

Mutations of the Act Promoter in *Myxococcus xanthus*^{∇†}

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Mutations within the –12 and –24 elements provide evidence that the *act* promoter is recognized by sigma-54 RNA polymerase. Deletion of the –20 base pair, which lies between the two conserved elements of sigma-54 promoters, decreased expression by 90%. In addition, mutation of a potential enhancer sequence, around –120, led to an 80% reduction in *act* gene expression. *actB*, the second gene in the *act* operon, encodes a sigma-54 activator protein that is proposed to be an enhancer-binding protein for the *act* operon. All *act* genes, *actA* to *actE*, are expressed together and constitute an operon, because an in-frame deletion of *actB* decreased expression of *actA* and *actE* to the same extent. After an initially slow phase of *act* operon expression, which depends on FruA, there is a rapid phase. The rapid phase is shown to be due to the activation of the operon expression by ActB, which completes a positive feedback loop. That loop appears to be nested within a larger positive loop in which ActB is activated by the C signal via ActA, and the *act* operon activates transcription of the *csgA* gene. We propose that, as cells engage in more C signaling, positive feedback raises the number of C-signal molecules per cell and drives the process of fruiting body development forward.

Sensing starvation, myxobacteria stop growing and start building fruiting bodies that fill with many thousands of spores (7). The cell-surface-associated, nondiffusible C signal is essential for both fruiting body formation and sporulation (39, 56). C signal is a morphogen, encoded by the *csgA* gene (38), translated as a 25-kDa protein, transported to the cell surface, and finally cleaved to the active 17-kDa signal molecule (40). The signal is transmitted from one cell to another by direct end-to-end contact between them (32–34, 52), which links signaling to the arrangement of cells. Mathematical simulation of C signaling in a population of moving cells has shown that C signal controls the shape of the *Myxococcus xanthus* fruiting body (57, 58). The particular shape of a myxobacterial fruiting body is a species characteristic and is inherited (49, 60).

Cell behavior changes as fruiting bodies form due to changes in the level of C signal, and successive stages of fruiting body development arise from increasing levels of C signal (31, 39). The low initial level of C signal induces traveling wave behavior (20, 37, 55, 70), a higher level produces streaming behavior, and the highest level induces spores to form. Cells in a stream are arranged end to end, all following the same trajectory, which leads to frequent end-to-end contacts (22–24). Streaming cells enlarge tiny initial aggregates by forming an onion-like succession of spherical shells around the aggregate. The cells within each shell stream in roughly circular orbits. This leads to a spherical outer domain of densely packed cells surrounding an inner domain of threefold-lower density, which is observed microscopically (28, 53). The level of C signal also regulates the expression of many genes involved in fruiting body devel-

opment (35), in a timely way. Sporulation genes have a higher threshold of C-signal intensity than genes for aggregation (31, 39), and the highest level of C signal induces the cells to differentiate into spores within the fruiting body (37). This property confines sporulation to the spherical mass of a *Myxococcus* fruiting body, while cells around the outside with lower C-signal levels do not sporulate (25, 46).

Thus, cell movement and gene expression are coordinated for *M. xanthus* fruiting body development by the rising number of C-signal molecules per cell (27). Transcription of *csgA* is initiated by starvation and the stringent response (5). Later *csgA* expression is enhanced by the *act* operon (17). Mutations in *act* were found to affect the expression of developmental gene reporters that are expressed after 6 h of development and are related to aggregation and sporulation (16). Moreover, deletion of the *actC* gene led to the premature formation of many small fruiting bodies (17)—an effect similar to a 10-fold overexpression of CsgA (37). By contrast, deletion of the *actD* gene led to delayed formation of large fruiting bodies. The kinetics of C-signal increase thus appears to control fruiting body size. After initiation of C-signal expression by starvation, how do the Act proteins regulate the rate and magnitude of the coordinating increase in the signal level?

MATERIALS AND METHODS

Cultures. Bacterial strains and plasmids are listed in Table 1. The conditions for growth; development, including sporulation; and electroporation have previously been described (17). Transduction was carried out as described previously (16).

Preparation of RNA from developing cells. A culture of 1×10^{10} to 2×10^{10} cells was induced to develop at 32°C on plates consisting of TPM buffer (10 mM Tris-HCl, pH 7.6, 1 mM KPO₄, pH 7.6, 8 mM MgSO₄) solidified with 1.5% Bacto Agar. Bacteria, or spores, were harvested at the specified time from the surface by scraping, suspension, and washing in TPM buffer. The solid material from harvest was resuspended in lysozyme-containing buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 2 mg/ml lysozyme) and then disrupted with a Branson 450 Tip sonifier, to ensure that both cells and prespores would be lysed and homogenized. RNA was isolated from the homogenate using RNeasy Midi Spin columns (QIAGEN).

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TABLE 1. Strains and plasmids

Strain or plasmid	Relevant characteristic(s)	Reference or source
<i>M. xanthus</i> strains		
DK1622	Wild type	26
DK5208	Tn5- <i>I32::csgA</i>	35
DK7853	<i>asgA476</i>	15
DK10602	DK1622::pTG021	This work
DK10603	$\Delta actB$	17
DK10604	$\Delta actC$	17
DK10605	$\Delta actA$	17
DK10627	Mx8[MS1512] \rightarrow DK10605 ^a	This work
DK10641	DK1622::pREG1727	This work
DK10642	DK1622 <i>att</i> Mx8::pTG027	This work
DK10643	Mx8[DK10642] \rightarrow DK5208	This work
DK10644	Mx8[DK10642] \rightarrow DK7853	This work
DK10645	DK10642::pEE106; SalI-NdeI	This work
DK10646	Mx8[DK10642] \rightarrow DK10605	This work
DK10647	Mx8[DK10642] \rightarrow DK10603	This work
DK10648	Mx8[DK10642] \rightarrow DK10604	This work
DK10649	Mx8[DK10642] \rightarrow MS1512	This work
DK10650	Mx8[DK10642] \rightarrow <i>fruA::TcΩV</i>	47 and this work
DK10651	Mx8[DK10602] \rightarrow DK10605	This work
DK10652	Mx8[DK10602] \rightarrow DK10603	This work
DK10653	Mx8[DK10602] \rightarrow DK10604	This work
DK10654	Mx8[DK10602] \rightarrow DK5208	This work
DK10655	Mx8[DK10602] \rightarrow DK7853	This work
DK10658	DK1622 <i>att</i> Mx8::pTG033	Table S2 ^b
DK10659	DK1622 <i>att</i> Mx8::pTG037	enh1-1 (Table S2)
DK10660	DK1622 <i>att</i> Mx8::pTG038	enh2-1 (Table S2)
DK10661	DK1622 <i>att</i> Mx8::pTG039	σ^{54} 5 C \rightarrow T (Table S2)
DK10662	DK1622 <i>att</i> Mx8::pTG040	σ^{54} 6 C \rightarrow Δ (Table S2)
DK10665	DK1622::pTG042	σ^{54} 1 T \rightarrow C (Table S2)
DK10666	DK1622::pTG043	σ^{54} 7 T \rightarrow C (Table S2)
DK10667	DK1622::pTG044	σ^{54} 2 G \rightarrow T (Table S2)
DK10668	DK1622::pTG045	σ^{54} 3 G \rightarrow T (Table S2)
DK10669	DK1622::pTG046	σ^{54} 4 C \rightarrow T (Table S2)
DK10670	DK1622::pTG047	enh1-2 (Table S2)
DK10671	DK1622::pTG048	enh2-2 (Table S2)
DK10672	DK1622::pTG049	enh3 (Table S2)
DK10673	DK1622::pTG050	C box 1 (Table S2)
DK10674	DK1622::pTG051	C box 2 (Table S2)
DK10675	DK1622::pTG052	σ^{54} 8 G \rightarrow T (Table S2)
MS1512	<i>sdeK</i> (Ω 4408)	11
Plasmids		
pBluescriptII SK+	3.0 kb; Amp ^r (<i>bla</i>) <i>lacZ</i> with multiple cloning site for blue-white screening	Stratagene
pEE106	Promoter and part of <i>fruA</i> gene in pBGS18 (59)	8
pJBZ281	7.2 kb; Kan ^r ; polylinker sites upstream of <i>lacZ</i> for translational fusions	M. R. K. Alley
pREG1727	20.7 kb; Amp ^r Kan ^r Mx8 attachment site <i>attP</i> ; transcriptional terminators followed by multiple cloning sites upstream of promoterless <i>lacZYA</i> genes for transcriptional fusions	R. Gill (10)
pSWU12	pBGS18 with Tn903 Kan ^r deleted and replaced with pKS-Tet ^r cassette; MluI/EcoRI	71
pTG021	pJBZ281::pTG108 (sequence bp 496 to 2192); SmaI-SmaI	This work
pTG025	pTG021::pBluescriptII SK+; EcoRI-BamHI	This work
pTG027	pREG1727::pTG025; EcoRI-XhoI	This work
pTG033	pBluescript/pREG1727::PCR fragment with <i>act</i> promoter region from upstream XhoI to downstream BamHI; BamHI-XhoI	This work
pTG108	Deletion of PstI fragment from pLAG66 (13); PstI-PstI	This work

^a Can be read as "a stock of Mx8 grown on MS1512 was used to transduce DK10605."

^b Table S2 can be found in the supplemental material.

Quantifying mRNA. To prepare slot blots, 25 μ l of each developmental RNA sample dissolved in water was diluted by adding 25 μ l 2 mM EDTA, 30 μ l 2 \times SSC (1 \times SSC is 0.15 M NaCl plus 0.015 M sodium citrate) (54), and 20 μ l 37% formaldehyde. The entire 100- μ l volume of the mixture was then heated at 60°C for 15 min and transferred to a Hybond-N nylon membrane (Amersham) with 10 \times SSC, according to the protocol for Bio-Dot SF (Bio-Rad). Finally the membranes were irradiated with UV light to cross-link the RNA to them. DNA

probes for *actA*, *actB*, and *actE* were labeled with ³²P (17). Probes were annealed onto the RNA-loaded membranes, as described for Southern blotting (54).

Site-directed mutagenesis. A two-step method of overlap extension PCR was employed, whose principles have been described previously (19). Two universal primers designated upstream XhoI and downstream BamHI were designed. Each universal primer had the indicated synthetic restriction site, following a terminal AAAG spacer. The upstream XhoI primer sequence is 5' AAAGCTC

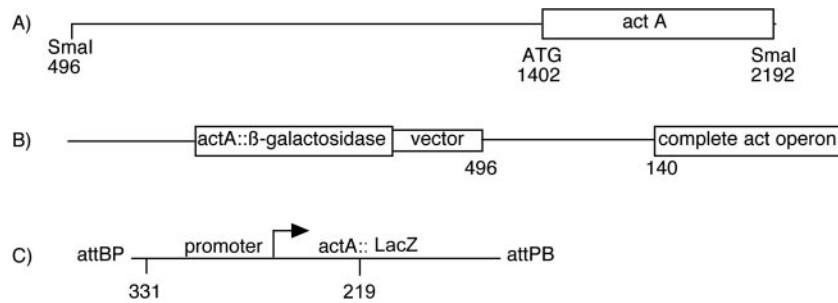


FIG. 1. Map of the *act* operon. (A) DNA fragment containing the region upstream of the operon and the N-terminal part of the ActA protein. (B) The fragment shown in panel A was cloned into pJBZ281, and the plasmid was inserted by homologous recombination into the *act* region of DK10602, a translational fusion. (C) The transcriptional fusion plasmid pTG027 was inserted in strain DK1622, creating DK10642. Position 331 is on the universal PCR primer, described in Materials and Methods.

GAGGCGGCTCGTCCGGACACCTTC 3'. The downstream BamHI primer is 5' AAAGGGATCCCACCGCCTCTTCGTCATCCAC 3', with the *act* operon sequences underlined in both sequences. From these two primers, the *act* promoter region starting at nucleotide (nt) 331 in the upstream region of the GenBank sequence AF350253 and running to nucleotide 2501 of that sequence near the beginning of *actB* was amplified by PCR. The XhoI and BamHI restriction sites oriented the PCR products for cloning into pBluescript or pREG1727.

To introduce particular deletions or point mutations at predetermined sites between nucleotides 331 and 2501, a set of mutagenic primers, shown in Table S2 in the supplemental material, was designed for both template strands. Each primer, located at the center of the cloned promoter region, contained 10 nucleotides of wild-type *act* sequence, followed by the desired mutant sequence, followed by another 10 nucleotides of the wild-type sequence. The first step of mutation induction involved two sets of 20 to 25 cycles of PCRs with one universal primer (either the upstream XhoI primer or the downstream BamHI primer) and one of each of the mutagenic primers. The mutagenic primer design ensures that pairs of first-step PCR products overlapped by at least 20 bp with the other member of the pair. The PCR products were purified by agarose gel electrophoresis. The second step involved the upstream XhoI primer, the downstream BamHI primer, and the two PCR products from the first step overlapping at the same mutation from the first step, and the complementary strands that formed a duplex in the overlapping mutant region were completed. This second-step PCR product, which contained a duplex version of the mutant sequence at the appropriate internal site, was digested with XhoI and BamHI and then cloned into the XhoI-BamHI site of pBluescript, for structure verification by sequencing. Verified clones were transferred from pBluescript into the XhoI-BamHI site of pREG1727. Plasmid pREG1727, constructed by Ron Gill of the University of Colorado and described in reference 10, forms transcriptional fusions between the cloned fragment and a promoterless *lacZ* gene. Cloning into the XhoI-BamHI site ensured that the promoter would be oriented properly for transcription of *lacZ*. The cloned mutant plasmids are described in Table 1. pREG1727 contains the prophage attachment site *attP* of myxophage Mx8 which promotes site-specific integration of this plasmid into the *attB* locus of the *M. xanthus* chromosome (12). pREG1727 also carries the NPTII gene, encoding kanamycin resistance, which can be used for selection of a plasmid integrant into DK1622, since pREG1727 cannot replicate as a plasmid in *M. xanthus* (12). That each new plasmid had integrated into the *attB* locus of DK1622 was confirmed by Southern blot hybridization of the products of restriction endonuclease digestion. The mutated sequence was again verified. Finally, each integrant strain was induced to develop, and its β -galactosidase activity was measured relative to that of wild type in the same experiment, as described previously (16). Each expression test was performed three to six times, to obtain a reliable average. Specific activities were measured as nanomoles of *o*-nitrophenol per minute per milligram of protein.

Nucleotide sequence accession number. The sequence of the *act* operon DNA is published in GenBank under accession number AF350253.

RESULTS

Dynamics of *act* operon expression. A translational fusion between ActA, the first protein of the operon, and LacZ was constructed for initial measurements of *act* operon expression.

A 1,696-bp *SmaI* restriction fragment that starts 906 bp upstream of the *actA* (MXAN3213) translation start, which is located at position 1402 on the fragment shown in Fig. 1A, was inserted into the *SmaI* site of plasmid pJBZ281. The *SmaI* site at position 496 lies within MXAN3211; consequently the cloned fragment includes MXAN3211, the 333-bp intergenic region upstream of *actA*, and the N-terminal 263-amino-acid polypeptide of ActA translationally fused to LacZ. This fusion plasmid integrated by homologous recombination into the *act* region of the chromosome of DK1622, the strain which provided a standard genetic background for this work. Insertion generated a tandem duplication strain, DK10602; its chromosomal structure, which was confirmed by Southern analysis, is shown in Fig. 1B. The fruiting body sporulation efficiency of DK10602 was equal to that of DK1622: at 3 days, there were 6.4×10^5 viable spores for DK1622 and 6.0×10^5 spores for DK10602. Therefore, the 906-bp upstream segment which includes MXAN3211 and the 333-bp intergenic region in DK10602 contains all the necessary positive elements for *act* expression (Fig. 1B).

Developmental expression of the *act* operon, as measured in DK10602, appears to rise in two distinct phases (Fig. 2). It increases gradually during the first 18 h of development and then rises rather steeply to a maximum at 21 to 24 h, when

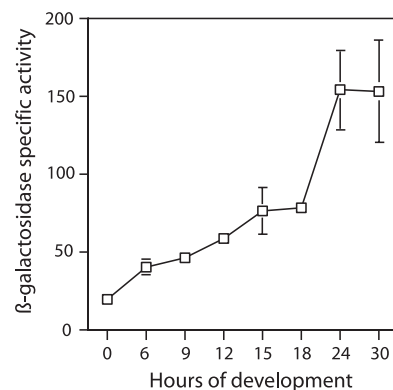


FIG. 2. Expression of β -galactosidase in the translational fusion strain DK10602. Specific β -galactosidase activity is shown during the first 30 h of development with error bars showing the standard deviations. Where error bars are not evident, they lie inside the square symbols centered on the average.

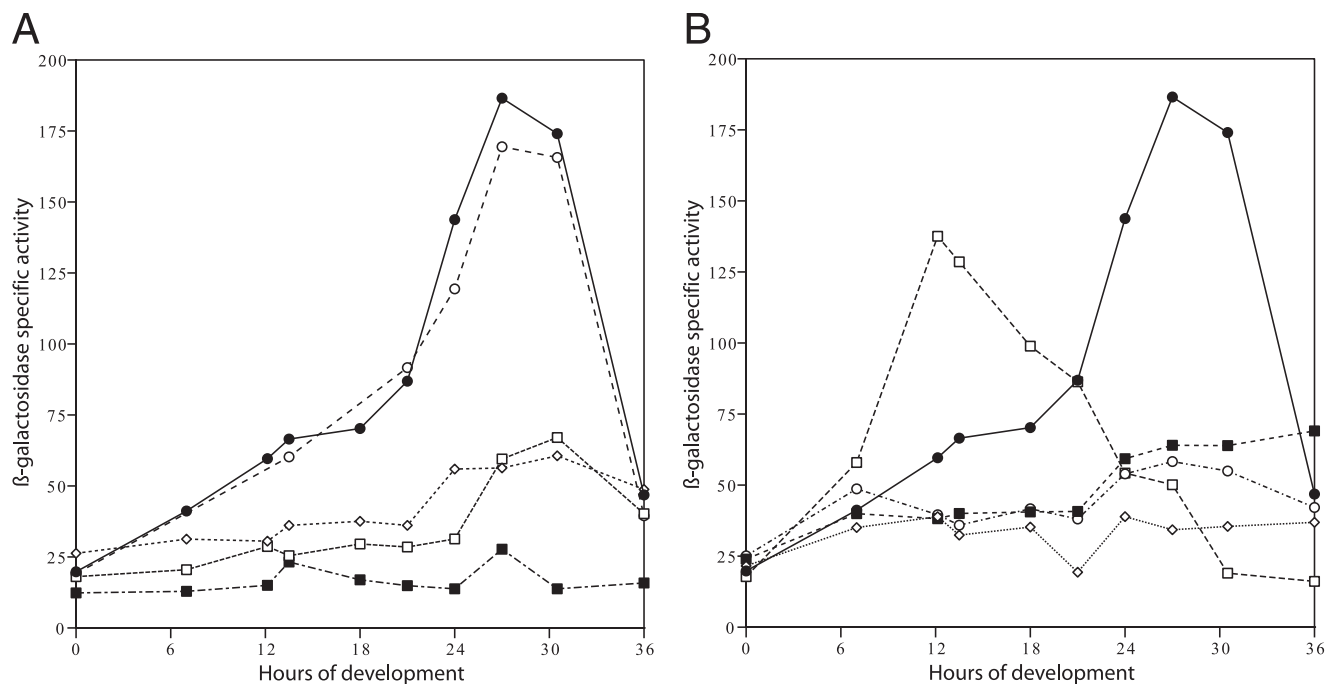


FIG. 3. A. Transcription of β -galactosidase in DK10642 and mutants defective in *asgA*, *fruA*, and *sdeK*. Closed circles, DK10642; open circles, *sdeK* mutant; open squares, *fruA* plasmid insertion mutant; closed squares, *fruA* Tn5 insertion mutant; open diamonds, *asgA* mutant. B. Transcription of β -galactosidase in DK10627 (closed circles) and mutants defective in *csgA* (open diamonds), *actA* (closed squares), *actB* (open circles), and *actC* (open squares). Specific activity is shown as nanomoles of *o*-nitrophenol per minute per milligram of protein.

heat-resistant spores begin to appear. A single, continuous rise in expression beginning at 6 h seems incompatible with the experimental data, given the size of measurement errors shown by the bars in Fig. 2. To probe the existence of two distinct phases of *act* expression, the same 906-bp upstream sequence of strain DK10602 was cloned into pREG1727, to generate a transcriptional fusion of *actA* to *lacZ*, and to build a reporter plasmid, pTG027. This fusion plasmid has translational stops in all reading frames ahead of *lacZ*, ensuring that the fusion of *actA* to *lacZ* is transcriptional, not translational. A transcriptional fusion eliminates any effects that translational fusion might have that limit protein synthesis. Plasmid pREG1727 is designed to integrate by site-specific recombination into the Mx8 *attB* site of *M. xanthus* (12), which is distant from the chromosomal *act* region. Figure 1C shows pTG027 inserted into the chromosome of DK1622 at the Mx8 prophage *attB* site, creating strain DK10642. The chromosomal *act* region was found to be undisturbed in this strain; its expression should be normal, and pTG027 should be an accurate reporter. Southern blot assays performed on DK10642 supported the idea that insertion had occurred within the *attB* site and not within the *act* operon. The spore titer of DK10642 (6.1×10^5) was the same as that of its DK1622 parent (6.4×10^5), confirming that the chromosomal *act* region was normal. The solid line with filled points in Fig. 3A shows measurement of DK10642 *lacZ* expression. Again, two phases of *lacZ* expression are evident. A gradual increase during the first 18 h is followed by a rapid increase to 24 h. Because the expression profiles of Fig. 2 and Fig. 3 are similar, it is likely that the two transcriptional phases seen with DK10642 are responsible for the two phases of translation seen with the fusion DK10602. Thus, Act protein

levels are primarily controlled at the level of transcription in these strains, and they can be used as sensitive reporters of transcription.

To find transcriptional regulators that are needed for *act* operon expression, pTG027 with the *act* insert was transduced or transfected into a series of candidate regulatory mutant strains derived from DK1622. Figure 3A shows evidence for two similar phases of *act* expression in an *sdeK* mutant background. The very similar time course of the Δ *sdeK* strain compared with DK10627, which has the wild-type allele of *sdeK*, indicates that *act* transcription does not depend on *sdeK*. The *sdeK* gene encodes a histidine protein kinase that has been shown to be essential for expression of many developmental genes (11, 36, 48). The substrate(s) for the putative *sdeK* kinase has not yet been identified, and the data in Fig. 3 imply that neither FruA, for which there is evidence of phosphorylation correlated with activity (8), nor any of the *act* products—among which ActB is likely to be phosphorylated—is a substrate for the *sdeK* kinase. However, it is evident from Fig. 3A that transcription of *act* depends on *asgA*, since expression is greatly depressed in that mutant. Expression of *act* also depends on the response regulator protein FruA, which has been verified with two different *fruA* insertion mutations (8, 47). The Ogawa *fruA* plasmid insertion mutant happens to be in a DZF1 background that contains a *pilQ1* motility mutation (47), which may account for the slightly lower level of *act* expression than in the Ellehaug Tn5 insertion mutant, which is *pilQ*⁺. Expression of *fruA* depends on the A signal (8). Together, the *asgA* and *fruA* defects in *act* operon expression imply a FruA requirement for the transcription of *act*.

The second phase of *act* expression that is evident in Fig. 2

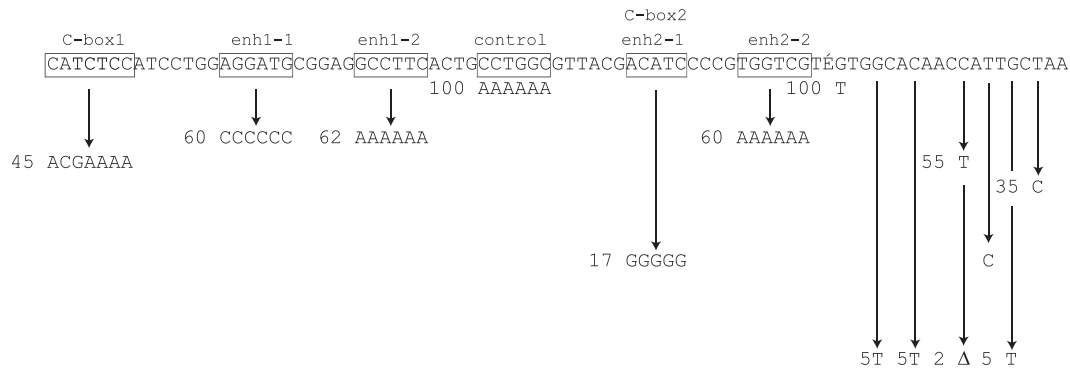


FIG. 4. DNA sequence of the *act* promoter region from residues -167 to -12 . The transcription start, AGC, is residue $+1$, and it is located to the right of the sequence shown. The -12 and -24 regions of a sigma-54 promoter, two potential enhancer sequences, and two potential C boxes are included. Boxed elements in the line were replaced by the sequences shown below them at the arrow tips, and the numbers next to the new sequence give the resulting level of *actA::lacZ* fusion reporter expression, expressed as a percentage of the unmutated sequence. Specific β -galactosidase activities relative to the wild-type promoter were measured as described in Materials and Methods. Site-specific mutations were introduced with the PCR primers presented in Table S2 in the supplemental material.

and Fig. 3A rises steeply at 20 h when CsgA protein is reaching its highest specific activity under the same conditions (17). The data of Fig. 3B indicate that transcriptional expression of *act* directly depends on C signal for several reasons. First, there is very little, if any, increase in *act* expression in a *csgA* mutant. Second, the $\Delta actC$ mutant, which gives precocious *csgA* expression (16), shows precocious expression of the *actA* transcriptional reporter. This argues that the level of C signal normally limits *actA* expression. The steepness of the early rise in the $\Delta actC$ expression profile in Fig. 3B recalls the second phase of *act* expression in DK10642 cells. For that reason, it appears that in the $\Delta actC$ mutant phase two is precocious while phase one is truncated. Together, the effect of the null *csgA* and of the $\Delta actC$ mutants implies that the steep phase of *act* expression depends on C signaling. The early phase is depressed in both the *asgA* mutant and in two *fruA* mutants (Fig. 3A).

A promoter for *act*. Figure 3B shows that *act* reporter expression is substantially and equally depressed in the $\Delta actA$ and $\Delta actB$ mutants. Inasmuch as *actB* encodes a sigma-54 activator protein (15), and *actA* encodes a response regulator that works with *actB* (17), the dependence of *act* expression on *actB* strongly suggests that the *act* operon is expressed from a sigma-54 promoter, and that inference has been tested by site-directed mutagenesis.

When the *actA*, *actB*, *actC*, and *actD* genes are expressed as an operon, they are transcribed from a single adenosine residue 57 nt ahead of the presumed *actA* translational start (17). The sequence ahead of the transcription start site, taken from the GenBank sequence with accession number AF350253, is shown in Fig. 4. The entire sequence was verified in the complete sequence of the *M. xanthus* genome, GenBank accession number CP000113. A proposed ribosome binding site (TGAG GAAGT) and the ATG translation start codon of *actA* (nucleotide $+1$) that lie to the right of the sequence shown in Fig. 4 are evident in the published sequence. Relative to the transcription start, the upstream region from -167 to -12 is spelled out in Fig. 4. It includes -15 to -29 with a TGGCA C-N₅-TTGC sequence resembling the sigma-54 consensus from a compilation of sigma-54 promoters (2). That compilation shows that transcription initiates from 8 to 21 nt down-

stream from the highly conserved GC element in the proximal (-12) element of a sigma-54 promoter (2). In the *act* operon, transcription starts 15 nt from that GC element. This resembles the extensively studied *M. xanthus* sigma-54 *pilA* promoter (71). It also resembles the *M. xanthus* $\Omega 4521$ promoter, for which sequence mutations have indicated sigma-54 specificity (29). The sigma-54 gene (*rpoN*), which is in single copy, is essential for *M. xanthus* development. Uniquely among bacteria checked, *rpoN* of *M. xanthus* is also essential for growth (30).

Individual nucleotides in the promoter-like sequence were altered and then tested for level of *act* expression. Mutant promoters were generated by sequence overlap extension PCR (Materials and Methods), using the oligonucleotides shown in Table S2 in the supplemental material. The mutated segments were then individually cloned into pREG1727, replacing the normal sequence with its mutant versions in pTG027. This cloning incorporated each of them into the same transcriptional fusion segment so that the activity of each mutant promoter could be quantified by measuring the β -galactosidase activity relative to that of the sequence without mutation. β -Galactosidase specific activity was measured 24 h after initiation of development, the time at which maximum *act* expression was observed from the normal promoter, as indicated in Fig. 2 and Fig. 3. To compensate for the biological variation in the level of gene expression, each mutant-to-wild-type comparison was made within the same experiment. Figure 4 presents the relative β -galactosidase activities observed in each mutant as relative promoter activity in percent. Changes in the -14 to -20 region, which is the downstream or proximal oligonucleotide element of the promoter (the -12 element), show that those bases are important for expression. By the change of residue -14 from T to C, activity fell to 34% of the original level; changing residue -16 from G to a nonconsensus T reduced activity to 7% of that of the wild-type promoter. Changing residue -18 from T to C reduced activity to 20% of the original level, and at -20 changing C to T decreased activity to 50% of the original level. Changes in the upstream promoter element (the -24 element) at residue -24 or -27 decreased activity to less than 10% of that of the wild-type promoter. By

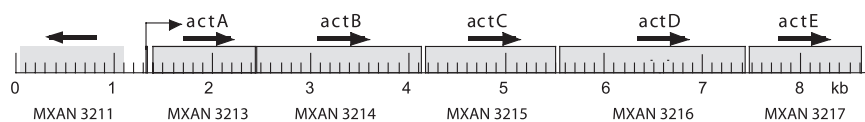


FIG. 5. The *act* operon, showing the location of all *act* genes, including *actE*. Horizontal arrows indicate the promoter and direction of transcription of each gene.

contrast, changing the -30 G residue to T had no effect, suggesting that -30 does not have a significant interaction with sigma-54 holoenzyme (Fig. 4). If the *act* promoter were recognized by a member of the sigma-70 family of factors, assuming that -30 is part of the -35 region, an effect would have been expected (18). *M. xanthus* is known to employ several members of the sigma-70 family of factors (67).

One of the largest mutant defects (93%) was obtained by deleting the residue at position -21 , which lies between the upstream and downstream elements of a sigma-54 promoter. This defect exemplifies a requirement for proper spacing between the -12 and -24 elements, as has been observed elsewhere (3, 29, 43–45). Strict spacing between promoter elements helps to distinguish sigma-54 promoters from sigma-70 promoters (2, 18). Considering the conserved residues, the interelement spacing, and the distance from the *act* transcription start, the data strongly support recognition by sigma-54.

Upstream regulatory sequences. Sigma-54 promoters often have upstream regulatory elements. To identify regulatory elements, five to seven base pairs were replaced in each of the potential regulatory sites, indicated at the top of Fig. 4. Kroos and coworkers have identified a set of septanucleotide sequences, named C boxes, near the transcription start of a number of established C-signal-dependent promoters (9, 41, 61, 66, 73–75). Expression of the *act* operon is C signal dependent (17), but C boxes have yet to be reported with sigma-54 promoters. Two sequences that resemble the C-box consensus CAYCCY (where Y = C or T) (41) are evident -167 and -120 nt upstream of *actA* (Fig. 4). Both sequences were mutated, replacing all seven or four of seven bases with nonconsensus residues. Replacing all five residues of potential box 2 lowered activity to 17% of the wild-type level, a substantial decrease (Fig. 4). Changing all seven residues of potential box 1 significantly decreased the activity to 45% of the wild-type level (Fig. 4). These changes may be contrasted with mutation of a hexanucleotide sequence, “control,” in the same general area, centered at -130 . Replacement of all six bases of this sequence, which lies between the two C boxes and between the

two enhancer-like sequences described below, was found to have no detrimental effect (Fig. 4).

Holoenzyme $E\sigma^{54}$ forms physically detectable closed-promoter complexes at known σ^{54} promoters, but the enzyme is unable to initiate transcription spontaneously. To separate the polynucleotide strands of an $E\sigma^{54}$ -DNA complex requires an enhancer sequence in the DNA (42). A promoter-specific activator protein binds the enhancer to interact, by DNA looping, with holoenzyme bound at the promoter (51, 62, 68). Two candidate sequences, which are boxed in Fig. 4, were identified by comparing the sequence upstream of the *act* promoter with several sigma-54 enhancers that have been studied—the enhancers for *glnAp₂* and *pspA* in *Escherichia coli* (50, 69) and for *nodD3* in *Sinorhizobium meliloti* (6). Two sequences upstream of the *act* promoter are indicated, one located around -150 (candidate enh1) and another sequence located around -110 (candidate enh2). Candidate enhancer 2 includes C box 2 (Fig. 4). Because the *glnAp₂* enhancers have dyad symmetry, the two half-sequences were separately mutated. Localized multibase mutations were obtained by substituting either A₆, C₆, or G₅ for the five or six resident bases (see Table S2 in the supplemental material). Candidate half-sequence 2-1 proved important for expression of the *act* operon, since expression in the substitution mutant fell to 17% of the wild-type level (Fig. 4). Mutation of candidate elements 1-1, 1-2, and 2-2 showed expression reduced to 60%, 62%, and 60% of wild-type levels, respectively (Fig. 4). The possibility that A₆ bends DNA should be noted.

actE. Examination of the *act* DNA sequence revealed an open reading frame that starts just downstream of *actD* and that could encode a polypeptide of 753 amino acids. Figure 5 presents a fine-scale map of the whole region. The newly recognized gene, MXAN3217, named *actE*, is cooriented with the other *act* genes and is likely to be cotranscribed with *actA* and *actB*, according to the time pattern of hybridization of developmental RNA to open reading frame-specific probes (Fig. 6). Deletion of *actB* decreases the amount of *actE* transcript to the same extent as it decreases *actA* and *actB* transcript levels, as if all were transcribed together. The *actE* stop codon is at

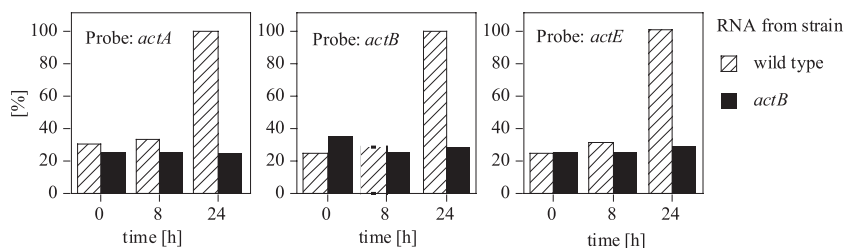


FIG. 6. Relative levels of *act* operon mRNA at 0, 8, and 24 h of development are shown expressed as percentages. The 24-h level was set as 100%. DNA probes specific for *actA* and *actB* were described in reference 17, and an *Apal* fragment was employed for *actE*. The mRNA levels were quantified by measuring ^{32}P on slot blots as described in Materials and Methods.

nucleotide residue 8591 in the GenBank sequence with the accession number AF350253. A 7.3-kb *act* mRNA has been reported elsewhere (17)—just long enough to extend from the beginning of *actA* through the end of *actE* (Fig. 5). Mutations in *actE* have not yet been sought, and no sequence homologs have been found to suggest function.

DISCUSSION

The effects of sequence changes in the *act* operon promoter of *M. xanthus* are strong evidence that the promoter is recognized by sigma-54. Considering the large number of sigma-54 activator proteins encoded by the genome (13, 21), and with this addition to the list of established sigma-54 promoters, it is evident that sigma-54 regulates a substantial part of fruiting body development. The mutations reported here lie in the right half of the 333-bp segment that separates the *actA* (MXAN3213) transcription start from the start of MXAN3211, an open reading frame which, if expressed, is expected to be transcribed in the opposite direction from *act* (Fig. 5). Two candidate sigma-54 enhancer elements were found, one around -150 and one around -110. Site-specific mutagenesis of those two elements led to an 80% activity reduction from the wild type when the -110 element was mutated and a smaller, but significant, reduction when the one at -150 was mutated. Direct evidence for enhancer function is lacking.

The product of *actB* is shown here to be an activator of *act* operon expression, since deletion of *actB* decreases expression of *actA* (Fig. 3B) and of *actE* (Fig. 6). ActB might interact with one or both of the potential enhancers, but no binding studies have been carried out. ActA and ActB appear to be part of a signal transduction pathway that receives input from C signal on the transmitting cell. The output of this signal transduction pathway would be a transcriptionally active ActB (perhaps phosphorylated ActB [ActB~P]), whose absence would account for the reduced *act* operon expression in a C-signal-deficient or an ActA-deficient mutant (Fig. 3B). Thus, the rapidly rising late phase of ActA and ActB expression evident in Fig. 2 and Fig. 3 could be explained by ActB binding to the enhancer of its own promoter. Because a positive feedback loop would thereby be created, an accelerated rise in expression would be expected.

Expression of the *act* operon starts by 6 h of development, after the traffic jams that nucleate aggregates have formed (28). Ellehaug et al. have shown that A signaling, but not C signaling, is required for *fruA* expression (8). FruA protein is a DNA-binding transcriptional activator that is necessary for expression of a number of developmentally regulated genes, *devRS*, $\Omega 4400$, $\Omega 4469$, $\Omega 4273$, $\Omega 4500$, and *fdgA* (8, 47, 64, 75), as well as for *act*. FruA belongs to the FixJ family, and it begins to be expressed between 3 and 6 h (8, 47), in preparation for aggregation gene expression, including the *act* genes. ActB belongs to the NtrC family of transcriptional activators, which have an ATPase domain located between their receiver and DNA-binding domains that FruA lacks. It is attractive to think that FruA is a positive transcriptional regulator of the *act* operon for the early phase of *act* operon expression that occurs at a lower rate than the late phase because ActB has an ATPase activity to drive promoter opening while FruA does not.

Early-phase expression of the *act* operon would produce some ActB protein. ActB could then bind one or more specific enhancers. Using its ATPase to open the sigma-54 promoter, ActB would initiate the second, rapidly rising phase of *act* transcription. In addition to *actB*, *csgA* and *actA* are also needed for the second, more rapid phase of *act* expression (Fig. 3B). p17 CsgA on the transmitting cell, via a (currently unknown) receptor on the receiving cell, would signal ActA, which, in turn, would activate ActB, presumably to ActB~P. With active ActB protein, expression of the entire *act* operon would accelerate, rapidly producing all the Act proteins, including ActC and ActD for timing.

Recently it has been suggested that the *actD* gene, MXAN3216, is a cysteine protease homolog that belongs to clan CD, which includes family C14 of caspase-like enzymes EC 3.4.22.36 and which is directed toward particular proteins (1, 4, 63, 65). Members of this family have a His/Cys catalytic dyad around which sequence is conserved. In *actD* the His/Cys residues are found in the consecutive sequences LVYYS $\underline{\text{GHS}}$ and LDS $\underline{\text{CASG}}$ (with putative catalytic residues underlined). Because deletion of *actD* delays C-signal-dependent gene expression, the normal timing of *csgA* expression is likely to involve proteolysis. Since deletion of *actC* leads to precocious C-signal-dependent gene expression, *actC* protein appears to inhibit the rise of *csgA* expression. One scheme for the temporal control of *csgA* expression would start with production of comparable amounts of ActC and ActD proteins from their coexpression, and ActC (MXAN3215) would inhibit *csgA* expression. ActD caspase could degrade ActC, releasing the inhibition and allowing the expression of *csgA* to rise at the appropriate time. The final level of *csgA* expression is set by *actA* and *actB* (17). Since C signaling stimulates *csgA* expression, ActB~P might be a transcription factor or an activator of a transcription factor for the *csgA* promoter.

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