

Caf1R gene and its role in the regulation of capsule formation of *Y. pestis*

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A new transcription unit of the *fl* gene cluster was found. The DNA sequencing revealed one long open reading frame. Deletion and frame shift mutation analyses have demonstrated the importance of a corresponding gene product for the F1 antigen biosynthesis. A homology of the deduced amino acid sequence with that of AraC family DNA-binding regulators was shown. A potential regulatory DNA region is discussed.

Gene regulation; Capsule antigen; Nucleotide sequence; *Y. pestis*

1. INTRODUCTION

Previously we have shown that at least three gene products are required for the formation of *Y. pestis* capsule: the *caf1* gene product (F1 antigen), which is a major capsule compound [1], the *caf1M* gene product, which plays a chaperone-like function during transport of the F1 subunit [2], and the *caf1A* gene product, which is important for anchoring of the capsule to the cell surface [3]. The genes are assembled into two transcription units oriented in the same direction: one for the *caf1* gene and the other for the *caf1M* and *caf1A* genes. Later another transcription unit, lying in the opposite orientation to these transcription units, was found. In the present study we show that this region contains a gene which is very important for the regulation of capsule biogenesis.

2. MATERIALS AND METHODS

The recombinant plasmids, containing the whole *fl* operon (pFS2) or its fragment (p12R) have been described previously [1,2]. The plasmids, pFSRM7 and pO6, were obtained by cloning of 1,110 and 1,054 bp fragments of pFS2 into pUC19. DNA cloning and sequencing by the Maxam–Gilbert and Sanger–Coulson methods were performed essentially as described by Maniatis et al. [4]. DNA and protein sequence analyses were made using the GENPRO software package from Riverside Scientific Enterprises and PCGENE package from Genofit. Anti-plague agglutinating antiserum and erythrocytes conjugated with polyclonal anti-F1 antibodies were obtained from the Anti-Plague Institute 'Microb' (Saratov, Russia). The internal primers were obtained from Immune-Biotech (Lyubuchany, Russia).

3. RESULTS AND DISCUSSION

The only long open reading frame was found 1.6 kb upstream from the *caf1M* gene (Fig. 1). A potential gene

product, designated Caf1R, was found to be 301 amino acid residues long with $M_r = 36,052$. A PIR'89 sequence database search revealed a homology between Caf1R and proteins of the AraC family (see references in [5]) (Fig. 2) which are known to be DNA-binding transcription regulators. These proteins are proposed to consist of helix-turn-helix (or homeodomain in eukaryotic analogs) regions containing positively charged amino acid residues. Recently the crystal structure of a homeodomain region of the Mata2–DNA complex has been determined [6]. As shown in Fig. 2, the Mata2 homeodomain displays about 20% primary structure homology with the corresponding region of VirF-Y., with 8 of 12 conserved hydrophobic positions in the AraC family proteins coincident with those of the Mata2 homeodomain, and the sole hyperconserved Arg residue of the AraC proteins coincident with Arg of the Mata2 protein. This Arg is a conserved residue in homeodomains of eukaryotic regulatory proteins, and is shown to be a component of the third homeodomain helix directly interacting with a large DNA groove [6].

The Caf1R protein seems to be highly basic, like other DNA-binding proteins. An isoelectric point of Caf1R was calculated to be 9.5.

A regulatory area of AraC was shown to be located in the C-terminal region of the protein [7]. We found that the plasmid, p12R, codes for a truncated form of Caf1R containing only 81 N-terminal amino acid residues with an additional tail of 19 amino acids encoded by the vector DNA (Figs. 1 and 3). A predicted isoelectric point of this Caf1R derivative was found to be 10.5.

Bacterial cells harboring p12R can produce very large amounts of the F1 antigen and are able to form a capsule. In order to prove that Caf1R and its N-terminal peptide possess positive regulatory activity we have introduced a frame-shift mutation into the plasmids, pFS2 and p12R, by digestion of DNA with *Xho*I, followed by T7 DNA polymerase filling-in of cohesive

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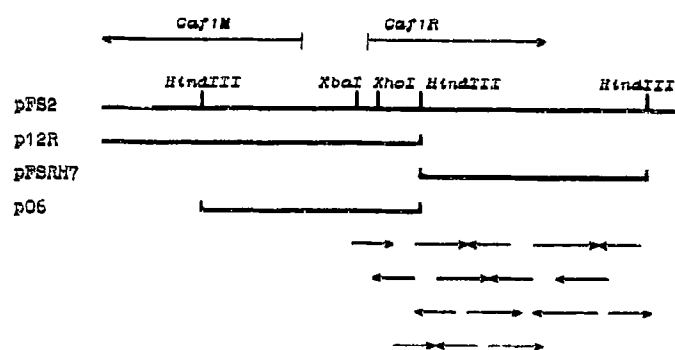


Fig. 1. Restriction map and sequencing strategy of the *cafIR* gene region. Bold bars indicate the cloned fragments. Arrows above the restriction sites indicate the open reading frames. Short arrows below indicate lengths and directions of the sequences determined.

ends and blunt-end ligating. The *E. coli* HB101 clones, containing plasmids lacking *Xho*I sites, failed to produce a detectable amount of the F1 antigen at 37°C. This frame-shift mutation leads to a truncated protein containing only the 18 N-terminal amino acids of CafIR. These data clearly demonstrate that the regulatory region is included in the N-terminal part of CafIR 81 amino acid-long region.

We can speculate about a possible Caf1R-determined regulation mechanism involved in the capsule formation, proceeding from the data on the other positive regulators sharing homology with Caf1R. The regulatory regions were shown to be located in the closest vicinity to the genes encoding such regulators. Therefore, the most probable regulatory region affected by

[illegible]

Fig. 2. Amino acid sequence alignment of the Caf1R protein with the AraC and Matα2 protein. Numbers indicate positions in the corresponding proteins. A possibility of random coincidence with the Caf1R is indicated against each sequence of the AraC proteins [11]. The predicted alpha-helix and beta-structure content is also indicated [12]. The hyperconserved hydrophobic and hydrophilic residues are indicated by asterisks and circles, respectively. Conserved hydrophobic residues are marked with crosses. The amino acid residues identical to those of Caf1R are indicated by capital letters. The identical and conserved residues in VirF-Y. and Matα2 sequences are indicated by colons and dots, respectively. Alpha-helical regions in the homeodomain of Matα2, revealed by the X-ray crystal analysis [6], are indicated by open boxes.

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verted repeats in the *cafIM* promoter region which might also be necessary for the regulation. We cannot exclude the possibility that proteins other than CafIR may be involved in the regulation process. An involvement of a histone-like protein in the transcription regulation of two divergent promoters was shown in a *pap* gene cluster [8]. The *pap* regulatory region was shown to contain a pair of inverted repeats like the *cafIR-cafIM* intergenic region. Finally, it should be noted that the synthesis of the CafIR protein is a prerequisite for F1 antigen production and hence formation of *Y. pestis* capsule. The CafIR protein seems to induce the *cafIM* gene transcription leading to the production of CafIM and CafIA proteins involved in capsule formation. The CafIR protein may be involved in signal transduction. Such signals may be temperature [9] and/or other environmental factors, such as Ca^{2+} and Mg^{2+} ions concentration [10]. Additional experiments should be carried out to elucidate the role of the C-terminal part of the CafIR protein.

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