LagC is required for cell–cell interactions that are essential for cell-type differentiation in Dictyostelium

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Strain AK127 is a developmental mutant of Dictyostelium discoideum that was isolated by restriction enzyme-mediated integration (REMI). Mutant cells aggregate normally but are unable to proceed past the loose aggregate stage. The cloned gene, lagC (loose aggregate C), encodes a novel protein of 98 kD that contains an amino-terminal signal sequence and a putative carboxy-terminal transmembrane domain. The mutant strain AK127 shows no detectable lagC transcript upon Northern analysis, indicating that the observed phenotype is that of a null allele. Expression of the lagC cDNA in AK127 cells complements the arrest at the loose aggregate stage, indicating that the mutant phenotype results from disruption of the lagC gene. In wild-type cells, lagC mRNA is induced at the loose aggregate stage and is expressed through the remainder of development. lagC− null cells aggregate but then disaggregate and reaggregate to form small granular mounds. Mature spores are produced at an extremely low efficiency (<0.1% of wild type), appearing only after ~72 hr, whereas wild-type strains produce mature spores by 26 hr. lagC− null cells accumulate reduced levels of transcripts for the prestalk-enriched genes rasD and CP2 and do not express the DIF-induced prestalk-specific gene ecmA or the cAMP-induced prespore-specific gene SP60 to significant levels. In chimeric organisms resulting from the coaggregation of lagC− null and wild-type cells, cell-type-specific gene expression is rescued in the lagC− null cells; however, lagC− prespore cells are localized to the posterior of the prespore region and do not form mature spores, suggesting that LagC protein has both no cell-autonomous and cell-autonomous functions. Overexpression of lagC from an actin promoter in both wild-type and lagC− cells causes a delay at the tight aggregate stage, the first stage requiring LagC activity. These results suggest that the LagC protein functions as a nondiffusible cell–cell signaling molecule that is required for multicellular development.

[Key Words: Dictyostelium discoideum; cell–cell interactions; signal transduction, developmental mutation]

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Removal of nutrients from Dictyostelium discoideum initiates a developmental process that culminates with the formation of a fruiting body consisting of spores held up by stalk cells. The first step in the formation of this multicellular structure is the aggregation of individual amoebae through chemotaxis to extracellular cAMP [Konijn et al. 1968]. Within the aggregate, cells fated to become stalk or spores can be distinguished by the expression of cell-type-specific genes (Mehdy et al. 1983; Krefft et al. 1984; Morrissey et al. 1984; Gomer et al. 1986; Williams et al. 1989; Esch and Firtel 1991; Haberstroh and Firtel 1990; Fosnaugh and Loomis 1993]. These initially intermixed prestalk and prespore cells then sort out to establish spatial patterns that persist throughout culmination. In response to extracellular cAMP, prestalk cells migrate through the aggregate to form the tip, which acts as an "organizer" during the rest of Dictyostelium development [Raper 1940; Traynor 92]. The tip of the aggregate elongates to form the first finger, which then falls onto the substratum to produce the migrating slug that eventually culminates, forming a mature fruiting body. The entire developmental process takes ~24 hr [Loomis 1982].

Extracellular cAMP plays a key role during subsequent development as a regulator of prestalk- and prespore-specific genes. Induction of these genes requires high levels of cAMP produced within the aggregate, which then binds to cell-surface receptors and activates G protein-coupled signaling pathways [for reviews, see Firtel 1991; Kimmel and Firtel 1991; Schaap 1991; Williams 1991]. Prestalk-specific and prestalk-enriched genes, which are induced early in this process [e.g. rasD and pst–cathepsin/CP2 (cprB)], can be induced to a low level in starving
cells in shaking culture by the addition of nanomolar pulses of cAMP that mimic signaling during aggregation [Mehdy et al. 1983; Reymond et al. 1984; Mehdy and Firtel 1985; Pears et al. 1985; Datta et al. 1986; Jermy et al. 1987]. These genes are then fully induced in response to high levels of cAMP. Prespore genes such as SP60 (cotC), SP70 (cotB), and pspB require cell–cell contact, in addition to high levels of cAMP, for expression [Mehdy et al. 83; Chisholm et al. 84; Mehdy and Firtel 85; Fosnaugh and Loomis 1991; Powell-Coffman and Firtel 1994]. The prestalk gene ecmA requires DIF (differentiation inducing factor) in addition to cAMP for maximal expression [Williams et al. 1987; Berks and Kay 1990]. lacZ reporter constructs driven by these cell-type-specific promoters have been used to study spatial patterning and temporal differentiation of cell types in wild-type and mutant strains, and to follow cellular differentiation in chimeras containing wild-type and mutant cells [Haberstroh and Firtel 1990; Esch and Firtel 1991; Mann and Firtel 1991, 1993; Hadwiger and Firtel 1992; Hooper 93; Fosnaugh and Loomis 1993].

Recently, a technique for insertional mutagenesis in Dictyostelium, termed restriction enzyme-mediated integration (REMI) was developed [Kuspa and Loomis 1992]. The technique involves introduction of an active restriction enzyme along with linearized vector DNA containing compatible cohesive ends into Dictyostelium cells. High-efficiency integration occurs at chromosomal sites recognized by the restriction enzyme. REMI is mutagenic in Dictyostelium, and developmental mutants have been isolated at a frequency of 0.3–1% of the transformants. The vector-tagged mutated genes in these strains can then be isolated by excising the vector and flanking DNA with a second restriction enzyme, followed by circularizing the DNA with ligase and transforming into Escherichia coli.

Strain AK127 was identified in the initial REMI screen for developmental mutants and shown to contain a single vector insert [Kuspa and Loomis 1992]. Morphogenesis in this strain is blocked at the loose aggregate stage. Because of the observed phenotype, the affected gene was designated lagC (loose aggregate mutant C). The mutant phenotype was regenerated by transforming linearized vector DNA carrying flanking sequences into wild-type Dictyostelium cells. Homologous recombination into the lagC gene indicated that phenotype results from disruption of this gene [Kuspa and Loomis 1992, Dynes et al. this paper]. This paper reports the analysis of strain AK127, and the cloning and molecular characterization of lagC. We show that lagC encodes a developmentally regulated, putative transmembrane domain protein that is essential for the expression of genes induced after aggregate formation in vivo and in shaking culture in response to high levels of cAMP. Analysis of chimeric organisms marked with cell-type-specific reporter genes shows that LagC has both cell-autonomous and non-cell-autonomous functions and indicates that lagC is involved in cell–cell interactions that are essential for cellular morphogenesis. Our results suggest that the lagC mutation identifies a previously unknown component of a signaling pathway that is essential for multicellular differentiation in Dictyostelium.

Results

lagC cDNA and protein sequence

Plasmid p127Cla contains Dictyostelium genomic sequences from around the vector insertion site (IS127) in strain AK127 [Kuspa and Loomis 1992]. These sequences were used to isolate a 3.0-kb lagC cDNA, termed c127.1, from a library that was constructed using RNA isolated at 12–16 hr of development [Fig. 1; Schnitzler et al. 1994; see Materials and methods]. In the cDNA clone, the DNA sequence around the unique BamHI site matched to the BamHI site at the insertion point [Fig. 1]. Analysis of the DNA sequence identified a single long open reading frame that encodes an 888-amino-acid protein with a molecular mass of 98 kD, beginning with the first methionine codon in the cDNA [Fig. 2, top]. The cDNA contains ~200 bp of 5′-untranslated sequence and ~30 bp of 3′-untranslated sequence. The base composition of the untranslated sequences is typical for Dictyostelium noncoding regions (~90% A + T) [Kimmel and Firtel 1983]. Hydrophobicity analysis of the protein sequence and comparison to published data bases predicted three structural features. There is a 17-amino-acid amino-terminal signal sequence and a 24-amino-acid carboxy-terminal glycine-rich hydrophobic putative transmembrane domain with a 14-amino-acid hydrophilic basic carboxy-terminal tail. The putative processed protein would be 871 amino acids, with a molecular mass of 96 kD, as predicted by the signal sequence cleavage algorithm of von Heijne (1986). We expect that the long amino-terminal region containing 17 cysteines would not be cytoplasmic [see Discussion].

Both the nucleotide and amino acid sequences of LagC were compared with other available sequences using the BLAST and BLOCKS programs [Altschul 1990; Henikoff and Henikoff 1991]. The putative transmembrane domain was found to be similar to the putative transmembrane domain of the protein cell–CAM105 (C-CAM),

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**Figure 1.** lagC genomic and cDNA map. The position of insertion of the REMI vector DIV2 into the BamHI site of the lagC gene as well as other restriction sites and start and stop codons is shown. DIV2 inserted ~560 bp from the start codon of lagC in the strain AK127, causing a lagC− null mutation.
The use of suspension cultures of starved cells allows one to test whether the expression of a gene might be regulated under different developmental conditions. Three protocols are used: [1] 30 nM pulses of cAMP, which mimics cAMP signaling during the aggregation phase of development; [2] pulses followed by high continuous levels of cAMP in the absence of continuous cell–cell contact, which induces classes of pretalk-enriched genes expressed during the multicellular stages; [3] pulses followed by high continuous levels of cAMP in the presence of continuous cell–cell contact, which induces classes of pretalk-specific and prespore-specific genes expressed during the multicellular stages (see introductory section and below). Cells remain separate when shaken at high speeds in the presence of EDTA (fast-shake culture) and form stable cell–cell contacts when shaken at low speeds (slow-shake culture) [Fig. 3B; Mehdy and Firtel 1985; Datta et al. 1986]. These three protocols were used to examine the effects of cAMP and cell–cell contact on lagC expression. Wild-type KAX-3 cells expressed lagC at low levels in response to nanomolar pulses of cAMP. The gene is expressed at significantly higher levels in fast-shake conditions in response to cAMP and in slow-shake cultures whether or not cAMP is present. The independence of lagC expression on exogenous cAMP under slow-shake conditions is often observed with genes that can be induced in fast-shake culture in the presence of cAMP. This presumably is attributable to the generation of endogenous cAMP.

which is a mammalian cell adhesion molecule and extracellular ATPase [Auriullius et al. 1990; Obrink 1991]. The identity in this 24-amino-acid region is 54% [Fig. 2B]. The transmembrane domain of the human herpes virus glycoprotein D and the predicted transmembrane domains of the sea urchin egg peptide (speract) receptor and several members of the murine carcinoembryonic antigen family share a similar glycine-rich transmembrane structure [McGeoch et al. 1985; Hinoda et al. 1988; Dangott et al. 1989; Dveksler et al. 1991]. No significant sequence homologies were found.

Regulation of lagC mRNA expression

A portion of the lagC-coding region was used to probe a developmental RNA blot to examine lagC expression kinetics [Fig. 3A]. No lagC transcripts were detected in RNA isolated from vegetative and 4-hr developing cells. A single 3.5-kb band was observed in RNA isolated from wild-type cells from the loose aggregate stage (8 hr) through culmination (24 hr), with the level of transcript peaking at 8–10 hr. When RNA blots from AK127 cells [lagC mutant cells] were examined, no lagC RNA was detected, indicating that the disruption of the lagC gene results in a null phenotype.

Extracellular cAMP acting through cell-surface receptors is known to regulate the expression of many genes expressed during development [see introductory section].
within the clumps (Mehdy and Firtel 1985). In cells shaken in the absence of cAMP, no expression is seen for the first 6 hr. Cells shaken for 9 hr in the absence of cAMP pulses started to clump because of low endogenous cAMP oscillations (Mann and Firtel 1987) and expressed lagC mRNA. lagC transcripts were absent in AK127 cells (lagC- null mutant cells) when assayed in shaking culture with cAMP (data not shown). When a lagC probe was hybridized to mRNA isolated from Percoll gradient-separated prestalk and prespore cells (Ratner and Borth 1983; Kubahara et al. 1993), the lagC transcript was more abundant in prestalk cells than in prespore cells by a ratio of 3:1 (data not shown).

*Developmental morphology of lagC- null cells*

The developmental morphology of lagC- null cells was analyzed by plating washed cells on non-nutrient agar. AK127 cells aggregated to form loose mounds with kinetics similar to those of wild-type KAx-3 cells but did not form hemispherical tight aggregates [Fig. 4A]. The lagC- aggregates started to dissociate at 12 hr and often formed ring-like structures [Fig. 4B]. These loose aggregates appeared to be in flux, with groups of cells splitting off and merging with other groups. By 17 hr, lagC- cells had dispersed to form a rough layer only a few cells thick [Fig. 4C]. The cells then synchronously formed streams and reaggregated by 24 hr [Fig. 4D]. These secondary aggregates did not fall apart again but, instead, became granular by 27 hr [Fig. 4E], developing multiple, small, tip-like structures by ~35 hr [Fig. 4F]. Depending on local conditions on the plate, most of the lagC- cells remained as aggregates at 60 hr [Fig. 4G], but a very small fraction of lagC- cells formed minute fruiting body-like structures [Fig. 4H]. Morphogenesis was also examined in four other, independently derived lagC- derivatives of DH1 generated by homologous recombination using the cloned gene. These strains showed a similar pattern of aggregation, ring formation, disaggregation, and reaggregation; however, the formation of later structures occurred to a significantly lesser degree.

The ability of lagC- null cells to form viable spores, as assayed by the presence of heat- and detergent-resistant cells, was tested (Loomis 1969; Hadwiger and Firtel 1992). When lagC- null cells were developed, no spores (<0.01% of cells developed on agar plates) were detected.

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**Figure 4.** Developmental morphology of AK127 cells. Cells were grown axenically, washed, and plated on non-nutrient agar for development as described in Materials and methods. (A) AK127 aggregates, 9 hr after starvation; (B) disaggregation of AK127 aggregates, 13 hr, showing ring-like intermediate structures; (C) completely disaggregated AK127 cells, 17 hr; (D) AK127 cells reaggregating, 24 hr, showing large streams; (E) AK127 granular aggregates, 27 hr; (F) AK127 aggregates with multiple tips, 40 hr; (G) AK127 final morphological stage, 60 hr [note that E, F, and G all represent the same region of the plate taken at different times.]; (H) AK127 small fruiting body, 60 hr. The magnification of H is approximately five times (linear dimension) greater than the other panels.
at 36 hr, a time when spores are present in wild-type strains. Moreover, when 36-hr structures were dissociated and examined microscopically, no spore-like cells were observed. When lagC− null cells were developed for 72–96 hr, a small number of oval, birefringent spore-like cells were seen in the few minute fruiting body-like structures. The total number of viable spores was very low [0.1–0.01% of cells developed on agar compared with >80% for wild-type cells]. These pseudo-fruiting bodies also contained some vacuolated stalk cells within the "stalk-like" region.

To examine morphogenesis further in AK127 cells and to see whether extracellular cAMP could rescue development, AK127 cells were plated on agar containing 1 mM cAMP and were compared with wild-type cells developed under the same conditions. Wild-type cells formed a ring that resulted from the outward chemotactic movement of cells as the cAMP in the agar was hydrolyzed by phosphodiesterase produced by the cells (Franke and Kessin 1992). The cells remaining in the center aggregated by 9 hr [data not shown] and formed fruiting bodies by 21 hr [data not shown]. Development was repressed in the outer ring of cells by the high cAMP. lagC− null cells showed a similar pattern of development at 9 hr, with aggregates in the center of the plated cells [data not shown]. In contrast to the situation in non-cAMP-containing agar, the lagC− aggregates did not disperse, but formed tight, tipless aggregates at 21 hr, at which time morphogenesis ceased [data not shown]. No evidence for cAMP suppression of the lagC− null phenotype was observed.

Developmental gene expression in lagC− null cells

The effect of LagC on the expression of genes that are differentially regulated during development was examined by RNA blot analysis in wild-type and lagC− null cells developed on filters and in shaking culture in the presence or absence of cell–cell contact and exogenous cAMP. In lagC− null cells developed on filters, transcripts for the prestalk-enriched gene rasD were barely detected, whereas transcripts for the prestalk gene ecmA and the prespore genes SP60 (cotA) and SP70 (cotB) were not detectable within 24 hr of starvation [Fig. 5; data not shown]. The prestalk-enriched gene pst–cathepsin/CP2 (cprB), which is induced at low levels in response to cAMP pulses during aggregation [A. Clark, G. Schnitzler, and R.A. Firtel, unpubl.] and at high levels in response to higher, continuous cAMP in the aggregate [Mehdy et al. 1983; Mehdy and Firtel 1985; Pears and Williams 1987], was expressed at significantly reduced levels in lagC− null cells compared with wild-type KA3-3 cells.

To examine the ability of exogenous cAMP to regulate prestalk and prespore gene expression, cells were starved in shaking culture (suspension), pulsed with cAMP for 6 hr, and then shaken in the presence or absence of cAMP. In lagC− null cells, CP2 was induced to a small extent, similar to the level seen in response to cAMP pulses in wild-type cells [Fig. 3], but further expression was not induced by high, continuous cAMP. Unexpectedly, CP2 mRNA accumulated under fast-shake conditions in the absence of added cAMP in lagC− null cells. rasD produces two transcripts [1.0 and 1.2 kb] from independent promoters [Louvion et al. 1991; Esch et al. 1992]. The rasD 1.2-kb transcript, which is induced in fast-shake culture in wild-type cells in response to high levels of cAMP Esch and Firtel 1991], accumulated in lagC− null cells but to a lower level than was observed in wild-type cells [Fig. 5]. Accumulation of the rasD 1.2-kb transcript was dependent on exogenous cAMP even in slow-shake conditions, unlike the pattern seen in wild-type KA3-3 cells. The 1.0-kb transcript, which is induced at the tight aggregate stage in wild-type cells (10-hr time point), was not observed in lagC− cells. Neither ecmA [a cAMP- and DIF-induced prestalk-specific gene] nor SP60 [prespore-specific] transcripts were detected under any of these assay conditions in lagC− null cells [Fig. 5]. The aggregation-stage, pulse-induced gene csaA (gp80) was normally expressed in response to cAMP pulses in mutant cells.
Mosaic analysis of lagC- null cells showed fewer stained cells than with the wild-type pattern seen in cells, suggesting that hr of development, mixtures of AK127 and wild-type past the loose aggregate stage on their own in the first 24 ing a prior period of cell-cell contact, high levels of ex-

types is not ordered, gates (data not shown).

Because histochemical staining for β-galactosidase ac-

tivity can be more sensitive than RNA blot analysis in that it can detect expression within single cells, the ac-

ivation of cell-type-specific genes was also examined using promoter-lacZ gene fusions. lagC- null cells transformed with rasD/lacZ showed very low levels of staining within the loose aggregate (data not shown). The number of stained cells and the intensity of staining decreased as the aggregates dispersed and then increased as they reformed. AK127 cells carrying an ecmA/lacZ reporter showed fewer stained cells than with the rasD construct, although we observed the same general temporal pattern of β-gal staining as was seen in rasD/lacZ-transformed cells (data not shown). The absence of detectable ecmA transcripts by RNA blot analysis is consistent with the relatively few ecmA/lacZ cells that detectably express β-gal and the longer time period of incubation necessary to see the staining. ecmA/lacZ staining is also seen in the pseudo-fruiting bodies observed at 72–96 hr. In these aggregates, β-gal staining is observed in cells scattered through all parts of the organism, including some in the stalk-like region. The staining in these cells does not show the highly defined pattern seen in wild-type cells, suggesting that these organisms the spatial patterning of the various cell types is not ordered. lagC- null cells transformed with SP60/lacZ showed no detectable expression until after the cells had reaggregated and not until development had proceeded for at least 60 hr. The expression level was very low and staining was seen only in scattered aggregates (data not shown).

Mosaic analysis of lagC- null cells developed with wild-type cells

lagC- null cells carrying various lacZ reporter constructs were washed, mixed with wild-type KAX-3 cells in a ratio of 1:3 (mutant to wild type), and plated for development on filters. Cells were fixed and stained for β-gal expression at various times to determine whether the mutant phenotypes seen in lagC- cells were altered by extracellular signals generated by wild-type cells or direct cell–cell interactions with them. To examine the general spatial pattern of lagC- null cells within the chimeras, lagC- null cells were marked by expressing the reporter construct act15/lacZ (Mann and Firtel 1993), which expresses β-gal in all cell types throughout development. Although lagC- null cells cannot proceed past the loose aggregate stage on their own in the first 24 hr of development, mixtures of AK127 and wild-type cells form chimeras that develop with normal morphology and timing. The lagC- null cells were randomly scattered throughout the newly formed loose aggregates [Fig. 6A]. By the slug stage, mutant cells were present throughout the organism, although there appeared to be a greater concentration of the cells toward the posterior one-third of the slug (Fig. 6B). In control mixtures of labeled and unlabeled wild-type cells, slugs were uniformly labeled (data not shown). In culminants, lagC- null cells were found in the sorus, stalk, upper and lower cups, and basal disc (Fig. 6C,D); however, whereas lagC- null cells appeared to be randomly dispersed in the stalk, they were restricted to the lower half of the spore mass (Fig. 6C). Some lagC- null cells expressing β-gal had either stalk cell-like or spore-like morphology (Fig. 6D). However, when cells from 26-hr culminants were assayed for mature spores (see above), no lagC- null spores were detected (<0.01% of wild type), indicating that this aspect of development cannot be rescued by contact with wild-type cells.

When the proportion of wild-type to lagC- null cells was reduced to 1:1, development became abnormal. The coaggregates formed multiple tips that developed into multiple fingers and slugs and then proceeded through development to form small, but normally proportioned, fruiting bodies. When the proportion of wild-type cells was reduced further to 3:1 (mutant to wild-type), multiple tips were again formed but the aggregates were more granular in structure. Although some of the cells proceeded through development (predominantly wild-type cells), most remained as loose aggregates. When the ratio was reduced further to 9:1, few tips were seen and most cells did not develop further, suggesting that the function provided by wild-type cells is limiting or nondiffusible.

To study the expression pattern of cell-type-specific genes and the spatial patterning of the individual cell types in chimeric organisms, lagC- null cells transformed with lacZ gene fusions driven by the ecmA, rasD, and SP60, spiA, and ecmB promoters were mixed with wild-type cells in the original 1:3 ratio, and the pattern of β-gal expression was examined during development [for promoter references, see Haberstroh and Firtel 1990, Ceccarelli et al. 1991; Esch and Firtel 1991, Jermyn and Williams 1991; Richardson and Loomis 1992, Mann and Firtel 1993, D. Richardson, W. Loomis, and A. Kimmel, in prep.]. ecmA/lacZ is strongly expressed beginning at the loose aggregate stage and is expressed in the prestalk zone (anterior 10–15% of wild-type slugs) and in anterior-like cells [ALCs] (Jermyn and Williams 1991); however, it was expressed at extremely low levels in lagC- aggregates (see above). When lagC-:ecmA/lacZ cells were mixed with wild-type cells, we observed a temporal and spatial pattern of expression of β-gal throughout development that was very similar to that seen in wild-type:wild-type chimeras expressing ecmA/lacZ, except that the boundary between the prestalk and prespore zone was not as sharp as in wild-type slugs (Fig. 6,E–G). Moreover, the rate of staining, an indication of the level of expression, was similar in lagC-:ecmA/lacZ and wild-type:ecmA/lacZ cells when mixed with unmarked wild-type cells (data not shown), suggesting that ecmA expression and the spatial
Figure 6. Cytological staining of act15/lacZ, ecmA/lacZ, and SP60/lacZ constructs in AK127 cells of chimeric organisms containing a ratio of 3:1 untransformed KAx-3 cells to labeled AK127 cell. Cells were mixed before washing, development and staining methods are described in Materials and methods. [A] AK127 cells carrying act15/lacZ mixed with KAx-3 at the aggregate stage, 10 hr of development; [B] act15/lacZ, slug stage, 16 hr; [C] act15/lacZ, fruiting body stage, 24 hr; [D] AK127 spores and stalk cells carrying act15/lacZ, 24 hr; [E] ecmA/lacZ, 10 hr; [F] ecmA/lacZ, 16 hr; [G] ecmA/lacZ, 24 hr; [H] SP60/lacZ, 10 hr; [I] SP60/lacZ, 16 hr; [J] SP60/lacZ, 24 hr.

patterning of prestalk A and prestalk O cells (found in the anterior and posterior halves of the prestalk zone, respectively) were normal in lagC⁻ null cells in the chimeras. In wild-type cells, rasD is detectably expressed in prestalk A and prestalk O cells, ALCs, and rear-guard cells [cells in the very posterior of the slug] [Esch and Firtel 1991]. rasD/lacZ in lagC⁻ null: wild-type chimeras was expressed at the expected stage of development but showed abnormal spatial regulation, with rasD/lacZ-expressing cells being less enriched in regions occupied by prestalk A cells [anterior of the prestalk zone] [data not shown].

When SP60/lacZ-transformed lagC⁻ null cells were mixed with wild-type cells, β-gal-expressing cells appeared at the correct developmental stage but with an altered staining pattern [Fig. 6H–J]. In contrast to the pattern in wild-type slugs in which the posterior 80% of the slug expresses β-gal uniformly [Haberstroh and Firtel 1990], stained lagC⁻:SP60/lacZ cells were present only in the posterior 50% of the slug [Fig. 6I] and were restricted to the lower half of the spore mass [Fig. 6J]. This pattern was similar to that described above with lagC⁻ null: act15/lacZ cells, indicating a general exclusion of lagC⁻ null cells from the top of the sorus [see above, Fig. 6C]. spiA/lacZ, a marker for terminal spore cell differentiation, showed a similar pattern of expression as SP60/lacZ in the sorus of chimeras [data not shown]. ecmAΔ89/lacZ, a marker for stalk cell differentiation, showed that lagC⁻ null cells participated normally in stalk cell formation [data not shown]. These results indicate that wild-type cells can fully rescue prestalk and stalk cell differentiation but can only partially rescue prespore differentiation.

Expression of lagC from an actin promoter in AK127 and KAx-3

The lagC⁻ cDNA was cloned behind the act15 promoter on an integrating plasmid and transformed into AK127 and KAx-3. The resulting transformants were examined for developmental phenotypes by plating on non-nutrient agar. AK127:act15/lagC, KAx-3:act15/lagC, and KAx-3-untransformed control cells made tight aggregates with similar morphology and timing. Both AK127:act15/lagC and KAx-3:act15/lagC cells remained at the tight aggregate stage for 2–6 hr longer than KAx-3 cells before continuing development and forming normal fruiting bodies complete with spores after 30 hr [data not shown]. The results indicate that act15-driven lagC expression in AK127 rescued the developmental arrest phenotype, consistent with the phenotype of AK127 being attributable to a loss of LagC protein. Developmental delay at the tight aggregate stage was seen in both mutant and wild-type backgrounds, indicating that
overexpression of LagC protein was probably responsible for this delay.

Discussion

We show that the cDNA encoding the LagC protein can complement the developmental defect in AK127, indicating that the mutant phenotype results from the loss of LagC protein. The derived amino acid sequence predicts that the LagC protein has an amino-terminal signal sequence for secretion and a putative transmembrane domain that shows similarity to the membrane spanning region of several mammalian and viral cell surface proteins, including C-CAM, suggesting that it is a transmembrane protein. The large number of cysteine residues in the long amino-terminal domain of LagC suggests that this is either extracellular or lumenal and thus likely cytoplasmic.

The developmental morphology of lagC– null cells suggests that the initial time of action of the LagC protein is during the loose aggregate stage. When lagC– null cells are plated at high densities on non-nutrient agar, the cells aggregated, disaggregated, and reaggregated synchronously, as if the mutant cells failed to receive the proper signal within the loose aggregate to continue morphological development, so they then disaggregated and adopted an alternative developmental program. We expect that this disaggregation is a result of a normal, developmental down-regulation of the signaling pathways involved in aggregation combined with an inability of the aggregate to proceed further through development (Kumagai et al. 1988; Devreotes 1989; Firtel 1991). A similar phenotype is also seen with strains carrying a gene disruption of the transcription G-box binding factor (GBF) that arrests at the loose mound stage (see below; Schnitzler et al. 1994). After reaggregation, lagC– null cells form granular aggregates and some late gene expression is observed. lagC mRNA first starts to accumulate in wild-type cells at 8 hr of development, just as the loose aggregate is forming. Because normal morphogenesis arrests at that stage in lagC– null cells, LagC protein appears to be required as soon as it is made. The absence of any effect of overexpressing LagC during growth and early development and a developmental delay in LagC– overexpressor cell lines at the tight aggregate stage argues for the specificity of the action of this gene at this stage.

Although there is an extremely low level of ecmA expression and no expression of SP60 (through 24 hr of development) when lagC– null cells are plated for development, these cell-type-specific genes are induced in chimeras with wild-type cells. This suggests that LagC function is, to a large extent, non-cell-autonomous. This nonautonomous function can be partially complemented only when the majority of the cells within chimeras are wild type, suggesting that LagC function is probably not catalytic or involved in regulating a diffusible molecule. We suggest that LagC acts as an essential component of a cell–cell signaling pathway, possibly a cell-surface ligand, that is necessary for later development. An alternate possibility is that LagC is an essential cell adhesion molecule. Although this latter possibility is not completely excluded, we observed no clumping of cells constitutively expressing lagC during the growth stage. It is possible that LagC functions as one component of a heterophilic cell–cell adhesion system. lagC– null cells that express prespore-specific genes are absent from the front of the prespore zone and the top half of the sorus in chimeras and fail to make normal spores that are resistant to heat and detergents, indicating a second, cell-autonomous role for LagC.

What role does LagC play in development?

Our results suggest that LagC protein is not required for cAMP signaling in early development. Null mutations in genes involved in pulsatile cAMP signaling such as cAR1, Go2, ACA, and PDE all lead to an aggregation-minus phenotype (Kumagai et al. 1989; Sun and Devreotes 1991; Franke and Kessin 1992; Pitt et al. 1992). Mutations that affect cAMP-mediated processes during the aggregate stage, such as PDE overexpression or car2 null mutations, arrest development at the tight aggregate stage, after the induction of cell-type-specific gene expression (Traynor et al. 1992; Saxe et al. 1993). The LagC protein acts between these two phases of cAMP-directed development. The lagC– null phenotype also does not appear to exclusively mediate DIF signaling, as both prestalk and prespore genes are affected in lagC– null cells.

The further development of some lagC– null strains to produce very small aggregates with some abnormal small fruiting body-like structures suggests that the lagC– null phenotype is slightly leaky with regard to development past the loose aggregate stage. When AK127 cells were grown for extended periods of time and the cells were then plated for clones, we identified strains that were capable of making abnormal-looking fruiting bodies containing heat- and detergent-resistant spores. Because these cells are still uracil prototrophs, the partial suppression is not attributable to a loss of the Pyr5-6 marker used to disrupt the lagC– gene. Presumably, the suppression is attributable to a second site mutation and suggests that LagC function can be bypassed in certain genetic backgrounds.

Studies with the G protein Ga4 reveal similarities to the processes affected in lagC– null cells (Hadwiger and Firtel 1992). Although Ga4 is expressed at low levels in vegetative growth, Ga4 transcripts accumulate to high levels at the tight aggregate stage. Overexpression of Ga4 [Ga4Hc cells] causes loose aggregates to disperse, forming rings before reaggregating to form mutant culminants. Likewise, mutants carrying a disruption of the gene encoding the transcription factor GBF aggregate, disperse, and reaggregate and do not express cAMP-induced cell-type-specific genes (Schnitzler et al. 1994). The gbf null mutation is cell autonomous; in mosaics with wild-type cells gbf null cells will coaggregate but will not participate further in development. The similarity of morpho-

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logical and gene expression phenotypes suggests that GBF, Ge4, and LagC could be linked in a common regulatory pathway.

Our data are consistent with a model in which the LagC protein either functions as a signal or is involved in establishing conditions essential for a signal critical to the transition from early to multicellular development in Dictyostelium. The synchronous disaggregation of lagC− mutant cells indicates that a critical signal is missing, with cells responding to its absence in a programmed way. This critical signal can be supplied, in large part, by wild-type cells in mixed populations. Moreover, the structure of the LagC protein is consistent with LagC acting as a cell-surface signal. In addition, it appears that a second cell-autonomous function of LagC is crucial for the transition of prespore cells to mature spores. Further elucidation of the function of LagC will require localizing LagC within the cell and the identification of interacting proteins and/or molecules that lie downstream from LagC in a possible signaling pathway.

Materials and methods

Strains and media

Cells were grown on plates or in shaking culture in HL5 medium [Sussman 1987] supplemented with vitamin B12 (Sigma) to 0.5 μM and folic acid (Sigma) to 0.5 μM, with or without 15–30 μg/ml of G418 [GIBCO/BRL]. Cells were also grown on plates in minimal media as described previously [Franke and Kessin 1977]. Clonal isolates were obtained by plating cells in association with Klebsiella aerogenes on SM nutrient agar plates [Sussman 1987]. Cells grown in HL5 or minimal medium were transformed as described previously [Dynes and Firtel 1989; Kuspa and Loomis 1992]. The following constructs were transformed into Dictyostelium cells: act15/lagC [described below], act15/ lacZ [Mann and Firtel 1993; G. Shaulsky and W. Loomis, unpubl.], rasD/lacZ, [Esch and Firtel 1991], S60/lacZ [Haberstroh and Firtel 1990], emcA/lacZ [Jermyn and Williams 1991], ecmA/ lacZ [Cecarelli et al. 1991], spiA/lacZ [D. Richardson, W. Loomis, and A. Kimmel, in prep.], and p127Cla [Kuspa and Loomis 1992].

DNA constructs and RNA blots

The cDNA c127.1 containing the LagC-coding region, 3′-untranslated region, and a portion of the 5′-untranslated region was isolated from a ZAP [Stratagene] cDNA library made from RNA isolated from cells developed for 12–16 hr [Schnitzler et al. 1994], using a flanking probe from p127Cla [Kuspa and Loomis 1992] and standard protocols [Sambrook et al. 1989]. c127.1 was cloned into the act 15 expression vector EXP4 [+] with Spel–XhoI to create act15/lagC. EXP4 [+] was constructed as follows. The act 15 promoter–2H3 terminator fragment from BS18 [Kumagai et al. 1989, 1991; Firtel and Chapman 1990] was cloned into pATSF [Dynes and Firtel 1989; Haberstroh and Firtel 1990] that had been digested with XhoI–EcoRI, generating EXP1. The Act6-neom selectable marker from BS18 was digested with BamHI, blunt-ended with Klenow, digested with EcoRI, and cloned into EXP1 digested with EcoRI and EcoRV. A NotI linker was put into the blunt end site of the pACT construct, creating EXP2. The EcoRI site in EXP2 was filled in and a NotI linker was inserted to flank Act6-neo, thus generating EXP3. A polynucleotide containing the following sites was inserted into the BglII site in EXP3 to generate EXP4 [+]. [Oligonucleotide restriction sites: BglII, Spel, EcoRI, BclI, XhoI, oligonucleotide sequence, top strand: 5′ GATCTACTAGTGAATTCTGATCACTCGAG 3′; DNA manipulation was performed using standard protocols [Sambrook et al. 1989]. RNA blots were performed as described by Mann and Firtel [1987]. Analysis of gene expression in shaking cultures with cAMP was carried out as described in Mehdy and Firtel [1985]. To produce the figures for publication, standard autoradiographs were scanned in an Apple Colorscanner at 300 dots per inch using Ofoto and saved as a PICT file without any manipulation of the image. The file was opened in Canvas, cropped and labeled, and saved as a PICT file. No other image manipulation was performed. The Canvas file was then printed on a Linotron 500 printer at 1270 lines per inch.

Analysis of morphology, β-galactosidase activity, spore production, and cell-type separation

Analyses of morphology, β-galactosidase activity, and spore production were performed on cells grown axenically in shaking cultures of HL5 medium supplemented with B12/folate (see above) to mid-log phase (1×10^6 to 3×10^6 cells/ml). To examine synchronous morphological development, cells were washed in 12 mM Na/K phosphate buffer (pH 6.1), resuspended at 3×10^7 cells/ml, and spread on non-nutrient agar plates.

For β-galactosidase staining, cells were spread on nitrocellulose filters that were laid on agar plates for development. Histochromalocalization of β-galactosidase activity was determined using the X-gal staining procedure described in Haberstroh and Firtel [1990] and Esch and Firtel [1991]. To assay mature spore production, cells developed for 36 hr or 5 days were washed off non-nutrient agar plates with a rubber scraper, spun down, resuspended in 10 mM EDTA (pH 7.2), 0.1% NP-40, and incubated for 45 min at 42°C. Cells were washed three times with phosphate buffer, resuspended in SM medium, diluted, and plated with K. aerogenes on SM agar plates. Plaques arose from single cells and were counted to quantitate the number of viable spore that were plated [Hadjwiger and Firtel 1992].

Prestalk and prespore cell populations were purified on Percoll gradients as described by Kubahara et al. [1993].

Nucleotide and amino acid sequence analysis

Nucleotide and amino acid sequences were analyzed using the BLAST and BLOCKS programs with available data bases [GenBank, PIR, Swissprotein] [Altschul et al. 1999; Henikoff and Henikoff 1991]. Hydropathy analysis was performed on a Macintosh computer using the Kyte–Doolittle method [Kyte and Doolittle 1982] with the program MacVector 3.5 [International Biotechnologies, Inc./Kodak]. Amino-terminal signal sequence cleavage site was determined by the matrix-weighted method of von Heijne [1986].

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References

Altschul, S.F., W. Gish, W. Miller, E.W. Myers, and D.J. Lipman.
1990. Basic local alignment search tool. J. Mol. Biol. 215: 403–410.

Aurivillius, M., O.C. Hansen, M.B.S. Lazzek, E. Bock, and B.
Öhrink. 1990. The cell adhesion molecule cell-CAM 105 is
an ecto-ATPase and a member of the immunoglobulin super-
family. FEMS Lett. 264: 267–269.

Berks, M. and R.R. Kay. 1990. Combinatorial control of cell
differentiation by cAMP and DIF-1 during development of
Dictyostelium discoideum. Nature 310: 977–984.

Chisholm, R.L., E. Barklis, and H.F. Lodish. 1984. Mechanism of
sequential induction of cell-type specific mRNAs in Dicty-
ostelium differentiation. Nature 310: 67–69.

Dangott, L.J., J.E. Jordan, R.A. Bell, and D.L. Garbers. 1989.
Cloning of the mRNA for the protein that crosslinks to the
egg peptide speract. Proc. Natl. Acad. Sci. 86: 2128–2132.

Devreotes, P. 1989. Dictyostelium discoideum: A model system
for cell-cell interactions in development. Science 245: 1054–1058.

Devksler, G., M. Pensiero, C. Cardellichio, R. Williams, G.
Jiang, K. Holmes, and C. Dieffenbach. 1991. Cloning of the
mouse hepatitis virus (MHV) receptor: Expression in human
and hamster cell lines confers susceptibility to MHV. J. Vi-
rol. 65: 6891–6891.

Dynes, J.L. and R.A. Firtel. 1989. Molecular complementation of
a genetic marker in Dictyostelium using a genomic DNA
library. Proc. Natl. Acad. Sci. 86: 7966–7970.

Esch, R.K. and R.A. Firtel. 1991. cAMP and cell sorting control
the spatial expression of a developmentally essential cell-
type-specific ras gene in Dictyostelium. Genes & Dev. 5: 9–21.

Esch, R.K., P.K. Howard, and R.A. Firtel. 1992. Regulation of the
Dictyostelium cAMP-induced, prestalk-specific DdrasD
gene: Identification of cis-acting elements. Nucleic Acids
Res. 20: 1325–1332.

Firtel, R.A. 1991. Signal transduction pathways controlling
multicellular development in Dictyostelium. Trends Genet.
7: 381–388.

Firtel, R.A. and A.L. Chapman. 1990. A role for cAMP-depend-
ent protein kinase A in early Dictyostelium development.
Genes & Dev. 4: 18–28.

Franke, J. and R. Kessin. 1977. A defined minimal medium for
axenic strains of Dictyostelium discoideum. Proc. Natl.
Acad. Sci. 74: 2157–2161.

---. 1992. The cyclic nucleotide phosphodiesterases of Dicty-
ostelium discoideum: Molecular genetics and biochemis-
try. Cell Signal 4: 471–478.

Gomer, R.H., S. Datta, and R.A. Firtel. 1988. A molecular analysis of G proteins and
control of early gene expression by the cell surface cAMP
receptor on stalk cell differentiation. Proc. Natl. Acad. Sci.
74: 2157–2161.

Havélock, L. and R.A. Firtel. 1990. A spatial gradient of ex-
pression of a cAMP-regulated prestalk cell-type-specific
gene in Dictyostelium. Genes & Dev. 4: 596–612.

Hadwiger, J.A. and R.A. Firtel. 1992. Analysis of Go4, a G-pro-
tein subunit required for multicellular development in Dicty-
ostelium. Genes & Dev. 6: 38–49.

Henikoff, S. and J.G. Henikoff. 1991. Automated assembly of
protein blocks for database searching. Nucleic Acids Res.
19: 6565–6572.

Hinoda, Y., M. Neumaier, S. Hepta, Z. Drzeniek, C. Wagem, L.
Shively, L. Hepta, J. Shively, and R. Paxton. 1988. Molecular
cloning of a cDNA coding for biliary glycoprotein I: Primary
structure of a glycoprotein immunologically crossreactive
with carcinoembryonic antigen. Proc. Natl. Acad. Sci.
85: 6959–6963.

Hooper, N.A., C. Anjard, C.D. Reymond, and J.G. Williams.
1993. Induction of terminal differentiation of Dictyostelium
by cAMP-dependent protein kinase and opposing effects of intracellular and extracellular cAMP on stalk cell differen-
tiation. Development 119: 147–154.

Jermyn, K.A. and J.G. Williams. 1991. An analysis of culmina-
tion in Dictyostelium using prestalk and stalk-specific cell
autonomous markers. Development 111: 779–787.

Jermyn, K., M. Berks, R. Kay, and J. Williams. 1987. Two dis-
tinct classes of prestalk-enriched messenger RNA sequences
in Dictyostelium discoideum. Development 100: 745–785.

Kimmel, A. and R. Firtel. 1983. Sequence organization in Dicty-
ostelium: Unique structure at the 5' ends of protein coding
genes. Nucleic Acids Res. 11: 541–552.

---. 1991. cAMP signal transduction pathways regulating
development of Dictyostelium discoideum. Curr. Opin.
Genet. Dev. 1: 383–390.

Konijn, T.M., D. Barkley, Y.Y. Chang, and J. Bonner. 1968. Cy-
clic AMP: A naturally occurring acrasin in the cellular slime
molds. Am. Nat. 102: 225–233.

Kreft, M., L. Voet, J.H. Gregg, H. Mairhofer, and K.L. Williams.
1993. Evidence that positional information is used to establish
the prestalk-preseore pattern in Dictyostelium discoideum
aggregates. EMBO J. 3: 201–206.

Kubahara, Y., M. Maeda, and K. Okamoto. 1993. Analysis of the
maturation process of prestalk cells in Dictyostelium dis-
coidium. Exp. Cell Res. 207: 107–114.

Kumagai, A., S.K.O. Mann, M. Pupillo, G. Pitt, P.N. Devreotes,
and R.A. Firtel. 1988. A molecular analysis of G proteins and
control of early gene expression by the cell surface cAMP
receptor in Dictyostelium. Cold Spring Harbor Symp.
Quant. Biol. 53: 675–685.

Kumagai, A., M. Pupillo, R. Gunderson, R. Miale-Lye, P.
Devreotes, and R. Firtel. 1989. Regulation and function of Go protein subunits in Dictyostelium. Cell 57: 265–275.

Kumagai, A., J.A. Hadwiger, M. Pupillo, and R.A. Firtel. 1991.
Molecular genetic analysis of two Go protein subunits in
Dictyostelium. J. Biol. Chem. 266: 1220–1228.

Kuspa, A. and W.F. Loomis. 1992. Tagging developmental genes
in Dictyostelium by restriction enzyme-mediated integra-
tion of plasmid DNA. Proc. Natl. Acad. Sci. 89: 8803–8807.

Kyte, J. and R.S. Doolittle. 1982. A simple method for displaying
the hydopathic character of a protein. *J. Mol. Biol.* 157: 105–132.

Louvion, J.F., J.C. Scholder, S. Pinaud, and C.D. Reymond. 1991. Two independent promoters as well as 5’ untranslated regions regulate *Ddras* expression in *Dictyostelium*. *Nucleic Acids Res.* 19: 6133–6138.

Loomis, W. 1969. Temperature-sensitive mutants of *Dictyostelium discoideum*. *J. Bacteriol.* 99: 65–69.

———. 1982. *The development of Dictyostelium discoideum*. Academic Press, New York.

Mann, S.K. and R.A. Firtel. 1987. Cyclic AMP regulation of early gene expression in *Dictyostelium discoideum*: Mediation via the cell surface cyclic AMP receptor. *Mol. Cell. Biol.* 7: 458–469.

———. 1991. A developmentally regulated, putative serine/threonine protein kinase is essential for development in *Dictyostelium.* *Mech. Dev.* 35: 89–101.

———. 1993. cAMP-dependent protein kinase differentially regulates prestalk and prespore differentiation during *Dictyostelium* development. *Development* 119: 135–146.

McGeoch, D.J., A. Dolan, S. Donald, and F.J. Rixon. 1985. Sequence determination and genetic content of the short unique region in the genome of *Herpes Simplex virus type 1*. *J. Mol. Biol.* 181: 1–13.

Mehdy, M. and R. Firtel. 1985. A secreted factor and cyclic AMP jointly regulate cell-type-specific gene expression in *Dictyostelium discoideum*. *Mol. Cell. Biol.* 5: 705–713.

Mehdy, M.C., D. Ratner, and R.A. Firtel. 1983. Induction and modulation of cell type specific gene expression in *Dictyostelium*. *Cell* 32: 763–771.

Morrissey, J.H., K.M. Devine, and W.F. Loomis. 1984. The timing of cell-type-specific differentiation in *Dictyostelium discoideum*. *Dev. Biol.* 102: 414–424.

Obrink, B. 1991. C-CAM (cell-CAM 105) — A member of the growing immunoglobulin superfamily of cellular adhesion proteins. *BioEssays* 13: 227–234.

Pears, C. and J. Williams. 1987. Identification of a DNA-sequence element required for efficient expression of a developmentally regulated and cAMP-inducible gene of *Dictyostelium discoideum*. *EMBO J.* 6: 195–200.

Pears, C., H. Mahhubani, and J. Williams. 1985. Characterization of two highly diverged but developmentally co-regulated cysteine proteinase genes in *Dictyostelium discoideum*. *Nucleic Acids Res.* 13: 8853–8861.

Pitt, G.S., N. Milona, J. Borleis, R.C. Lin, R.R. Reed, and P.N. Devreotes. 1992. Structurally distinct and stage-specific adenyl cyclase genes play different roles in *Dictyostelium* development. *Cell* 69: 305–315.

Powell-Coffman, J.A. and R.A. Firtel. 1994. Characterization of a novel *Dictyostelium discoideum* prespore-specific gene, *PspB*, reveals conserved regulatory elements. *Development* (in press).

Raper, K. 1940. Pseudoplasmodium formation and organisation in *Dictyostelium discoideum*. *J. Elisha Mitchell Sci. Soc.* 59: 241–282.

Ratner, D. and W. Borth. 1983. Comparison of differentiating *Dictyostelium discoideum* cell-types separated by an improved method of density gradient centrifugation. *Exp. Cell Res.* 143: 1–7.

Reymond, C., R. Gomer, M. Mehdy, and R. Firtel. 1984. Developmental regulation of a *Dictyostelium* gene encoding a protein homologous to mammalian ras protein. *Cell* 39: 141–148.

Richardson, D.L. and W.F. Loomis. 1992. Disruption of the sporulation-specific gene *spiA* in *Dictyostelium discoideum* leads to spore instability. *Genes & Dev.* 6: 1058–1070.
LagC is required for cell-cell interactions that are essential for cell-type differentiation in Dictyostelium.

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