Allele-specific suppression of the temperature sensitivity of *fitA/fitB* mutants of *Escherichia coli* by a new mutation (*fitC4*): isolation, characterization and its implications in transcription control

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The temperature sensitive transcription defective mutant of *Escherichia coli* originally called *fitA76* has been shown to harbour two missense mutations namely *pheS5* and *fit95*. In order to obtain a suppressor of *fitA76*, possibly mapping in *rpoD* locus, a Ts⁺ derivative (JV4) was isolated from a *fitA76* mutant. It was found that JV4 neither harbours the lesions present in the original *fitA76* nor a suppressor that maps in or near *rpoD*. We show that JV4 harbours a modified form of *fitA76* (designated *fitA76**) together with its suppressor. The results presented here indicate that the *fit95* lesion is intact in the *fitA76** mutant and the modification should be at the position of *pheS5*. Based on the cotransduction of the suppressor mutation and/or its wild type allele with *pps, aroD* and *zdj-3124*::Tn10 kan we have mapped its location to 39·01 min on the *E. coli* chromosome. We tentatively designate the locus defined by this new extragenic suppressor as *fitC* and the suppressor allele as *fitC4*. While *fitC4* could suppress the Ts phenotype of *fitA76** present in JV4, it fails to suppress the Ts phenotype of the original *fitA76* mutant (harbouring *pheS5* and *fit95*). Also *fitC4* could suppress the Ts phenotype of a strain harbouring only *pheS5*. Interestingly, the *fitC4* Ts phenotype could also be suppressed by *fit95*. The pattern of decay of pulse labelled RNA in the strains harbouring *fitC4* and the *fitA76** resembles that of the original *fitA76* mutant implying a transcription defect similar to that of *fitA76* in both these mutants. The implications of these findings with special reference to transcription control by Fit factors *in vivo* are discussed.

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1. Introduction

Regulation of gene expression in prokaryotes occurs mostly at the level of transcription. This involves use of multiple sigma factors, antisigma factors and several macromolecular elements collectively called "accessory transcription factors" (Yura and Ishihama 1979; Ishihama 1988, 1993, 2000). These factors interact either with DNA or RNA

polymerase or both and confer selectivity on the process and thus regulate gene expression. Our laboratory has been studying the control of transcription by accessory transcription factors in *Escherichia coli* for a long time. Our earlier work in this area has been reviewed (Jayaraman 1994). Specifically, two genes have been identified designated *fitA* and *fitB*, mapping close to each other at 38.7 min. The products of these genes are believed to interact with each other

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as well with RNA polymerase and control the expression of few classes of genes (which might include some of the genes coding ribosomal proteins). A tentative model has been proposed to explain the interaction between FitA and FitB products with each other as well as with RNA polymerase. These conclusions stem from the initial isolation and characterization of a temperature sensitive transcription defective mutant (fitA76) and two of its suppressors (fitA24 and fitB) and modulation of growth properties of these mutants by four rpoB mutations (rpoB240, rpoB7, rpoB55 and rpoB42) in an allele specific manner (Jabbar and Jayarman 1976, 1978; Jayaraman and Jabbar 1980; Dass and Jayaraman 1985a,b, 1987; Munavar and Javaraman 1987; Munavar et al 1993). Polar effect of Tn5 insertions in fitA on fitB expression suggested that fitA and fitB could form an operon by themselves or be components of one, the direction of expression being $fitA \rightarrow fitB$ (Munavar 1991). Initially, it was believed that the fitA and fitB mutations defined hitherto unidentified genes and the lesions could be single base change in the respective

Molecular characterization of a recombinant plasmid clone harbouring 2.1 kb fragment from wild type E. coli which complemented the Ts phenotype of the *fitA76* mutant and the corresponding chromosomal region of fitA76 mutant revealed that fitA76 harbours two missense mutations: a $G_{293} \rightarrow A_{293}$ transition in *pheS* locus (which codes for alpha subunit of phenylalanyl tRNA synthetase) and the other named fit95 (possibly located in the pheT locus which codes for the beta subunit of the phenylalanyl tRNA synthetase). The presence of the same $G \rightarrow A$ transition at position of 293 of pheS gene in the temperature sensitive translation defective pheS mutant namely pheS5 (Kast et al 1992) and absence of transcriptional abnormalities characteristic of fitA76 in either pheS5 or fit95 mutants justify the need for both mutations to elicit the phenotype characteristic of the fitA76 mutant (Ramalingam et al 1999; Sudha et al 2001; B Praveen Kamalakar and M H Muanvar, unpublished results). These results and the similarity in the organization and expression of fitA and fitB genes vis-à-vis pheS and pheT genes (Springer et al 1982) led to the proposal that fitA and fitB genes could be same as pheS and pheT and that the subunits of phenylalanyl tRNA synthetase could also function as selective transcription factors interacting perhaps with β subunit of RNA polymerase (Ramalingam et al 1999; Sudha et al 2001).

In this investigation, in an effort to know whether the selective transcription regulation by Fit factors stems from their interaction with the σ subunit of RNAP, we sought for mutation(s) in rpoD capable of suppressing the fitA76 Ts phenotype. Starting from a fitA76 mutant, we isolated a Ts⁺ derivative (JV4). This report describes the genetic and physiological characterization of this derivative.

2. Materials and methods

2.1 Media and chemicals

M9 minimal media and LB medium were prepared according to Miller (1972, 1992). Antibiotics were used at the following concentrations: kanamycin 30 μ g/ml; tetracycline 10 μ g/ml; rifampicin 50 μ g/ml. Rifampicin was obtained from Sigma Chemical Company, USA while all other antibiotics and other chemicals were from local sources. [³H] uridine was from Bhabha Atomic Research Centre, Mumbai. The ready-made scintillation fluid (INSTA-GEL) was from Packard Instrument Company Inc., USA.

2.2 Bacterial strains, and bacteriophages

The *E. coli* strains used in this study are listed in table 1. The P1 phage (P1 *vir*) used in this investigation is from our laboratory collection.

2.3 Methods

All genetic techniques were according to Miller (1972, 1992).

2.4 Curing of Tn10

This was done according to the method of Maloy and Nunn (1981). Fresh overnight cultures to be cured of Tn10 were sub cultured into fresh LB medium and allowed to grow till midlog phase. One ml of the cells were centrifuged down and resuspended in the same volume of saline. Approximately $10^5 - 10^6$ cells were plated on Bochner's medium and incubated at 37°C . The colonies which appeared were segregated twice on Bochner's medium and checked for the Tet's phenotype.

2.5 Pulse labelling of RNA and decay of pulse-labelled RNA

Cells were grown to midlog phase at 30°C with shaking in glucose minimal medium, centrifuged, resuspended in the original volume of 0.01 M Tris-HCl, pH 8 containing 1 mM disodium EDTA and aerated at 30°C for 10 min. The Tris EDTA treatment was given to allow rapid permeation of rifampicin that is used to arrest RNA synthesis after pulse labelling. The cells were again centrifuged down, resuspended in the original volume of pre-warmed growth medium, divided into two halves and shaken at 30°C and 42°C for 1 h. After pre-incubation, RNA was pulse labelled with [3 H] uridine (16,500 mCi/mmol; 0.5 μ Ci/ml) for 20 s. The pulse was terminated by adding rifampicin (100 μ g/ml) and non-radioactive uridine (10 mM). After this 0.2 ml aliquots

Table 1. List of *E. coli* strains used in this study.

Strain	Relavent genotype	Source/Reference
CSH57	F ⁻ arg his trp ilv metA leu purE rpsL	Laboratory collection
AB1157	F^- his $G4\Delta$ (gpt-pro) leuB6 thr1 argE3 rpsL31	Laboratory collection
JAJ572	F- fitA76 (pheS5 fit95) derivative of CSH57	Jabbar and Jayaraman 1978
JV4/Tn10	A Tet ^r Ts ⁺ derivative of JAJ572	This study
HfrC K10	HfrC wild type	Berlyn, CGSC, USA
K10 Tet ^R	pps::Tn10 derivative of HfrC	This study
CAG18578(K34)	zdj3124::Tn10kan derivative MG1655	Berlyn, CGSC, USA
RT500	F- his pps aroD pyrD edd pfkA? rpsL	Laboratory collection
JV4	Tet ^s derivative of JV4/Tn10	This study
JV41	pps::Tn10 fitA+ fitB+Ts (fitC4) derivative of JV4	This study
JV43	zdj 3124::Tn10kan fitC4 fitA+ fitB+ pps::Tn10 derivative of JV4	This study
JV57	aroD pps::Tn10 derivative of CSH57	This study
JVS5	pps::Tn10 pheS5 derivative of CSH57	This study
JV fitA76*	aroD ⁺ Ts (fitA76* fitC ⁺) derivative of JV57	This study
PMJfitA76*	zdj 3124::Tn10kan fitC+ fitA76* derivative of JV4	This study
PMJ02	pps::Tn10 fitA76* fitC ⁺ derivative of HfrC K10	This study
PMJ50	pps::Tn10 derivative of PMJ fitA76*	This study
SMJ01	Same as K10 but has pps::Tn10 fitA76 (pheS5-fit95)	Sudha et al 2001
NP37	Same as HfrC K10 but pheS5	Bachmann, CGSC, USA
SMJ02	Same as HfrCK10 but has pps::Tn10 pheS5	Sudha et al 2001
PKM01	Same as HfrCK10 but has pps:: Tn10 fitC4	This study
PKM01 pps ⁺	Same as PKM01 but pps ⁺ Tet ^S	This study
PKM57	pps::Tn10 fitC4 fitA+ fitB+derivative of CSH57	This study
PKM57 pps ⁺	Same as PKM57 but pps ⁺ Tet ^S	This study
PMJ95	Same as AB1157 but has fit95 pps::Tn10 rpoB201	This study

were removed at different time intervals, added to 0.5 ml of ice cold 10% TCA, and kept chilled. The precipitate was collected on Whatman glass microfibre filters, washed 5 times with 5% TCA containing 50 μ g/ml of non-radioactive uridine and once with 95% ethanol and air dried. The dried filters were counted in a liquid scintillation counter at an efficiency of 65%.

3. Results

3.1 Isolation of a temperature insensitive derivative from a fitA76 mutant

When the work reported in this paper was initiated it was neither known nor suspected that the fitA76 mutant could harbour two mutations (pheS5 and fit95), although its transcription defects at 42° C were well documented (Jayaraman and Jabbar 1980; Dass and Jayaraman 1985 a,b; Munavar et al 1993). We thought it would be of interest to see if a mutation in the rpoD gene (coding for the σ^{70} subunit) could suppress the Ts phenotype of the fitA76 mutant. Therefore, a derivative of E. coli C600 ($rpoD^+$) having a Tn10 insertion close to rpoD was mutagenized with MNNG and grown overnight in LB. Phage P1 propogated on the mutagenized culture was used to transduce Tn10 (Tet $^{\circ}$) to a fitA76 mutant

(JAJ572). The transduced cells were plated on LB-Tet medium and incubated at 30°C until the Tet^r transductants appeared as tiny colonies. Four plates containing approximately 2000 tiny Tet^r transductants in total were shifted to 42°C and incubated at that temperature for 24 h more. Of the four colonies, which grew bigger in size (see table 2, cross 1), one that grew well at 42°C in both LB and minimal medium was designated as JV4/Tn10 and used for further studies.

3.2 The suppressor mutation is not located in the rpoD region in JV4/Tn10

In order to verify whether the Ts phenotype of the *fitA76* is suppressed by mutation(s) in or near the *rpoD* locus, the Tn10 (linked to *rpoD*) from JV4/Tn10 was transduced again into a *fitA76* recipient and the Tet^r transductants were screened for Ts⁺ phenotype. None of the 230 Tet^r transductants checked was Ts⁺ implying the absence of a suppressor mutation in or near *rpoD* (cross 2, table 2). It is possible that while selecting for Ts⁺ derivatives we might have picked up either a true revertant or a colony harbouring a suppressor elsewhere on the chromosome. This is reminiscent of the observations of Dass and Jayaraman (1985a) who isolated *fitA24* as an intragenic suppressor of *fitA76* while attempting to isolate suppressor mutation(s) in *rpoB*. A similar

Table 2. Isolation of a Ts⁺ derivative (JV4) from a fitA76 mutant and demonstration of the presence of fitA76 in it.

Donor	Recipient	Selected marker/ character	Unselected phenotype	Cotransduction (%)
P1/ C600(has Tn10 near rpoD+) MNNG mutagenized	JAJ572(fitA76) (Ts)	Tet ^r	Ts ⁺	0.2 (4/2000)
P1/ JV4/ Tn10 (a Ts ⁺ colony from cross 1)	JAJ572(fitA76)	Tet ^r	Ts ⁺	< 0.40 (0/230)
P1/ JAJ572(fitA76)	RT500(aroD)	$aroD^+$	Ts	45.00 (46/103)
P1/ JV4/ Tn10	RT500(aroD)	$aroD^+$	Ts	40.00 (137/340)
P1/ JV4 (Tet ^s derivative of JV4/ Tn <i>10</i>)	RT500(aroD)	aroD ⁺	Ts	43.00 (71/165)

Ts, temperature sensitivity; Ts⁺, temperature insensitivity.

observation was made by Munavar and Jayaraman (1987), who isolated *fitB* as an extragenic suppressor of *fitA76* while trying to isolate suppressor mutation(s) in *gyrA*. Therefore, we looked for the retention of the *fitA76* lesion in JV4/Tn10.

3.3 The fitA76 lesion is intact in JV4/Tn10 and the suppressor mutation maps close to fitA76 to its left

The fitA76 lesion cotransduces approximately 50% and 60% with aroD and pps, respectively (Dass and Jayaraman 1985a; see also cross 3, table 2). When P1 propagated on JV4/Tn10 was used as the donor to transduce the aroD+ allele into RT500 (the same recipient used in cross 3), 40% of the aroD+ transductants became temperature sensitive (cross 4, table 2). This result showed the presence of a Ts lesion, possibly fitA76, in JV4/Tn10 near aroD and also implied that the suppressor mutation could lie elsewhere on the chromosome. Before proceeding to know the position of the suppressor mutation it was necessary to cure JV4/Tn10 of its Tn10 because such a Tets derivative would allow mapping of the suppressor mutation by replacing it with its wild type allele along with linked Tn10 or Tn5 insertions from the Carol Gross collection (Nicholas et al 1998). The curing was done as described under §2. The cured (Tets) derivative was designated JV4. The curing of Tn10 had no obvious effect on the phenotype of JV4 because it grew as well at 42°C as did JV4/Tn10. Also, when P1 propagated on JV4 was used to transduce the aroD⁺ marker into RT500, 40% of the aroD+ transductants became temperature sensitive, as expected (see table 2, crosses 4 and 5).

The previously isolated both intra and extragenic suppressors of fitA76 Ts, namely, fitA24 and fitB by themselves conferred a Ts phenotype. In order to know if this will be true in the present instance also the effect of introduction of the $fitA^+$ allele into JV4 was studied. P1 propagated on HfrC pps::Tn10 ($fitA^+$) was used to transduce $fitA^+$ via the linked

pps::Tn10 into JV4 and the Tetr transductants obtained were screened for growth at 42°C on LB medium. Since both the donor and recipient in this cross are Ts⁺ (the former being fitA+ is Ts+ and the latter being fitA76 harbouring a suppressor is phenotypically Ts⁺) replacement of fitA76 by fitA+ allele would not be expected to give temperature sensitive transductants in this cross, unless the suppressor mutation by itself confers temperature sensitivity in a fitA+ background. The fraction of temperature sensitive colonies (if obtained) among Tetr transductants should equal the cotransduction between pps and fitA (~60%), provided the locus of the suppressor and fitA are not linked. The fitA-pps cotransduction frequency using pps::Tn10 fitA+ as donor and an authentic fitA76 as recipient was observed to be approximately 66% (table 3, cross 1). However, when same donor P1 was used to transduce pps::Tn10 linked fitA+ to JV4 (Tet^S) only 34% of the Tet^r transductants became temperature sensitive (table 3, cross 2). This showed that the suppressor mutation is located in the vicinity of the fitA, possibly to its left (see below) and confers temperature sensitivity in fitA⁺ background. It could be linked to fitA since its cotransduction frequency with pps is only 34% as against the expected cotransduction frequency of 60%, if unlinked. This cross is schematically illustrated in figure 1A.

In order to verify these conclusions, we randomly picked up a temperature sensitive Tet^r transductant from the above cross and designated the same as JV41 with a presumptive genotype *sup-fitA*⁺-*pps*::Tn10 aroD⁺. When P1 propagated on JV41 was used to transduce the *aroD*⁺ marker into an *aroD fitA*⁺ recipient (RT500), 21% of the *aroD*⁺ transductants became Ts (cross 3, table 3). When the same P1 lysate was used to transduce *pps*::Tn10 in to the *pps*⁺*fit*⁺ strain CSH57, 40% of the Tet^r transductants became Ts (table 3, cross 4; figure 1B,C). The *aroD*-suppressor and *pps*-suppressor cotransduction frequencies (21% and 40% respectively) place the suppressor mutation approximately

Table 3. Location of the suppressor of *fitA76* near *fitA* in JV4.

Donor	Recipient	Selected marker/ character	Unselected phenotype	Cotransduction (%)
P1/HfrC fitA+ pps:: Tn10	JAJ572 (fitA76)	Tetr (pps::Tn10)	Ts^+	66.00 (71/107)
P1/HfrC fitA ⁺ pps:: Tn10	JV4 Tet ^s (derivative of JV4/Tn <i>I0</i>)	Tet ^r (pps::Tn10)	Ts	34.00 (77/230)
P1/JV41(a <i>pps</i> :: Tn10 Ts transductant of JV4 from cross 2)	RT500 (aroD fitA+)	$aroD^+$	Ts	21.0 (19/91)
P1/JV41	CSH57 (<i>fit</i> ⁺ <i>pps</i> ⁺)	Tet ^r (pps::Tn10)	Ts	40.0 (53/134)

Ts, temperature sensitivity; Ts⁺, temperature insensitivity.

0.83 and 0.61 min to the left of *aroD* and *pps* respectively. It is known that fitA is located 0.35 min to the left of aroD and 0.2 min to the left of pps (reviewed by Jayaraman 1994). Therefore, the suppressor should be located to the left of aroD, pps and fitA, approximately at 39.01 min on the E. coli linkage map (see below for details), leading to the following order of markers in the Ts⁺ derivative JV4 or JV4/Tn10: suppressor- fitA76 - pps - aroD counterclockwise on the E. coli genetic map (Berlyn 1998). However, it was shown earlier (see table 2) that JV4 harbours a Ts mutation, which cotransduces 40% with aroD possibly fitA76 itself (for the sake of clarity this Ts lesion will be called fitA76 until we show below that it is indeed a modified form of fitA76). After the position and phenotype of the suppressor were known it can be retrospectively inferred that the Ts transductants obtained in cross 5 table 2 should be the result of an event leading to the inheritance of only fitA76 (excluding the suppressor allele) along with selected marker $(aroD^+)$ as shown in the figure 1D.

3.4 Unmasking the latent temperature sensitivity of JV4 by the introduction of the wild type allele of the suppressor

The data presented so far shows that the relevant genotype of JV4 is: $sup^--fitA76-pps^+-aroD^+$. It was also shown that replacement of fitA76 by $fitA^+$ allele in JV4 also results in temperature sensitivity (see above). Therefore replacement of the mutant suppressor allele by the wild type allele in JV4 should also result in Ts phenotype. For this purpose the choice of the donor is very crucial. The donor should have a Tn10/Tn5 insertion located within transducible limits to the left of suppressor but beyond transducible limits from $fitA^+$. Such a donor could be used to co-transduce only the locus of the suppressor along with the Tn10/Tn5 insertion and exclude the cotransduction of $fitA^+$. Testing five different strains carrying Tn10::kan insertions located near the fit

3.5 The map position of the suppressor mutation in relation to other fit mutations and its implications

The map positions of fitA and fitB alleles relative to their neighbours, especially *aroD* and *pps*, have been worked out very well (reviewed by Jayaraman 1994). Now it is known that fitA is the same as pheS and fitB could possibly be pheT (Ramalingam et al 1999; B Praveen Kamalakar, M H Munavar and R Jayarman, unpublished results) fitA and fitB map at 38.7 and 38.6 min respectively, on the E. coli chromosome. The aroD-suppressor and pps-suppressor distances (0.83 and 0.61 min, respectively; see above) place the suppressor locus at 39.01-39.03 min. Also the type suppressor allele cotransduces zdj3124::Tn10 kan around ~ 4% (see above) and the mutant allele cotransduces with same insertion around 12% (see below). The average cotransduction (8.0%) places the suppressor 1.3 min away from the zdj3124::Tn10 kan insertion, at 39.01 min. The map position of the suppressor mutation indicates that it could be an extragenic suppressor. The Ts phenotype of the suppressor mutation implies that it defines an essential function. We have tentatively designated the locus defined by the suppressor mutation as fitC and the mutation will henceforth be referred to as fitC4. Figure 2 summarizes the position of fitC in relation to fitA and fitB, on the linkage map.

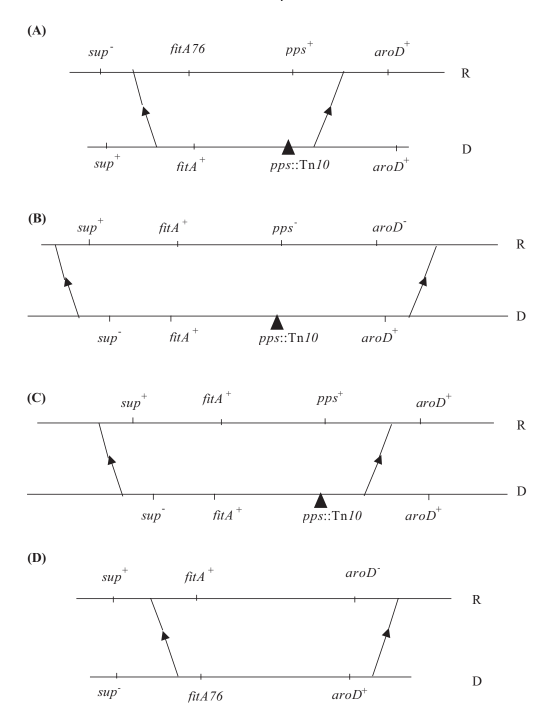


Figure 1. Schematic illustration of the transductional cross described in cross 2, table 3 (**A**), cross 3, table 3 (**B**), cross 4, table 3 (**C**) and cross 5, table 2 (**D**). (A) HfrC sup^+fitA^+pps ::Tn10 (D) x JV4 $sup^-fitA^76 pps^+$ (R). The two cross over events needed to generate the transductants of the type sup^-fitA^+pps ::Tn10, one to the right of the selected marker (pps::Tn10) and the other to its left between sup^+ and $fitA^+$ are shown. (**B**) JV41(sup^-fitA^+pps ::Tn10 $aroD^+$) (D) x RT500 (sup^+fitA^+pps - $aroD^-$) (R). The two cross over events needed to generate the transductants of the type $sup^-fitA^+aroD^+$ one to the right of the selected marker ($aroD^+$) and the other to its left, beyond sup^- are also shown. (**C**) JV41(sup^-fitA^+pps ::Tn10 $aroD^+$) (D) x CSH57 ($sup^+fitA^+pps^+aroD^+$) (R). The two cross over events needed to generate the transductants of the type sup^-fitA^+pps ::Tn10 one to the right of the selected marker (pps::Tn10) and the other to its left, beyond sup^- are also shown. (**D**) JV4 $sup^-fitA^+aroD^+$ (D) x RT500 sup^+fitA^+aroD (R). The two cross over events needed to generate the transductants of the type sup^+fitA^+6 $aroD^+$ one to the right of the selected marker ($aroD^+$) and the other to it's left between sup^- and $sup^-fitA^+aroD^+$ one to the right of the selected marker ($aroD^+$) and the other to it's left between sup^- and $sup^-fitA^+aroD^+$ one to the right of the selected marker ($aroD^+$) and the other to it's left between sup^- and $sup^-fitA^+aroD^+$ one to the right of the selected marker ($sup^-fitA^+aroD^+$) and the other to it's left between sup^- and $sup^-fitA^+aroD^+$ one to the right of the selected marker ($sup^-fitA^+aroD^+$) and the other to it's left between $sup^-fitA^+aroD^+$ are also shown (figure not drawn to scale). R, Recipient; D, donor

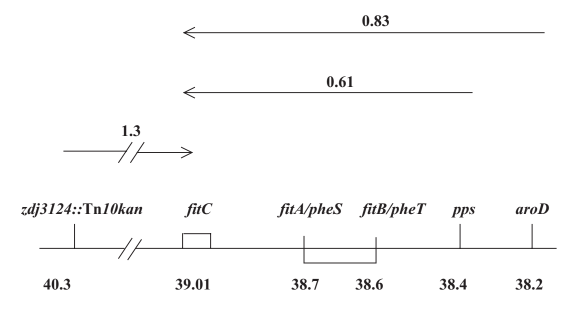


Figure 2. The map postion of *fitC* (in minutes) calculated from co-transduction frequencies using Wu's formula (Wu 1966). The position given for *zdj3124*::Tn*10kan* is from Nicholas *et al* (1998). The positions of *fitA* and *fitB* are from our previous publications (reviewed by Jayaraman 1994) and *pps aroD* positions are from Berlyn (1998).

3.6 Differential suppression of the Ts phenotype of fitA76 (pheS5 fit95) and pheS5 mutants by the fitC4 mutation

It was shown above that introduction of fitC⁺ allele into JV4 renders the latter Ts. The reciprocal experiment, namely, introduction of the fitC4 allele into fitA76 would be expected to render the latter Ts⁺ (reconstruction of JV4). Since fitA76 is now known to harbour two lesions pheS5 and fit95 (Ramalingam et al 1999) it would be interesting to look into the effect(s) of the fitC4 mutation on pheS5 and if possible with fit95 also. These experiments required a strain from which fitC4 could be mobilized into desired recipients. Such a strain of genotype zdj3124::Tn10kan fitC4 fitA+ fitB+ pps::Tn10 was constructed as follows. JV4 was transduced with P1 grown on a $fitA^+B^+C^+pps$::Tn10 and the TetR transductants screened for temperature sensitivity to select the transductants that retained fitC4 but received fitA+B+ (fitC4 fitA+B+ pps::Tn10). zdj 3124::Tn10kan was then introduced and Ts colonies (which retained *fitC4*) among the kan^R transductants were picked up. The final construct, zdj3124::Tn10kan fitC4 fitA+ fitB+ pps::Tn10 was named JV43. When P1/JV43 was used to transduce a fitA76 recipient (JAJ572) none (out of 160) of the kan^r transductants were Ts⁺ (table 4, cross 1) whereas one would expect approximately 4% to have become Ts⁺ (see § 3.5). However, 12% of the kan^r transductants obtained using a *pheS5* strain as recipient were Ts⁺ (table 4, cross 2). The inability of fitC4 to suppress the fitA76 lesion was totally unexpected since fitC4 was isolated as a suppressor of a fitA76 strain. Moreover a Ts lesion was shown to be present in JV4 at a site expected of fitA76 (see table 2, crosses 3–5). In order to clarify the discrepancy the ability of fitC4 to suppress the Ts lesion present at the position of fitA76 in JV4 was tested. As shown in table 4, crosses 3 and 4, fitC4 does suppress the same at the same frequency as it suppresses pheS5 (12%). The above data shows that the lesions present in original fitA76 and that present in the position of fitA76 in JV4 are not the same. The Ts phenotype due to former could not be suppressed by the fitC4 mutation whereas the latter (henceforth referred to fitA76*) could be.

3.7 Reconstruction of JV4 (fitC4 fitA76*) in another genetic background

As could be seen from the above the JV4 was isolated as a Ts⁺ derivative of JAJ572 (a CSH57 derivative; see table 1) during an attempt to isolate suppressor mutations(s) in *rpoD*. Genetic characterization of JV4 reported above clearly indicate that it neither harbours a suppressor of *fitA76* in/near *rpoD*, nor harbours the original *fitA76* lesion. Instead, it is shown to harbour a modified form of *fitA76* (*fitA76**) together with its suppressor *fitC4*. Since most observations made during the characterization of JV4 were unexpected, it was imperative to move these two mutations (*fitC4* and *fitA76**) into a totally new genetic background such as HfrC, and reconfirm that suppression occurs independent of genetic background. Since JV4 is a Ts⁺

Table 4. Transductional crosses illustrating the differential suppression of *fitA76* and *pheS5* Ts mutants by *fitC4*.

Donor	Recipient	Selected marker/ character	Unselected phenotype	Cotransduction (%)
P1/JV43(zdj 3124:: Tn10 kan fitC4 fitA ⁺ B ⁺ pps::Tn10)	JAJ572 (fitA76)	Kan ^r	Ts ⁺	< 0.6 (0/160)
P1/JV43(<i>zdj 3124</i> :: Tn <i>10 kan fitC4 fitA</i> ⁺ <i>B</i> ⁺ <i>pps</i> ::Tn <i>10</i>)	JV5S (pheS5)	Kan ^r	Ts ⁺	12 (10/85)
P1/JV4 (fitC4 ⁻ fitA76 ? aroD ⁺)	JV57 ($fitC^+$ $fitA^+B^+$ pps::Tn10 aroD)	$aroD^+$	Ts	40 (43/106)
P1/JV43(zdj3124::Tn10 kan fitC4 fitA+B+ pps::Tn10)	an <i>aroD</i> ⁺ -Ts transductant obtained in cross 3 (JV <i>fitA76</i> *)	Kan ^r	Ts ⁺	12 (15/125)

Ts⁺/Ts refers growth or no growth on LB plates at 42°C.

Table 5. Transductional crosses involved in the reconstruction of fitA76* fitC4 (Ts⁺) in HfrC and CSH57 backgrounds.

Donor	Recipient	Selected marker/ character	Unselected phenotype	Cotransduction (%)
CAG18578 (<i>zdj 3124</i> ::Tn <i>10</i> <i>kan fitC</i> ⁺ <i>fitA</i> ⁺ <i>fitB</i> ⁺)	JV4 (fitA76* and fitC4)	Kan ^R	Ts	3 (8/250) ^a
HfrC K10 (fitC+fitA+fitB+ pps::Tn10)	PMJ fitA76* (kan ^r fitC ⁺ , fitA76* from cross 1)	Tet ^R	Ts	16 (14/85) ^b
PMJ50(pps::Tn10 fitA76* fitC+kan ^r from cross 2)	HfrC K10	Tet ^R	Ts	50 (44/88) ^c
PMJ02 (fitC+fitA76*Ts pps:: Tn10 from cross 3)	PKM01 <i>pps</i> ⁺ (HfrC <i>fitC4</i> Ts <i>pps</i> ⁺ Tet ^S)	Tet ^R	Ts^+	~ 1(3/316)
PMJ02 (fitC+fitA76*Ts pps:: Tn10 from cross 3)	PKM57 pps ⁺ (CSH57fitC4 Ts pps ⁺ Tet ^S)	Tet ^R	Ts ⁺	~ 1 (2/208)

^a Ts transductant of this cross (kan^r fitC⁺ fitA76*) was named PMJfitA76*.

derivative of CSH57, the same experiment can also be done in CSH57. These were done as follows.

P1 prorogated on the strain K34 (zdj3124::Tn10kan; figure 2) was used to transduce the Kan^R marker into JV4. The Kan^R transductants that received $fitC^+$ but retained $fitA76^*$ would be Ts. It was observed that $\sim 3\%$ (8/250) of the Kan^R transductants exhibited Ts phenotype as could be expected of the cotransduction between zdj3124::Tn10kanR and fitC4 (see above and also cross 1, table 5). One of the Ts colonies (PMJ $fitA76^*$) was purified and was used as a recipient to transduce pps::Tn10 using P1/HfrC $fitC^+A^+B^+$ pps::Tn10 (K10 TetR, table 1). In this cross any transductant receiving only pps:: Tn10 but not $fitA^+$ will continue to have $fitA76^*$ thus would be Ts. As can be seen from the table 5 cross 2, 14 out of 85 TetR colonies retained the $fitA76^*$ allele

and were Ts at 42°C. One of the resultant Ts colonies was named PMJ50 (see table 5). When P1/PMJ50 was used to mobilize the *fitA76** allele into wild type HfrCK10 with linked *pps*::Tn10, 50% of the Tet^R transductants were Ts at 42°C as was expected (table 5 cross 3). One Ts (PMJ02), one Ts⁺ colony was purified and used for further studies.

First in order to make sure that the *fitC4* confers a Ts phenotype regardless of genetic background, P1/JV41 [pps::Tn10 fitB+ fitA+ fitC4 (Ts) derivative of JV4] was used to transduce fitC4 Ts with linked pps::Tn10 into HfrC, AB1157 and CSH57. In all the three genetic backgrounds around 40% of TetR colonies became Ts (data not shown) as expected of the cotransduction between *fitC4* and pps::Tn10. This confirms that the Ts phenotype of *fitC4* is

^b Ts transductant of this cross (fitC⁺ fitA76* pps::Tn10) was named PMJ50.

^c A Ts Transductant of this cross was named PMJ02.

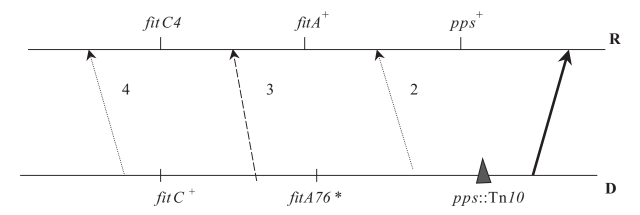


Figure 3. Schematic illustration of the transductional cross involved in the reconstruction of *fitA76* fitC4* and hence Ts⁺ strain (similar to JV4) using relevant donor and recipient strains, see crosses 4 and 5 of table 5. The relevant genotype of both the recipient stains (PKM01*pps*⁺ and PKM57 *pps*⁺) are *pps*⁺ *fitA*⁺ *fitC4* and genotype of the donor strain is *pps*::Tn*10 fitA76* fitC*⁺ (PMJ02). To generate the Ts⁺ (*fitC4 fitA76**) transductant, one crossover (1) should occur to the right of *pps*::Tn*10* locus (bold line) and the other second cross over should occur in between *fitC*⁺ and *fitA76** (3) (figure not drawn to scale). For other details see text.

not grossly affected by genetic background although there is some minor variation (B Praveen Kamalakar and MH Munavar; unpublished results). We chose one pps::Tn10 fitC4 Ts derivative of HfrC (named PKM01) and one pps::Tn10 fitC4 derivative of CSH57 (named PKM57) and intended to transduce fitA76* into both to make sure that in both cases the resultant fitC4 fitA76* transductants become Ts⁺ phenotype. However, the presence of pps::Tn10 in PKM01 and PKM57 rendered them unsuitable for use as recipients to introduce fitA76* along with pps::Tn10. Therefore it was imperative to eliminate pps::Tn10 from PKM01 and PKM57. This was done by transducing them to pps^+ by P1 propogated on a wild type $(pps^+fitA^+fitB^+fitC^+)$ strain. Among the pps⁺ (lactate⁺) transductants from both the crosses, one that retained *fitC4* and hence Ts were saved. To the pps⁺fitC4 derivatives of PKM01 (PKM01 pps⁺) and PKM57 (PKM57 pps⁺), we introduced fitA76* with linked pps::Tn10 using P1 made on PMJ02. Such a cross is schematically illustrated in figure 3. As could be seen from the figure in such crosses, transductants which received only pps::Tn10 (1:2 crossovers) will exhibit Ts phenotype due to fitC4 mutation. Also the transductants which received both fitA76* and fitC+ with pps::Tn10 will also become Ts due to fitA76* (1:4 crossovers). However, the transductants which receive only fitA76* with pps::Tn10 (1:3 crossovers) will have the genotype fitC4 fitA76* and thus should exhibit Ts⁺ phenotype due to suppression of each others' Ts phenotype. But a fraction of such colonies will be low because the second crossover should occur between fitC+ and fitA76* (figure 3). As was expected, about 1-2% of the TetR transductants in both crosses became Ts+ (see table 5). These results confirm that the introduction of fitC4 and fitA76* in any genetic background confers Ts⁺ phenotype.

3.8 fit95 is intact in JV4

Considering the fact that fitA76 has two mutations (pheS5 and fit95; Ramalingam et al 1999), we speculated that the fitA76* in JV4 might have arisen due to a change either at the pheS5 position or at the fit95 position. Initially we wanted to check whether fit95 is intact in JV4. It is known that fit95 when present alone confers a Ts phenotype on rich media devoid of NaCl (referred as - salt Ts phenotype, Ramalingam et al 1999). In the fitA76 mutant the order of markers is pheS5-fit95-pps-aroD, counter-clockwise on the E. coli chromosome. The fit95 and pheS5 could be separated at a frequency of approximately 1-2% in transductional crosses (Ramalingam et al 1999 see also cross 1, table 6). If JV4 harbours intact fit95 we could expect that in a cross P1/JV4 X aroD fitA+ recipient like JV57, 1-2% of the aroD⁺ transductants should be fitA⁺ fit95-aroD⁺ and hence Ts on – salt media. As can be seen from the data presented in table 6, cross 2, approximately 2% of the aroD+ transductants were Ts on LB-NaCl media. These results confirm the presence of the fit95 mutation in JV4. Therefore, the modified form of fitA76 (fitA76*) present in JV4 might have acquired a change at the position of pheS5. Sequence analysis has confirmed this conclusion (also see §4).

3.9 Evidence that fit95 could also suppress the fitC4 Ts phenotype

When we transduced a *fitA76* (*pheS5 fit95*) recipient for Kan^R using P1 grown on *zdj3124*::Tn*10kan fitC4* strain (JV43) none of the Kan^R transductants became Ts⁺ (see table 4 cross 1; this cross is also shown as cross 1 in table 7 for the sake of clarity). However, in a reciprocal cross (in

Table 6. Transductional crosses showing the presence of the *fit95* mutation in the Ts⁺ derivative JV4.

Donor	Recipient	Selected marker/ character	Transductants exhibiting the –salt Ts ^a (fit95) phenotype (%)
P1/HfrC <i>fitA76</i> (pheS5 fit95 pps::Tn10)	AB1157 (<i>pps</i> + <i>fît</i> +)	Tet ^r	1 (10/735)
P1/JV4 fitC4fitA76* aroD+	JV57 (aroD pps:: Tn10 fit+)	$aroD^+$	2 (6/285)

^a –Salt Ts refers to temperature sensitivity on rich media devoid of sodium chloride at 42°C. For cross 1 data reproduced from Ramalingam *et al* (1999).

Table 7. Evidence for suppression of *fitC4* Ts phenotype by *fit95*.

Donor	Recipient	Selected marker/ character	Unselected phenotype	Cotransduction (%)
P1/JV43 (<i>zdj 3124</i> ::Tn <i>10</i> <i>kan fitC4 fitA</i> ⁺ <i>fitB</i> ⁺ <i>pps</i> ::Tn <i>10</i>)	JAJ572 fitA76 (pheS5-fit95)	Kan ^R	Ts ⁺	< 0.6 (0/160)
P1/JAJ572 (fitA76 pps+)	JV43(zdj 3124:: Tn10kan fitC4 fitA ⁺ fitB ⁺ pps::Tn10)	pps^+	Ts ⁺	5 (10/202)
P1/JAJ572 (fitA76 pps+)	RT500(pps ⁻)	pps^+	Ts	70 (148/211)

which donor and recipient of cross 1 were interchanged) 5% of the pps⁺ transductants became Ts⁺ (see table 7, cross 2). These results are unexpected due to the following reasons. If the fitA76 Ts phenotype could be suppressed by the fitC4 mutation, then 60-70% of the pps⁺ transductants in cross should have become Ts⁺ because it is known that the fitA76 cotransduces with pps around 60-70% (Jabbar and Jayaraman 1978; Dass and Jayaraman 1985a; see also cross 3 of table 7). On the other hand if the fitA76 Ts phenotype could not be suppressed by the fitC4 mutation then none of the transductants should have become Ts⁺. Therefore, 5% Ts⁺ transductants obtained in cross 2 could be explained by postulating that the fit95 mutation present in the fitA76 (between pheS5 and pps), could suppress the fitC4 Ts phenotype. Figure 4 represents schematic illustration of the transductional cross 2 of table 7. The predicted genotypes and phenotypes of various possible classes of pps⁺ transductants are shown in table 8. As can be seen from table 8 all the pps⁺ transductants that arise due to second crossovers in all possible intervals other than between fit95 and pheS5 should be temperature sensitive. However, a second crossover event between fit95 and pheS5 would yield a class of transductants with genotype fitC4- pheS+ fit95 whose phenotype could not be predicted. Also the frequency of generation of such transductants would be much low (see above). Therefore, the 5% Ts⁺ transductants obtained in the above cross could be of the type fitC4 pheS⁺ fit95 pps⁺ only. These results not only reconfirm that the original fitA76 mutant indeed harbours a second mutation, fit95, in addition to the pheS5 but also clearly indicates that this mutation

suppresses the Ts phenotype due to fitC4.

We have reconfirmed that *fit95* indeed suppresses *fitC4* Ts phenotype by transducing *fit95* along with linked *pps*::Tn10 using P1 made on a *fit95 pps*::Tn10 strain (PMJ95) into the recipient *pps*⁺ *rpoB201*, a derivative of PKM01(harbouring *fitC4*). As was expected the transductants which received both *fitC4* and *fit95* mutations exhibited Ts⁺ phenotype regardless of media (data not shown) confirming the above results. In this cross both donor and recipient strains carry the *rpoB201* mutation known to stabilize the *fit95* – salt Ts phenotype (Ramalingam *et al* 1999; Rukmani 1996).

3.10 Kinetics of decay of pulse labelled RNA in strains bearing fitC4 and the modified form of fitA76 (fitA76*): Evidence that both mutants behave like the original fitA76 mutant

Upon a temperature shift from 30°C to 42°C in isogenic *fitA76* and *pheS5* mutants, it was observed that RNA synthesis was inhibited earlier and faster than protein synthesis in *fitA76* mutant and the reverse was seen in the *pheS5* mutant. This strongly suggested that the *fitA76* mutant is primarily transcription defective and *pheS5* mutant is primarily translation defective (Ramalingam *et al* 1999). Kinetics of decay of pulse labelled RNA in the *pheS5* and *fitA76* mutants showed the following. In the *pheS5* mutant, the pattern of decay of RNA pulse labelled at 30°C and 42°C was found to be triphasic (similar to that found with the wild type strain) consisting of fast decaying and slow

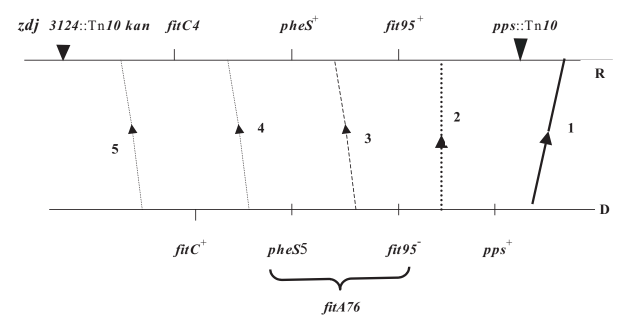


Figure 4. Schematic illustration of the transductional cross 2 shown in table 7. JAJ572 (*fitC*⁺ *pheS5 fit95*⁻ *pps*⁺) (D); X JV43 (*zdj3124*::Tn*10 kan fitC4 pheS*⁺ *fit95*⁺ *pps*::Tn*10*) (R). The first cross over is shown as a solid line and the second crossover as broken lines. It should be noted that only the second cross over occurring between *pheS5* and *fit95*⁻ (1:3) would yield transductants of the type *fitC4 pheS*⁺ *fit95*⁻. All the other second cross overs will yield only Ts transductants (see text and table 8).

Table 8. Predicted genotypes of the various classes of *pps*⁺ transductants in the transductional cross-illustrated in figure 4. JAJ572 (*fitC*⁺ *pheS5-fit95* (*fitA76*) *pps*⁺ (donor); X JV43 (*zdj 3124::*Tn*10 kan fitC4 pheS*⁺ *fit95* + *pps*::Tn*10* (recipient).

Interval of the second cross over	Genotype of the <i>pps</i> ⁺ transductants	Predicted phenotype of the <i>pps</i> ⁺ transductants
pps ⁺ -fit95 ⁻ (2)	fitC4 pheS5+ fit95+ pps+	Ts ^a
fit95-pheS5 (3)	fitC4 pheS5+ fit95- pps+	?
$pheS5$ -fitC $^+$ (4)	fitC4 pheS5 fit95 (fitA76) pps+	Ts^b
Beyond $fitC^+$ to its left (5)	fitC+ pheS5 fit95- pps+	Ts^c

The numbers shown in brackets refer to the second cross over intervals.

decaying species (representing mRNAs) and stable species (representing rRNAs and tRNAs) implying that all the three species of RNAs are made in this mutant at both temperatures. In the *fitA76* mutant RNA pulse labelled at 30°C decayed triphasically like that observed in a wild type strain. However, RNA pulse labelled at 42°C decayed biphasically comprising of only fast and slow decaying species (Sudha *et al* 2001); the stable species could not be detected at all in the *fitA76* mutant at 42°C. This together with earlier results of Jayaraman and Jabbar (1980) indicated that the absence of detectable stable RNA species in the *fitA76* mutant at 42°C could be due to lack of protection of nascent stable RNA at 42°C which in turn could be due

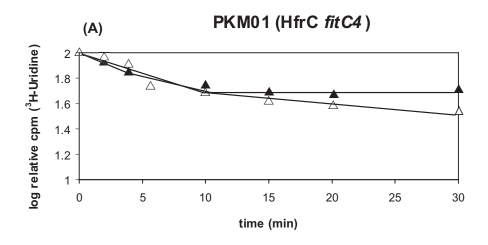
to lack of expression (transcription) of genes coding for (at least some) ribosomal proteins required to stabilize nascent ribosomal RNA from decay. Based on this, it was postulated that Fit factors function as selective transcription factors regulating the expression of few classes of genes; the genes coding for ribosomal proteins could be among them (Jayaraman and Jabbar 1980; Munavar *et al* 1993; Ramalingam *et al* 1999; Sudha *et al* 2001). Thus absence of detectable stable RNA synthesized at 42°C is one of the characteristic properties of the *fit476* mutant.

In order to know whether *fitC4* mutant and also the strain bearing the modified form of *fitA76* (*fitA76**) behave like the original *fitA76* mutant or like the *pheS5* mutant, we studied the

a Ts because these transductants are same as the recipient (fitC4) but pps⁺.

^b Ts because fitC4 mutation cannot suppress the Ts phenotype of the original fitA76 (pheS5-fit95) mutant.

^c Ts because these transductants are same as fitA76 but pps⁺.



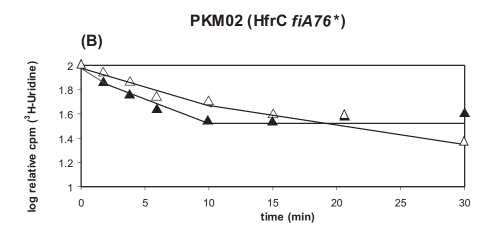


Figure 5. (A) Kinetics of decay of pulse-labelled RNA in HfrC fitC4 mutant (PKM01) at $30^{\circ}C(\blacktriangle)$ and $42^{\circ}C(\triangle)$. The initial CPM is taken as 100% and the values at other time points are normalized with respect to it. Each point in the graph indicates an average of three experiments. See text for other details. (B) Kinetics of decay of pulse-labelled RNA in HfrC $fitA76^*$ mutant (PKM02) at $30^{\circ}C(\blacktriangle)$ and $42^{\circ}C(\triangle)$. The initial CPM is taken as 100% and the values at other time points are normalized with respect to it. Each point in the graph indicates an average of three experiments. See text for other details.

kinetics of decay of pulse-labelled RNA in both. The relevant strains [PKM01 and PKM02 (PKM02 is genotypically same as PMJ02)] were grown in M9 medium and divided into two portions. One was kept at 30°C and the other was shifted to 42°C and incubated for 1 h. The two mutants at both temperatures were pulse-labelled for 20 s with ³H-uridine and pattern of decay was followed (see § 2.3 for details). We found the pattern of decay of RNA pulse-labelled at 42°C to be biphasic, and triphasic when pulse-labelled at 30°C. In both the mutants stable RNA species could not be detected when pulse-labeled at 42°C as the case with original *fitA76* (*pheS5 fit95*) mutant (see figure 5). For comparison the patterns of decay of pulse-labelled RNA in the original *fitA76* and *pheS5* mutants are also given (figure 6).

4. Discussion

Our earlier work provided genetic, physiological and biochemical evidence for the involvement of the fitA and fitB gene products in selective regulation of transcription in $E.\ coli$. It was originally believed that fitA and fitB were unidentified genes and their products interact with each other as well as with RNAP and regulate the expression of few classes of genes (Jabbar and Jayaraman 1978; Dass and Jayaraman 1985a,b, 1987; Munavar and Jayaraman 1987; Jayaraman 1994). It has been shown recently (Ramalingam $et\ al\ 1999$) that fitA is same as pheS, coding for the α subunit of phenylalanyl-tRNA synthetase (PheRS). There is suggestive evidence (genetic mapping, polarity of transposon

fitA76 vs pheS5 log relative cpm (H-Uridine) 1.8 1.6 1.4 1.2 1 0 5 10 15 20 25 30 35 40 45 50 55 60 time (min)

Figure 6. Kinetics of decay of pulse-labelled RNA in *fitA76* and *pheS5* mutants at 30°C (\triangle , \bullet) and 42°C (\triangle , \circ) (adapted from Sudha *et al* 2001).

insertions in fitA on fitB expression and complementation with Kohara phages) to show that fitB could be pheT, coding for the β subunit of PheRS (Munavar 1991; B Praveen Kamalakar and M H Munavar, unpublished results). Taken together it was proposed that fit function is a second to function associated with phenylalanine-tRNA synthetase, the subunits of which also function as selective transcription factors possibly by interacting with the β subunit of RNA polymerase (Ramalingam et al 1999; Sudha et al 2001).

In the present investigation, started at a time when the molecular details of the lesions present in the fit A76 mutant were not known, we attempted to isolate a suppressor of fitA76 mapping in rpoD. Starting from the fitA76 mutant we isolated a Ts⁺ derivative (JV4). We have shown here that JV4 neither harbours a suppressor of fitA76 mapping in/near rpoD nor is the original fitA76 lesion (pheS5 fit95) is intact in JV4. It is shown here that JV4 harbours a modified form of fitA76 (referred to as fitA76*) together with its suppressor. The suppressor mutation by itself confers a Ts phenotype and maps around 39.01 min. The map position of the suppressor clearly indicates that it is extragenic. We have tentatively designated this locus as fitC and the mutation as fitC4. While fitC4 could suppress the Ts phenotype due to fitA76*, pheS5 and fit95 mutants, it failed to suppress the Ts phenotype of original fitA76 (pheS5-fit95) mutant. The presence of the fit95 mutation in JV4 implies that some modification could have occurred at the position of *pheS5* to generate *fitA76**. This postulate has been verified to be true. The new *pheS* mutation present in JV4 (named pheS4) has been cloned, sequenced and shown to be a G₂₉₃-C₂₉₃ transversion (S Vidya, B Praveen Kamalakar, M H Munavar and R Jayaraman, unpublished results).

Extensive molecular analysis revealed that pheS4 could suppress fitA76 when present on a multicopy plasmid (S Vidya, B Praveen Kamalakar, M H Munavar and R Jayaraman, unpublished results). The inability of fitC4 to suppress the Ts phenotype of the transcription defective fitA76 mutant but its ability to do so in a translation defective pheS5 mutant raises the question whehter fitC4 is primarily transcription defective (like fitA76) or translation defective (like pheS5). The same question may be extended to fitA76* also. Results on the kinetics of decay of pulse labelled RNA in isogenic fitC4 and fitA76* mutants reported herein clearly indicates that both the mutations behave like the original fit A76 mutant. Moreover, the phenotype characteristic of fitA76 mutant viz the escape of phage T7 growth at 42°C after a brief period of incubation of infected cells at 30°C (Jabbar and Jayaraman 1976, 1978), has been found to be true of both fitC4 and fitA76* mutants (Saini 1997). Based on the above we propose that both fitC4 and fitA76* mutants are primarily transcription defective. Measurement of gross RNA synthesis at 30°C and 42°C and the pattern of inhibition of RNA and protein synthesis after a shift to 42°C would help to know the actual extent of transcription defect in these mutants. It would be worthwhile to know how the fitC4 mutation, which confers a primary defect in transcription, suppresses the phenotype of translation defective pheS5 mutant. Similarly it would be interesting to study the effect of fitC4 mutation in fitA24 and fitB genetic backgrounds. Such studies are currently underway.

Although, based on map position we might postulate that the *fitC4* mutation defines a new unidentified locus (*fitC*), it remains to be clarified whether this is true or *fitC4* is an allele of a known gene present at that position. However, the suppression properties of the *fitC4* mutation imply that the

product coded by it directly or indirectly interacts with the fitA/pheS and/or fitB/pheT products. How far would such interaction influences the transcription specificity of FitAB complex? Does the fitC gene product by itself interact with subunits of RNA polymerase? We believe that molecular cloning of fitC gene, identifying the fitC4 lesion and isolation of suppressor(s) of fitC4 mapping in the locus/loci coding for subunit(s) of RNAP would help elucidate the actual mode of functioning of fitC product. These are some facets of our current efforts.

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