought to be involved in these processes. The family of receptor-like kinases, and more specifically the wallassociated kinases, are transmembrane proteins able to sense the cell wall integrity and to regulate machineries involved in cell wall modification. They are involved in several morphogenetic events (Kohorn, 2001). Wall-associated kinases contain two EGF-like repeats in their extracellular domain, and their interaction with the cell wall polysaccharide pectin as well as Gly-rich proteins (Steinwand and Kieber, 2010) regulates the expression of numerous downstream genes, such as transcription factors and phytohormone-responsive genes (Kohorn et al., 2009). Similarly, structural cell wall components, such as Leu-rich repeat extensin proteins, were shown to regulate proper cell wall organization, and the presence of a Leu-rich repeat domain in these proteins suggests that they act as regulator and/or receptor of extracellular signals (Baumberger et al., 2003).

Altogether, in both the green and the opisthokont lineages, the study of the functional interactions between the ECM and transmembrane receptors is still an emerging field, and the peculiar evolutionary history of the brown algal ECM emphasizes the interest in deciphering them in *E. siliculosus* using *etl* as a tool.

The putative receptors identified in this study illustrate once more the special features of brown algae. Most of the transmembrane proteins repressed in *etl* (ERG1-4) contain a Lin-Notch domain. On one hand, this domain is absent from the Plantae phylum, with the exception of one nontransmembrane protein (Q016E4_OSTTA) in the prasinophyte *Ostreococcus tauri* (Chlorophyta). On the other hand, the Notch domain is specifically found in metazoan transmembrane receptors called Notch receptors, where it regulates the interaction between the extracellular and the intracellular regions (Fiúza and Arias, 2007). These Notch receptors are fundamental for animal development, in which they control lateral inhibition, boundary formation, and cell fate choice upon direct interaction with another transmembrane protein expressed at the surface of the adjacent cells (Delta or Jagged/Serrate; Fortini, 2009). Deregulation of Notch proteins is responsible for numerous cancers, and their involvement as both an oncogene and a tumor suppressor has been demonstrated frequently (Bolós et al., 2007; Koch and Radtke, 2007; Dotto, 2008). *ERG2-4* do not contain EGF-like repeats, which mediate the interaction of metazoan Notch receptors with their extracellular ligand (Artavanis-Tsakonas et al., 1995). The lack of these EGF-like domains in the *E. siliculosus ERGs* precludes solid presumptions about a putative conservation of function. However, the combination of the domains present in Notch receptors seems to result from a shuffling of ancestral domains specific to metazoans (Gazave et al., 2009). Hence, brown algae might have evolved alternative strategies to assemble domains at the cell surface. As a matter of fact, several predicted excreted or transmembrane EGF-like domaincontaining proteins are encoded by the *E. siliculosus* genome (see Supplemental Data Set 4 online), and homologs were shown to be secreted from embryonic cells of the brown alga *Fucus distichus* (Belanger et al., 2003). Therefore, *E. siliculosus* could have evolved structurally different receptors involved in cell–cell signaling while conserving Notch domains. The chemical features of the brown algal ECM could easily support the hypothesis that these organisms have evolved different cell–cell communication strategies than metazoans. This hypothesis is compatible with the absence of homologs of the Notch ligands Delta/ Serrate/Lag-2 in the genome of *E. siliculosus* (see Supplemental Data Set 5 online).

Many questions related to the *etl* mutant remain unanswered. In particular, whether the *E. siliculosus* ECM influences ERG signaling or vice versa is not clear, as several examples of reciprocal ECM-receptor interactions have been reported in animals and plants (Humphrey et al., 2007; Friedl and Zallen, 2010). Nevertheless, because of the unique features of their ECM, brown algae are interesting models to study ECM-transmembrane receptor interactions and their impact on cell differentiation. The presence of integrin genes and associated proteins (e.g., talin and α -actinin) recently reported in the genome of *E. siliculosus* illustrates the unexplored potential of brown algae as models to unravel novel ECM-cytoplasm communication systems (Cock et al., 2010). In this context, *etl* is a particularly relevant tool to start deciphering them. The identification of the potential ligands of the Lin-Notch-ERG transmembrane proteins is among the most challenging goals for future studies. The characterization of their diffusion through the ECM and their interaction with Notch domain proteins will enable a better understanding of the mechanisms accounting for the developmental pattern of *E. siliculosus* filaments. The other major challenge is the identification of the *ETL* locus, as we have shown it to be a major regulator of the developmental pattern in *E. siliculosus*, as a probable actor in the ECM receptor signaling processes. Positional cloning of *ETL* has recently been initiated, and the availability of other branching mutants in the laboratory might help to identify *ETL*.

In conclusion, this study complements the recent discovery on the role of auxin in the development of *E. siliculosus* (Le Bail et al., 2010) and illustrates that the development of brown algae is likely based on a combination of strategies with molecular actors conserved in either metazoans or land plants, although in each case, variations were found at the molecular level.

METHODS

Algal Strains, Culture Conditions, and Production of Mutants

The wild-type strain used was Ec32 (accession CCAP 1310/4; origin San Juan de Marcona, Peru). The *etl* mutant (CCAP 1310/337) was generated from unfertilized gametes of the Ec32 strain irradiated for 11 min (LD50) with UV-B (316 nm) and selected after 2 weeks of growth for its modified morphology compared with wild-type parthenogenetic sporophytes (parthenosporophytes). The Ec568 female strain (accession CCAP 1310/334; origin Arica, Chile) was used to perform crosses with the *etl* mutant. Thalli were grown in half-strength Provasoli-enriched (Starr and Zeikus, 1993), autoclaved natural seawater (pH 7.8) in Petri dishes located in a controlled environment cabinet at 13°C with a 14:10 light: dark cycle (light intensity 29 μ mol photon·m⁻²·s⁻¹).

Zygote Production

etl (Ec32 genetic background) was crossed with a wild-type sister strain (Ec568). Fertile gametophytes of each sex were placed in the dark in a small amount of medium for a few hours to induce synchronized gamete release after the addition of natural seawater. Five microliters of each gamete solution was then mixed in a single drop on a cover slip, which was inverted on a silicone joint fixed on a slide with Vaseline to isolate the drop. Zygotes, which can be distinguished from unfertilized gametes due to the presence of two stigmata, were localized in the drop and monitored for 2 weeks in the hanging drop with change of medium every 2 d. Thereafter, the cover slip with germlings was immersed in a Petri dish and further development was followed under an inverted microscope. Germlings from the germination of unfertilized gametes were removed. The sex of gametophytes was determined by microscopy observation of zygote formation in hanging-drop preparations in which the strains to be tested had been combined with fertile thalli of male and female reference strains. The heterozygous state of each zygote was confirmed by PCR amplification with markers specific to the two parents (see below). The 10 zygotes obtained from this cross all showed a wild-type phenotype, indicating that the *etl* mutation is recessive.

Cross-Verification

DNA was extracted from the diploid heterozygous sporophytes with the NucleoSpin Plant II kit (Macherey-Nagel). Nuclear DNA was amplified in a microsatellite region whose length differs by 40 bp between Ec32 (genetic background of the *etl* mutant) and Ec568 (Heesch et al., 2010; primer sequences are ESM27F, 5'-TGTAAAACGGCCAGTGCAACTGCCCTCG-GAAAATAAG-3', and ESM27R, 5'-GCGCGGTCGATGCTAATAATAA-CTA-3'). The amplification was performed in an Applied Biosystems thermocycler (35 cycles, 0.5 min 94 $^{\circ}$ C, 0.5 min 65 $^{\circ}$ C, 1 min 72 $^{\circ}$ C) using GoTaq DNA polymerase (Promega).

Morphometric Analysis

Observations were performed using an Olympus IX 51 inverted microscope. Cell sizes were measured using ImageJ software (http://rsbweb. nih.gov/ij/).

Cells in a filament were numbered from the closest extremity. A cell within a filament of *L* cells in length was assigned the number *c*, where, for reasons of symmetry, *c* is an integer such that $0 \le c \le (L-1)/2$. To compare positions between primary filaments of unequal length, the cell position was converted into a relative index using a procedure adapted from Le Bail et al. (2008a): a number of intervals, *N*, was chosen (in this study, we used $n = 10$). The whole filament was graduated into 2*N* intervals so that each extremity was marked as 0, and the center was marked *N* (note that *N* represents the limit between two cells if *L* is even, and the middle of the central cell if *L* is odd). If an event (such as an axial division or the emergence of a secondary filament) occurs at cell number *c*, then this event is shared between the intervals that overlap with this cell, each interval receiving an amount proportional to the extent of overlap. Mathematically, cell *c* overlaps with the *n*th interval (the interval between graduations $n-1$ and n) if and only if $2c/L \le n/N$ and $(n-1)/n \le n$ $(2c+1)/L$; then the *n*th interval undergoes an amount of event $A_n = min(n/N,$ $(2c+1)/L$) - max((n-1)/N,2c/L). For each interval, the amounts computed in all individuals of the observed sample are added, giving a total amount of observed events for each interval, *O*n. In parallel, for each observed individual, *L* expectations were computed by assigning the same probability (1/*L*) of event to each cell and shared between intervals using the same procedure. These expected amounts are also added over the whole population, giving a total amount of expected events for each interval: *E*n. The value $S_n = (O_n - E_n)/(O_n + E_n)$ represents the bias between observed and expected events in the *n*th interval.

TEM

Parthenosporophytes were fixed with 4% glutaraldehyde and 0.25 M sucrose at room temperature and washed with 0.2 M sodium cacodylate buffer containing graded concentrations of sucrose. The samples were postfixed in 1.5% osmium tetroxyde, dehydrated with a gradient of ethanol concentrations, and embedded in Epon. Polymerization was performed at 60°C for 24 h. Ultrathin serial sections were cut with a diamond knife on an ultramicrotome and were mounted on copper grids. After staining with 2% uranyl acetate for 10 min and 2% lead citrate for 3 min, the grids were examined with a Jeol 1400 transmission electron microscope.

Scanning Electron Microscopy

Twenty microliters of a gamete release (Le Bail et al., 2008a) was loaded and fixed on a polycarbonate filtration membrane (diameter 13 mm; Nuclepore, Whatman). After 2 weeks of growth, parthenosporophytes were fixed in artificial seawater (ASW) with 3% paraformaldehyde for 1 h and then washed for 10 min in ASW:water (3:2), ASW:water (2:3), and water. The parthenosporophytes were then dehydrated successively in a gradient of ethanol concentrations (30, 50, 70, and 90%, two times 95%, and three times 100%) and finally dried using a critical point dryer (Baltec CPD 030; Balzer). The membranes were then fixed on a glass plot and covered with a 25-nm-thick gold layer. Filaments were finally observed with a JEOL JSM 5200 scanning electron microscope.

Measurements of the cell wall thickness were performed using the ImageJ software (http://rsb.info.nih.gov/ij/). Because sections are not necessarily perpendicular to the cell wall, calculations of the mean were performed on both five random measurements per cell and a single value corresponding to the smallest thickness ($n = 89$ for wild-type and $n = 82$) for *etl* cells) and were statistically analyzed with the R software (http:// www.r-project.org/; R Development Core Team, 2011). The two measurement protocols yielded similar results.

Modeling

The development of a wild-type primary filament of *Ectocarpus siliculosus* was initially modeled by the stochastic one-dimensional cell automaton ECTOMAT (Billoud et al., 2008). Filaments of *etl* mutants could not be simulated by the same model, and observations showed that in this mutant, new transition events had to be taken into account. As the initial configurations of these new rules were expressed by patterns of various lengths, we reimplemented ECTOMAT in the more general form of a stochastic L-system (for review, see Prusinkiewicz and Lindenmayer, 1996). The probabilities for the five or eight transitions were optimized using a random descent algorithm, in which the value to minimize was the sum of squared differences between the distributions of R cell number in simulated versus observed filaments at each stage, normalized by the total number of cells (=indicator of discrepancy).

Microarray Experiment

Microarray experiments were performed on four independent wild-type, *etl*, and mutant A cultures. Each of the 12 cultures was grown in a single Petri dish, in identical growth conditions (culture cabinet), and for different culture durations (between 20 and 40 d, until they reached similar developmental stages corresponding to immature parthenosporophytes). To avoid the putative effects of diurnal oscillations on gene expression, all replicates were harvested at the same hour of day (10:30 AM). Inoculation and collection dates were different. These samples were frozen in liquid nitrogen and RNA extractions were performed as described by Le Bail et al. (2008b). cDNA synthesis and amplification were performed following the protocol outlined by Dittami et al. (2009). The array design contained probes for a total of 17,119 unigenes, each represented by four 60-mers, and was created by Roche NimbleGen, as described by Dittami et al. (2009). Roche NimbleGen performed cDNA labeling and hybridization (single-channel platform) as part of their Gene Expression Array Service. These experiments were based on amplified double-stranded cDNAs. Differentially expressed genes were identified in the data set with a significance analysis of microarrays test (Tusher et al., 2001) using TigrMEV 4.3 (Saeed et al., 2003; false discovery rate = 0.05). Normalized expression values were generated using quantile normalization (Bolstad et al., 2003) and the Robust Multichip Average algorithm (Irizarry et al., 2003a, 2003b) (see Supplemental Data Set 1 online). The data obtained for the *etl* mutant were uploaded to ArrayExpress under accession number E-TABM-799.

qRT-PCR

Validation of the microarray data was performed by qRT-PCR amplification (SYBR green) of the target genes as described by Le Bail et al.

Sequence Analysis

The peptide sequences of the ERG proteins were initially obtained from the automated annotation of the *E. siliculosus* genome (Cock et al., 2010) and checked in the wild type by sequencing of PCR-amplified or cloned full-length cDNAs for ERG 1, 2, 3, and 5. Using a proofreading PCR-based approach, each of the five ERG coding sequences framed by \sim 1 kb upstream and downstream was obtained in the *etl* mutant. Conserved domains were identified using InterProScan (Zdobnov and Apweiler, 2001). BLASTP analyses were performed using the blast 2.2.21 package (Altschul et al., 1997).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Expression and DNA Copy Number of the Retrotransposon in Wild-Type and *etl* Genomes.

Supplemental Table 1. Oligonucleotides Used in the Quantitative RT-PCR Analysis of the ERG Genes.

Supplemental Data Set 1. Normalized Signals of the Hybridized Microarrays.

Supplemental Data Set 2. Statistical Analysis of the Normalized Microarray (SAM) Data.

Supplemental Data Set 3. *E. siliculosus* Proteins with a Notch Domain.

Supplemental Data Set 4. Presence of EGF Patterns in the Proteome of *E. siliculosus* (Results from Interproscan).

Supplemental Data Set 5. Best Matches for the Uniprot Entries Annotated Delta/Jagged/Serrate Proteins in the Proteome of *E. siliculosus*.

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