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Identification and characterisation of behavioural genes of
Agrobacterium tumefaciens.

By

Sayyed Hassan Marashi

A thesis submitted to the Department of Biological Sciences
University of Durham

In accordance with the requirement for the degree of

Doctor of Philosophy

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Identification and characterisation of behavioural genes of *Agrobacterium tumefaciens*.

Sayyed Hassan Marashi

Abstract.

Three behavioural mutants (*fla*-8, *mot*-6 and *cheL*) generated by transposon Tn5 mutagenesis and localised on cosmid pDUB1905 were studied. The cosmid pDUB1905, from a representative genomic library of the *Agrobacterium tumefaciens* C58C1 chromosome has previously been partly mapped and found to contain genes concerned with flagella. In this study a region of 5860 nucleotides from a 12 kb *BamH1* fragment of cosmid pDUB1905 was sequenced completely in both directions. Homology searches of this sequence with sequence databases and other computer analysis revealed two flagellar-related genes (*flhA* and *fliR*), a chemotactic gene (*cheL*) and four open reading frames (*orfX*, *orfW*, *orfY* and *orfZ*) with no significant sequence identity to any open reading frame in databases. A putative promoter-like sequence was also found upstream of *orfZ*. The FlhA and FliR are the inner members of type III flagellum-specific export apparatus which are responsible for delivering the flagellar subunits lacking a signal peptide leader to the surface of the cell. These have counterparts in the type III secretion proteins system responsible for transporting pathogen proteins to host cells. The function of CheL has not yet been identified. Three ORFs have chaperone characteristics.

A mutant was created by insertion of a neomycin resistance cassette in the *fliR* homologue to determine the effects of the gene on motility. Phenotype analysis of the mutant showed no flagella and motility with small swarming pattern comparing to wild type, indicating that *fliR* is indeed a flagellar gene.

In this study two more members of flagellum-specific export, a chemotactic gene, three open reading frames which could have specific chaperon activity, and an unknown open reading frame were identified in *A. tumefaciens* C58C1.
TO:

My mother and Father,

Wife, Hoda,

Son Hossien,

Sister and Brothers

For their love and encouragement
Declaration

I declare that the work contained within this thesis submitted by me for degree of Doctor of Philosophy is my own original work, except where otherwise stated, and has not been submitted previously for a degree at this or any other University.

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Abbreviations.

Amp = ampicillin
ATP = adenosine triphosphate
Bp = base pairs
BSA = bovine serum albumin
CAMP = cyclic adenosine monophosphate
Cm = chloramphenicol
DMF = N,N-dimethyl formamide
dNTPs = deoxyribonucleoside triphosphates
DTT = dithiothreitol
EDTA = ethylenediaminetetraacetic acid
Gm = gentamycin
IPTG = isopropyl-β-D-thiogalactopyranoside
Kan = kanamycin
Kb = kilobase pairs
KDa = kilodaltons
MOPS = 3-(A^-morpholino)propane sulphonic acid
Neo = neomycin
OD = optical density
PEG = polyethylene glycol
p.s.i = pounds per square inch
Rif = rifampicin
RNAase = ribonuclease
SDS = sodium dodecyl sulphate
Strep = streptomycin
Tc = tetracycline
Tris = tris(hydroxymethyl)aminomethane
UV = ultraviolet
wrt = with respect to
X-Gal = 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside
5' = 5' terminal phosphate of DNA molecule
3' = 3' terminal hydroxyl of DNA molecule
1. Introduction
1.1 *Agrobacterium tumefaciens*: Characteristics and Classification

By any means *Agrobacterium tumefaciens* is one of the most successful organisms. It is one of the dominant bacterial species in the rhizosphere and it is capable of colonising and pathogenising plant hosts, causing epidemics of crown gall disease. Agrobacteria appear to have a great advantage in their capability for pathogenicity; making them one of the best natural vectors for introducing foreign genes into higher plants.

The most important feature of the pathobiology of this organism is its wide host range. *Agrobacterium tumefaciens* is a pathogen that can infect over 300 genera of dicotyledonous plants, at least four families of monocots and over 40 species of gymnosperms (54, 65).

Agrobacteria morphologically are aerobic, Gram-negative, short, encapsulated, rods, and usually 0.6-1.0 \( \mu m \) wide by 1.5-3.0 \( \mu m \) long. They are motile and have an unusual flagellation pattern with a polar tuft of two flagella and 2-4 lateral flagella (see fig 1.1) (97). They produce copious extracellular polysaccharide when grown on carbohydrate-containing medium. Frequently, they produce cellulose fibrils which are thinner and longer than their flagella. The colonies of Agrobacteria are non-pigmented and smooth. The optimum temperature for growth of *Agrobacterium tumefaciens* is 28°C.

*Agrobacterium* together with *Sinorhizobium* (formerly *Rhizobium*), *Rhizobium*, *Rhodobacter*, *Caulobacter* and *Azospirillium* are a member of the \( \alpha \)-subgroup of the class Proteobacteria. The family Rhizobiaceae contains soil bacteria which associate with plant cells and carry large plasmids.

The classifications of the genus *Agrobacterium* is divided into four species based on their phytopathogenicity (147). The pathogenic species are: *A. tumefaciens* which causes crown galls, *A. rhizogenes* which produces hairy root, *A. rubi*, the agent inducing cane galls on rubus species and *A. radiobacter* which is non-pathogenic. Each species is further divided upon the production of specific chemicals by the galls or infected roots called opines. This method of classification has been shown to be unrealistic since the phytopathogenicity of species is based on the production of opines. The genes coding for opine synthesis are on a large tumour-inducing (Ti) or
Figure 0.1.1.1 Electron micrograph of a wild type *A. tumefaciens* C58C*\(^1\)* cell showing flagella. Photograph taken by Dr. C. H. Shaw.

root inducing (Ri) plasmid that can be transferred easily from pathogenic strains to non-pathogenic strains resulting in the recipient strains acquiring the pathogenic properties of the donor strains. Therefore, phytopathogenicity is not suitable for species identification. In addition, it has been shown that all known Ti plasmid contain a variety of combination of opine catabolic genes and they can catabolise more than one opine (236).

An alternative system to classify the species has recently been established using phenotypic, physiological and genotypical differences, chromosomal characteristics, comparisons of protein patterns, and other tests such as growth at different temperatures and the sensitivity to sodium chloride (99 , 100). In this system of nomenclature, the genus *Agrobacterium* is divided into three biovars (biotypes) 1, 2, and 3. The three biovars are genetically and phenotypically different from each other and they contain both pathogenic and non-pathogenic strains. The strains of biovars can clearly be separated based on differences in 16s rRNA sequences. This
method of classification is more stable and valid for species identification than phytopathogenicity.

Members of the genus all possess two chromosomes, many of which demonstrate different topologies such as having linear and circular chromosomes (96). Each biovar has also a specific genome size.

Based on the kind of opines produced in the galls, Agrobacteria are also classified as octopine, nopaline, and leucinopine strains.

Despite of the new system of classification, the old nomenclature is still used and will be used here. The strains used in this project are derivatives of *A. tumefaciens* C58 which possesses four replicons: one circular and one linear chromosome (which is smaller) and two plasmids. This strain contains the Ti plasmid pTiC58.

1.1.2 Soil Ecology

*A. tumefaciens* can be isolated from both cultivated and uncultivated soils around the world and especially found in association with the rhizosphere. The numbers of bacteria present around the roots are vastly greater than those found in nearby soil (148). Pathogenic strains are usually only isolated from galls and soils surrounding the infected roots. Generally the populations of avirulent bacteria are much higher than the virulent strains in the soil (98). Since the Ti plasmid, the means of pathogenicity, is easily lost without selection, however many reports have shown that certain factors can affect the proportion between virulent and avirulent strains. In an experiment carried out to isolate *Agrobacterium* strains from a field with loamy soil that had not been cultivated for five years, the virulent number of Agrobacteria were much higher than expected (31). It was proposed that the loamy soil possibly increased the number of pathogenic strains according to its capacity for water retention and acid pH. The presence of weeds could also have a positive effect on virulent strain populations. Data from experiments on other related bacteria suggests that the soil type affects survival of the bacterial strains, the stability of plasmids, motility, growth and pathogenicity (31, 91). The number of virulent strains also can be affected by opines, which can be used by *Agrobacterium* but not by the host plant or majority of other microorganism. This capability of *Agrobacterium* to utilise
opines provides an environment favouring growth of the pathogen and transfer of the Ti plasmid. This hypothesis, is called the opine concept (175).

1.1.3 Crown Gall Tumours

The crown gall is characterised by uncontrolled and rapid plant cell proliferation at the wound site following infection by Agrobacterium. In nature invasion by Agrobacterium usually occurs at ground level near the junction of the stem and roots. The size of galls may vary in diameter from pea-size to 5-15 cm and can be up to 22.7 kg in weight (54).

The gall is composed of a mass of undifferentiated cells resulting from unlimited division and an increase in size of the vascular and parenchyma tissue (146).

Gall formation by Agrobacterium on dicot plants requires wounding which produces a wound meristem. Deeper wounds facilitate infection and usually result in large galls. Development of a tumour in a particular species of plant is dependent on the type of Ti plasmid possessed by the pathogenic Agrobacterium strains.

The cells of the gall synthesise specific compounds formed by condensation of an amino acid, a keto acid or a sugar and generically called opines which are absent from non-transformed cells. These compounds are specific growth substrates for Agrobacterium. Since growth of gall tumour is a benefit to virulent bacteria, the process has been named "genetic colonization".

After the induction of a tumour by the pathogenic Agrobacterium, the tumour cells continue to proliferate in the absence of bacteria. Also tumours induced by Agrobacterium, carrying wild-type Ti plasmids generally remain undifferentiated following transfer to culture. Analysis has shown that gall cells contain auxin and cytokinin sufficient for growth and cell division (54, 239, 242). The level of indole acetic acid in tumour cells is 2-500 fold higher than in non-transformed cells. The ratio between auxin and cytokinin in tumour tissue is likely to be a major determinant of the gall's phenotype. The proliferation of transformed plant cells results mostly from the expression of iaa and cyt genes, located on the Ti plasmid and responsible for the biosynthesis of auxin and cytokinin respectively.
1.1.4 Opines

One of the most significant characteristics of tumour cells is that they synthesise low molecular weight metabolites called opines. More than twenty opines have been identified, comprising nine structural families. Most opines are condensation products of amino acids with hexose sugars or with carbonyl compounds. A crown gall tumour synthesises only a limited number of opine molecules.

Based on the kind of opine produced by tumours, Agrobacterium tumefaciens strains are classified into different groups, octopine and nopaline being the most studied opine families. Octopine (N²-[1-D-carboxyethyl]-L-arginine) is synthesised by reductive condensation of a basic amino acid such as arginine, and pyruvate within plant tumours and degraded by octopine oxidase to arginine and pyruvate. This family consists of opines: octopine, lysopine, histopine, octopinic acid and mannitylopin (54).

Nopaline (N²-[1,3-D-dicarboxypropyl]-L-arginine) is formed by reductive condensation of 2-ketoglutarate with L-arginine, and similar to octopine, nopaline is degraded by nopaline oxidase to arginine and α-ketoglutarate. This family comprises of nopaline, nopalinic acid, and agrocinopine A and B (102).

The genes required for formation, utilisation and degradation of opines are on the Ti plasmid. According to the kind of opine synthesised by tumours, Ti plasmids are categorised into different types. Octopine and nopaline formation is carried out by octopine-type and nopaline-type Ti plasmids respectively.

Based on the effect of opines on the bacteria, opines often have been classified into two classes; the nutritional opines, which are used as carbon and energy sources, and the conjugal opines, which are used as inducer molecules for conjugation (61). Opines as energy sources are specifically metabolised by the virulent Agrobacterium strains, but not by most other soil bacteria or plant itself. Although the transformed plant cells produce these compounds. This ability creates an advantage for growth in the opine-rich environment by the tumour for those bacteria able to catabolise opines over non-catabolisers.

Production and utilisation of opines are binary in nature. The genes for biosynthesis of opines are located on a part of Ti plasmid called the T-DNA that is transfered to plant cells, and integrated into plant genome, driven by plant-active
promoters, while the genes that are on the Ti plasmid and allow the bacteria to use the specific opines, are activated by Agrobacterium promoters.

Occ and noc are the regions that encode octopine and nopaline utilisation respectively. Each region consists of different genes. The occ region carries the genes required for opine transportation, catabolism and also chemotaxis to opines (234, 235). Induction of these genes are carried out specifically by their cognate opines and are under control of proteins that have positive and negative regulatory functions called OccR and NocR for octopine and nopaline strains respectively (234). However the opine is not the direct inducer of these genes, involved in transport but is converted to the direct inducer during catabolism. The occ region encodes the regulator OccR and the transport proteins OccQ, M, P and J which allow the bacteria to uptake specific opines and then catabolise them (216).

In the octopine-type opine only one gene is required for synthesis of the opines while several genes encode enzymes involved in opine catabolism (ooxAB which converts octopine to arginine and pyruvate, orcA converts arginine to ornithine) (103).

Some opines acting as a conjugation inducer, stimulate Ti plasmid conjugation by enhancing expression of the tra gene which in turn activates other tra genes that are necessary for conjugation converting non-pathogenic strains into pathogenic strains. Opines, therefore seem to influence bacterial communities in the immediate vicinity of the tumour.

It has been shown that opines can translocate from tumour to non-transgenic part of the plant and released into the rhizosphere (175). This exuded opine can then be utilised by those Agrobacteria that have not induced the tumours but are present in the rhizosphere or on a part of the plant far from the site of the gall. Interestingly, certain soil isolates of Pseudomonas and Gram-positive coryneforms can utilise opines as sole sources of carbon and energy as well (175).

There is some evidence showing that opines can be beneficial to the galled plant in plant protection via antagonistic effect on insects and allelopathy against weed seeds (175).

Some opines are chemoattractants, and this chemotaxis is specific to the type of Ti plasmid present in the cell (103). Functions for chemotaxis require the regions encoding catabolism of that opine. occ and noc loci confer chemotaxis to their cognate opines. Chemotaxis to opines might be important in the interaction between
Agrobacteria and their plant hosts, and can play an important role in determining the
distribution of Agrobacteria in the tumour. Opines play a fundamental role in
recognition of plant hosts.

1.1.5 Ti plasmid

Ti plasmids are extra chromosomal elements whose host range is restricted to
the rhizobiaceae. Ti plasmids are generally 190-240 kb in size and present at a low
copy number (1-3 copies per cell) but can be transferred to other bacteria (90).
Although tumourigenic strains usually possess Ti plasmid but many non-tumourigenic
Agrobacterium, possessing plasmids, have also been isolated from soil (140).

Pathogenic Agrobacterium tumefaciens carries a specific Ti plasmid. According to the kind of opines produced by tumours, there are five Ti plasmid types:
1-Octopine type, 2-Nopaline type, 3-Agropine type, 4-Succinamopine type and, 5-
grapevine type(p) (54).

Despite differences between Ti plasmids, they each consist of five distinct
areas including three regions essential for tumourogenesis of bacteria (see fig. 1.4.1).
The regions are vir region, T-DNA region and opine catabolism region plus a region,
for conjugation containing tra and trb loci, and the rep region for replication
incompatibility (see figure 1.1.5.1). The plasmid also contains genes for conjugation
stimulated by opine, and sensitivity to the bacteriocin agrocin 84 in the nopaline and
succinamopine plasmids. Nopaline plasmids, such as pTiC58, also have a T-DNA-
independent locus, tzs which encodes an enzyme that produces the phytohormone
cytokinlin in the bacteria under vir-inducing conditions.

One part of the Ti plasmid is occupied by T-DNA which contains the genes
that are responsible for tumour formation (oncogenes) on the infected plants. T-DNAs
of different Ti plasmids have different compositions. Nopaline Ti plasmids carry T-
DNA as a contiguous stretch of ~ 22 kb, whereas the T-DNA in an octopine-type Ti
plasmid is composed of three independently transported segments that are left (TL)
with 13 kb, central with 1.5 kb, and right (TR) with 7.8 kb size (54). The borders that
flank and define the transferred region to the infected plant cells consist of a
conserved 25bp imperfect direct repeat sequence (86, 231). They serve as a target for
cleavage by the VirD2 endonuclease and play an important role in T-DNA
transformation (233). Experiments showed that the T-DNA borders can be replaced
by a bacterial origin of transfer (239). Mutation analysis showed that the right border is absolutely required for *Agrobacterium* pathogenicity, whereas the left border is not (179). Inversion of the right border has been shown to reduce virulence and result in transfer of nearly the entire Ti plasmid instead of the T-DNA region (239). These results suggest the polarity of T-DNA transformation from right to left, as determined by the orientation of the T-DNA border repeats. On the other hand, only the TL region is oncogenic and contains all the genes necessary for tumour formation, while the TR region has no genes for phytohormone production and is not oncogenic. Interestingly, the T-DNA itself (the sequences between the borders) has no effect on the efficiency of transfer (181, 239). Consequently, T-DNA sequences can be replaced by DNA sequences of interest for engineering plants. Now a days, non-oncogenic (disarmed) Ti plasmids are used widely as vectors for genetic transformation.
The T-DNA contains 8 to 13 genes, including a set for production of phytohormones, which are responsible for formation of the characteristic tumours on infected plants. These genes have features of plant genes rather than bacterial genes; e.g. they contain polyadenylation signals, are transcriptionally controlled by plant promoter and activator sequences and can be expressed in most dicot and some monocot plants. These genes encode enzymes which catalyse the initial steps in the conversion of adenine to zeatin and tryptophan to auxin, two plant hormones critical for cell growth, and also convert amino acids to opines. The TL region of octopine-type Ti plasmid contains the genes auxA and auxB that are responsible for auxin biosynthesis, cyt encodes cytokinin biosynthesis, and ocs controls octopine synthase. The TR region of these plasmid lacks the genes coding phytohormones and contains genes responsible for synthesis of further opines such as agropine and mannopine. In some cases, T-DNA transfer is enhanced by the so-called "overdrive sequence", a 24bp sequence, located to the right of the right border in octopine Ti plasmids.

The vir region which is 35-40 kb, consists of eight vir operons, virA to virG. Unlike octopine Ti plasmids, the nopaline type lacks the virF and virH operons but instead contains another accessory gene called tzs. The product of these genes, termed virulence (Vir) proteins, encode proteins involved in the detection of compounds that are released from wounded plants in the rhizosphere and mediate transfer of T-DNA into the host cells. Mutations in the virA, B, D, and G operon abolish tumour formation on all plant species, while mutation in the other vir genes reduces tumourgenicity and restricts the range of plant hosts.

The opine catabolism region, occ region, consists of genes that allow virulent *Agrobacterium* to take up opines and utilise them. The region contains the genes that are involved in the transportation and metabolism of octopine. These genes are under the control of an opine-inducible promoter, activated by the product of occR acting as a positive regulator. Likewise octopine type Ti plasmids, a similar set of genes are on the nopaline type Ti plasmid in the noc region. Another gene, ons, encodes a protein required for transportation of octopine across the plant cell membrane. Chemotaxis to opines is also carried out by function of the opine catabolism operon. In nopaline-type *A. tumefaciens* noc also confers chemotaxis to their cognate opines.

Replication and incompatibility region also maps to the Ti plasmid. In terms of compatibility, there are two Ti plasmid groups: incRH-1 and incRH-2. Another
gene which is responsible for replication is ein. All octopine plasmids belong to the same ein-group, whereas nopaline plasmids belong to one of two different groups.

1.2 Virulence in Agrobacterium tumefaciens

Agrobacteria infect their hosts at wound sites (109), which exude specific compounds into the rhizosphere. These chemicals switch on the virulence engine of pathogenic Agrobacterium tumefaciens which possess a Ti-plasmid to infect the plants. Although, wounded cells are necessary for bacterial attachment, however there is evidence showing that Agrobacteria are able to interact with unwounding plant cells and eventually to transform them (67). Agrobacterium can enter leaves using their stomata. The normal invasion process occurs in three levels. Those carried out in the rhizosphere, resulting in attachment of the bacteria to plant cells, those that occur in Agrobacterium itself, including preparation of the T-DNA for transformation, and finally the processes that happen in the host plant, leading to transformation and integration of the T-DNA into the plant genome. The virulence process can also be divided into different steps, as discussed in the next section.

1.2.1 Host-pathogen contact

Cell-cell contact between pathogen and host cells is necessary for many diseases including gall tumour formation. The first step in plant infection is attachment of pathogenic Agrobacterium to the surface of wounded plant cells. Non-attaching mutants are avirulents (54, 115).

The attachment process consists of two stages:
1. Direct attachment of bacteria to plant cells. This initiates a loose binding and is mediated by a bacterial Ca$^{2+}$-binding protein called Rhicadhesin. This protein is a major determinant of attachment and virulence. Ca$^{2+}$ plays a role in anchoring, stabilizing and for activation (130, 186).

2. Anchoring of bacteria to each other and to the plant cell surface using bacterial cellulose filaments, cellulose fibrils, stabilizes the initial binding. This generates a tight irreversible binding between the bacteria and the host cells (129). In this stage other bacteria not directly attached to the plant cell, create a large aggregation of bacteria and fibrils around plant cell surfaces. In fact only a small fraction of the
bacteria are directly bound to the host cell. The cellulose fibrils have an important role in pathogenesis. Mutants which cannot synthesise cellulose are reduced in virulence to 1/10th to 1/1000th that of the wild-type bacteria. Cellulose synthesis in \textit{Agrobacterium tumefaciens} involves two operons of the bacterial chromosome, containing five genes (\textit{cel} genes). Cellulose fibrils are produced at random by \textit{Agrobacterium tumefaciens} from all sides of the cell surface (130).

Direct attachment of only a few numbers of bacteria to the host cell, suggests that plant cells could have specific receptor sites. There is evidence showing that the plant cells have a receptor of protein and pectin located on their surface, and a second binding site of unknown composition, on the plasma membrane (167). The bacteria also have a binding site composed of protein and lipopolysaccharide, a component of the cell wall, located on their outer membranes (115).

The genes involved in attachment are located in the \textit{Agrobacterium} chromosom, and include \textit{chvA}, \textit{chvB}, \textit{pscA} (or \textit{exoC}) and \textit{Att} (181).

The success of interaction between \textit{Agrobacterium tumefaciens} and the host cells depends on the ability of the bacteria to go through the plants defenses. In the rhizosphere, \textit{Agrobacterium tumefaciens} also must compete effectively with other microorganisms as it attaches to the plant cell.

\subsection*{1.2.2 Plant Signals}

The next step in the virulence process is induction of the \textit{vir} genes which brings about transfer of the T-DNA into the plant cell. Activation of the \textit{vir} genes requires the presence of specific inducers which are released from wounded plants into the rhizosphere (177). The sap from wounded plants has a characteristic acidic pH of 5.0-5.8, and contains a high amount of different aromatic and aliphatic chemicals such as phenolic compounds, which serve as plant signals, and specifically stimulate \textit{Agrobacterium vir} gene expression. Phenolics also serve as attractants, activating chemotaxis in \textit{Agrobacterium} (182, 187, 188). In addition, thousands of healthy somatic cells, with intact cell walls, which are released from the roots of many plant species into the rhizosphere everyday, are capable of producing extra cellular signals that induce microbial gene expression (237).

Phenolic structures alone are sufficient to induce the expression of the \textit{vir} genes. Acetosyringone, a monocyclic aromatic hydrocarbon and \(\alpha\)-hydroxy{
acetosyringone, which are released naturally from injured plants, but are not found in undamaged plants, are examples of effective phenolic inducers. Other phenolic inducers are lignin precursors coniferyl alcohol, sinapyl alcohol, ferulic acid, and sinapinic acid (187). Some of these inducers are intermediates in cell repair biosynthetic pathways and for cell wall repair (181). So far, more than 40 low-molecular-weight phenolic inducers have been identified. Most of the monocots do not release such phenolic compounds; hence the virulence by Agrobacterium cannot be processed in these plants.

It is worth noting that most of the compounds released from wounded plants are bactericides at high concentrations and many also contain inhibitors of vir induction. Monocots also release some of these inducer compounds suggesting that Agrobacterium is able to recognise monocot wound sites, but further steps in the infection process might be blocked (13). Induction of the vir genes is dependent on a number of environmental factors including pH, presence of some phenolic compounds, presence of some monosaccharides, phosphate levels and presence of some heavy metals. The optimal conditions required by various strains for vir-induction are slightly different, however all require particular phenolic compounds and a pH of 5.0-5.8 (214).

1.2.3 Signal transduction

Detection of the plant signals that induce the vir genes is mediated by a two-component sensory regulatory system. The microorganisms use this kind of system to monitor their surroundings and make appropriate responses as they face high competition for limited resources and have little ability to change their environment. These systems respond, with a high sensitivity and efficiency, to changes in chemical environments around the bacteria. Changes in chemical composition are translated to signaling pathways often involving two families of signal-transducing proteins. One family is a membrane sensor protein that is responsible for recognising the changes in extracellular signals and transduced into appropriate intracellular events. These compounds surrounding the bacteria, and the other member that is a cytoplasmic protein, that transduces information from the membrane sensor to the cell. It has been suggested that a bacterium contains about 50 pairs of these sensors that regulate many of fundamental functions, such as motility and pathogenicity (197). The key function
acetosyringone, which are released naturally from injured plants, are examples of effective phenolic inducers. Other phenolic inducers are lignin precursors coniferyl alcohol, sinapyl alcohol, ferulic acid, and sinapinic acid (187). Some of these inducers are intermediates in cell repair biosynthetic pathways and for cell wall repair (181). So far, more than 40 low-molecular-weight phenolic inducers have been identified. Most of the monocots do not release such phenolic compounds; hence the virulence by Agrobacterium cannot be processed in these plants.

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of signal transduction involves phosphorylation of the membrane sensor protein that is a protein kinase.

Agrobacteria employ a two-component signal transduction system, composed of the virulence proteins VirA and VirG, to regulate infection of plants (223, 224, 226). Both VirA and VirG are the Ti-plasmid encoded genes required for chemotaxis and virulence induction mediated by phenolic compounds. The VirA/VirG system is thought to have multifunctional role, since at different phenolic concentrations it is known to affect both chemotaxis and Vir-induction. Low concentration (nonomolar) trigger chemotaxis alone, higher concentration (micromolar) also bring about Vir-induction and can suppress chemotaxis, thereby restricting the bacteria to the wound site and hence increasing, the infection likelihood. In *Agrobacterium* therefore, compounds released from plants trigger two specific activity, chemotaxis and virulence.

VirA is a sensor protein acting directly or indirectly as a receptor for phenolic compounds, like acetosyringone and specific monosaccharides, which are plant metabolites or monomers of plant cell wall polysaccharides (13). It has autokinase and phosphotransferase activity in vitro, resulting in autophosphorylation on amino acid histidine 474 (93, 226). It is an inner membrane protein, composed of four structural and functional domains: i) a receptor domain consisting of an N-terminal periplasmic domain and two transmembrane regions, designated TM1 and TM2, ii) a linker domain (cytoplasmic domain), iii) a transmitter domain that contains protein kinase activity, and iv) a phosphoryl receiver domain (40, 111, 133). Investigation into the properties of VirA have revealed that the periplasmic domain may not be the sensor domain for phenolics and direct interaction between VirA and acetosyringone requires that this metabolite diffuse through the outer membrane to reach VirA, which lacks extracellular domains (226). The periplasmic domain has a role in the detection of changing of pH and temperature of the environment. It has been suggested that VirA itself may not bind phenolics, rather plant phenolics may initially interact with chromosomally encoded protein of 10 and 21 kDa called p10 and p21. Binding to these complexes of phenol-binding proteins then activates VirA (166).

Some sugars, in particular the acidic monosaccharides D-galacturonic acid and D-glucuronic acid (both monomers of major plant cell wall polysaccharides) and other sugars such as glucose, arabinose, and galactose enhance vir-induction when acetosyringone are absent or present in small amounts (<10μm) in the environment.
The product of the chromosomal chvE gene is required to interact with these sugars and in turn interact with the periplasmic domain of VirA (51). This reaction results in supersensitivity of VirA binding to phenolic compounds. Transcription of chvE is regulated by the gene gbpR which lies adjacent to chvE on the chromosome. (64) In contrast, low amount of opines that are attractants for certain Agrobacterium, enhance vir gene expression in the presence of acetosyringone in the environment. This process is Ti-plasmid dependent and requires functions encoded by the opine catabolism operons of the Ti-plasmid.

As a general rule, conditions appropriate for vir gene induction are unsuitable for bacterial growth, whereas conditions that support bacterial growth are insufficient for vir gene induction. Therefore, it has been proposed that in low opine concentrations, Agrobacterium stay in the optimal state for vir gene induction, whereas at high levels of opine (produced in mature tumours) Agrobacterium are able to grow and use the opines as an energy source (181).

Interaction between plant signals and VirA leads to autophosphorylation of VirA at its histidine 474 residue in the transmitter domain. The phosphohistidin high-energy phosphate bond is then transferred to the ASP-52 or possibly the ASP-8 residue of VirG. The phosphorylated VirG is very stable, this is thought to facilitate maximal levels of vir gene induction (181).

The VirG protein is a cytoplasmic protein, but may also be partially associated with the Agrobacterium membrane. It acts as a transcriptional regulator of the other vir genes. To activate expression of the other vir genes, the C-terminus of VirG interacts with the vir-box, a conserved 12-bp sequence, which is located in the 5'-non-coding regions of all vir operons including virA and virG (94, 158). Each vir operon possesses up to five vir-boxes. The absence of a vir-box from the virA promoter is one factor restricting the number of plant species on which this strain can produce tumours (215). Specific nonconserved sequences downstream of the vir-boxes are required for perfect VirG function. These sequences prevent interference from other transducer proteins (165). Phosphorylated VirG is thought to have a high binding affinity for vir-boxes. There is also a possibility that VirG-P interacts with RNA polymerase, increasing RNA polymerase efficiency to recognise the vir promoter, or to open the closed DNA complex (89). The presence of vir inducers enhances the expression of virA and virG. Phosphate starvation also upregulates
expression of virG. In addition, the virG promoter is pH-inducible through a second promoter (17, 128, 225).

It is not known whether VirG is phosphorylated before or after binding to the vir-box. The actual role of phosphorylation is not clear, since unphosphorylated VirG can specifically bind vir promoters, however in vivo, VirG phosphorylation is required for vir gene expression. Phosphorylation of VirG may increase its DNA binding affinity (83).

The levels of expression of the vir genes are determined by the pool size of VirG. VirG also activates virA and virG transcription and therefore forms a positive feedback loop enhancing the infection process. Regulation of the vir regulon is also temperature dependent. The levels of expression decrease significantly when the temperature is raised above 22°C.

In conclusion, the compounds that are released from wounded plants into the rhizosphere act as plant signals and inducers. These signals are detected and transduced by A. tumefaciens activating vir gene expression. Sensing and transducing of the signals are mediated by VirA and VirG members of two-component regulatory system.

1.2.4 T-strand generation

The T-DNA carries all the information necessary for tumourgenesis. Preparation of the T-DNA is carried out by the vir genes and involves isolation of the T-DNA from the Ti-plasmid. A linear, single-stranded DNA molecule, called the T-strand is generated (181, 193). This process occurs in the bacterial cell. The T-strand is transferred into the plant cell in a form called a T-complex.

The first step of T-strand formation is isolation of the T-DNA region from the Ti-plasmid. Two VirD proteins, VirD1 and VirD2, encoded by the virD operon are together responsible for nicking the 25-bp border repeats that serve as nick site for cleavage of the T-DNA (193, 239). The virD operon contains four open reading frames. VirD1 encodes a topoisomerase enzyme that catalyses the conversion of supercoiled DNA to relaxed DNA (72, 89, 181). VirD2, encodes an endonuclease that recognises the right T-DNA border. These two proteins act as a specific border repeat nickase and cut at specific sites, between the third and forth base pairs of the bottom strand of the borders (181) Excision is thought to occur in a 5' to 3' direction, starting.
at the right T-DNA border and terminating at the left border. This initiates a free single-stranded T-DNA molecule or T-strand. Following cleavage VirD2, but not VirD1, remains tightly attached to the 5' end of the nicked T-strand at the right border nick and to the 5' end of the remaining bottom strand of the Ti-plasmid at the left border nick (47, 235). VirD2 possibly binds via the tyrosine 29 residue (181, 218, 239). This binding is thought to protect the T-strand from exonucleolytic degradation and probably prevents closure of the nick by ligase activity. In addition the bound VirD2 is thought to act as a pilot protein and direct the T-DNA to the plant nucleus.

When the T-strand is removed, the resulting gap in the Ti-plasmid will be repaired by generation of single-stranded copies. The replacement process removes the VirD2 molecule attached to the 5' end of the left border, allowing reformation of the circular DNA molecule of the Ti-plasmid. Recent data also suggests that VirD2 may have a role in ligation of the left border (154).

The final step in T-strand formation involves a 26 kDa protein from virC operon, called VirC1. This protein binds specifically to the overdrive sequence, which is a natural enhancer, located adjacent to the right border repeat. Overdrive sequence functions effectively in an inverted orientation and up to about 7 kb away from the right border repeat in either directions. This sequence is necessary for optimal T-strand formation when VirD1 and VirD2 are limited. VirC1 is thought to facilitate nicking of the T-DNA border and may play a role in T-strand export (236).

1.2.5 T-complex formation

There is some evidence indicating that the T-strand is transferred from the bacterium into the plant cell in a nucleoprotein form, called the T-complex. The T-complex is composed of at least three components. The single-stranded T-DNA molecule, carrying all the information for tumourgenesis, associated with two types of Vir proteins, VirD2 and VirE2 (64).

The endonuclease activity of VirD2 lies entirely within the N-terminal of the protein, since this domain of the VirD2 protein is sufficient to cut the border repeat and T-strand formation, but is unable to import the T-DNA into the host cells. It seems that the C terminus of VirD2 must be involved in a step after T-strand generation. Genetic analysis has shown that this domain contains a bipartite nuclear
localization signal (NLS) and conserved sequence called omega (DGRGG), located adjacent to the NLS. They are shown to be important, but not essential for tumourgenesis (88). The omega sequence may interact with other Vir proteins involved in transport of the T-DNA-VirD2 complex into the plant nucleus or may involve in the integration process (32).

The virE operon encodes a 7.0 and a 60.5 kDa polypeptide in octopine Ti-plasmids and a 64 kDa polypeptide in nopaline Ti-plasmid. VirE2 is a single-stranded DNA binding protein that attached at its 5'-end to the VirD2 protein. It is thought to protect the DNA against degradation by bacterial and plant exonucleases and endonucleases during transformation, and shape the T-complex into a thin and unfolded structure (transferable form). It may also provide specific targeting signals. Based on electron microscopy data, the nopaline-specific T-complex is proposed to be 3600nm long and 2nm wide. This suggests that the T-complex would contain approximately 600 molecules of VirE2 (47)and only one of VirD2, with a molecular mass of 50,000 kDa (181). It was reported that ssDNA-binding protein, AcvB (or its functional homologue VirJ), may also associates with the T-complex (59).

It is thought that the T-complex is formed in Agrobacterium and then all components of the T-complex together transfer to a recipient plant cell. The T-complex model proposes that the 5' end of the T-strand is associated with the VirD2 molecules, whereas the 3' end probably has a VirE2 protein attached in its proximity. It has also been suggested that VirE2 and the T-strand are transported independently from Agrobacterium into the plant cell, and that VirE2 functions inside the host cells (49, 77, 181).

Recently, data indicated involvement of another VirE product, named VirEl, in transportation of VirE2 from the bacterium into the plant cell. VirEl acts as an export chaperone for VirE2 (58, 202). VirEl, which is located adjacent to its targeted protein with no identified enzymatic activityIt, belongs to a new class of chaperones required for the export of virulence factors via a typeIII secretion pathway. It shares a number of properties with the typeIII secretion chaperones (for more details about chaperones see section 1.4.3 and Deng, 1999). This small protein is specifically required for the export of the VirE2. VirEl prevents VirE2 to bind to T-strand by forming VirE2-VirEl complex (24), inhibits VirE2 from aggregation, enhances the stability of VirE2 and perhaps maintain VirE2 in an export-competent state. This means that within Agrobacterium, VirE2 may not bind to the T-strand to form the T-
complex; rather may VirE2 enter the plant cell with assistance from VirE1 independently from the bacterium and formation of the T-complex and then bind to the T-strand-VirD2 complex. Another possibility is that VirE1 may facilitate export of the entire T-complex (77, 110, 202).

1.2.6 T-DNA transformation to the plant cell

It is not clear exactly how the T-DNA leaves the bacterium, crosses the membrane barrier, and enters the plant cell. Transportation of the T-DNA from donor bacterium into recipient cells (see fig 1.2.6.1) requires a passageway between the two cells. It is thought that the T-complex crosses the bacterial membrane and plant cells through a channel built by protein-protein connection of vir gene products (see fig. 1.2.6.2). It seems that some of the proteins encoded by the virB operon are involved in this bacterium-plant cell channel. The virB operon is the largest vir operon and contains 11 genes (44). It has recently been shown that the vir genes belong to a family of export system, called typeIV secretion system. The function of this system, which have members in α, β and ε subgroups of proteobacteria, is direct transport of protein or DNA-protein complex across the cell envelopes of bacteria to host cells (239). Ten of them have been shown to associate with the bacterial membrane, where they may form a structure, a pilus/pore, necessary for transfer of the T-complex. The typeIV secretion system comprises two major structural components: a filamentous appendage or pilus, and a membrane associated complex that translocates substrate across both bacterial cell membranes and possibly across the host cell membranes. Based on recent studies, it has been shown that the T-DNA transfer machinery is homologous to bacterial conjugal DNA transfer system. This has been supported by the fact that most of the VirB proteins show homology with different Tra proteins of the bacterial conjugal transfer systems. In fact vir genes belong to a family of export system, called typeIV secretion system. The function of this system, which have members in α, ε and γ subgroup of proteobacteria (108), is direct transport of protein or DNA-protein complex across the cell envelopes of bacteria to bacteria or from bacteria to eucaryotic cells (35, 45).

The products of two vir genes, virB9 and virB10, are known to form a high molecular-weight complex with other membrane proteins. VirB9 is thought to stabilize or facilitate formation of a VirB10 complex. VirB7 and VirB8 stabilize
VirB9 in turn. The stabilization of some VirB proteins, in the presence of other VirB proteins, suggests that the stabilized proteins may form a multiprotein channel structure necessary for T-DNA departure.

T-complex transformation is likely to require energy. Two proteins, VirB4 and VirB11, are associated with energy supplement activity. VirB4 has a nucleotide-binding site, whereas VirB11 is an ATPase and a protein kinase, both are inner membrane proteins (44, 181). VirB1 a low molecular-weight protein, is thought to interact with the recipient plant cells facilitating anchorage of the VirB channel at the infection site on the cell surface. Recent data has shown that a protein from the VirD operon, VirD4, is required for T-DNA transfer and tumourgenesis. This protein which

1.2.6.1 Diagram showing main steps of transfer of T-DNA to plant cell. Figure adapted from (68).
is anchored in the inner membrane of *Agrobacterium*, may play a role in the transformation of the T-complex by forming a link between the T-complex and the VirB channel. It contains a possible ATPase consensus sequence and therefore could generate the energy for T-complex transportation through the channel (114, 151).

Briefly, the T-complex is thought to be transferred from *Agrobacterium* to plant cell via a membrane pilus/pore channel (perhaps T-pilus) made up of the VirB and VirD4 proteins (108).
1.2.7 T-DNA integration into the plant genome

When the T-DNA enters a host plant cell T-complex form consists of the T-DNA bound with one VirD2 on its 5’ end (60, 105) and hundreds of VirE2 molecules bound along its length (233, 239, 241, 242), it needs to be directed to the nucleus. It has been proposed that an endogenous plant cellular pathway is used to import the T-complex into the cell nucleus. As a rule, the active nuclear import system requires a specific nuclear localisation signal (NLS) as a targeting signal that is recognised by the plant. Since the T-strand itself does not have an NLS, the transformation of the T-strand is mediated by functions of the VirD2 and VirE2 proteins, which both contain nuclear localisation signals (48, 49, 84, 181). It has been suggested that the T-strand is directed to the nuclear pore in a polar fashion via the bound VirD2 that function as a pilot protein within the plant cell. Each of the two proteins, VirD2 and VirE2, has two nuclear targeting signals (181, 211, 239).

Some evidence has shown the accumulation of T-strands in vir-induced *Agrobacterium* and presence of single-stranded, but not double-stranded T-DNA in the host plant cells soon after infection (210). These results together with an experiment on properties of double-stranded and single-stranded DNA, suggest that the T-DNA enters the plant cell nucleus in a single-stranded form.

The mechanism of integration of the T-DNA into the plant genome is still not well known. Data available from different plant species indicates that the integration site is random and active transcriptional chromosomal regions are targeted. Insertion appears to be randomly distributed between translated and untranslated regions (209). T-DNA integration is not site specific, however it seems that there is a preferences for integration into transcriptionally active loci (89).

T-DNA integration is a multistep process involving several activities, such as recombination, replication and repairing. Unlike other mobile DNA elements such as transposons and retroviruses, T-DNA does not contain enzymes enabling integration. Thus T-DNA insertion into the plant genome must be mediated by proteins accompanying the T-DNA into the host plant cells and/or by host cell factors. The only bacterial proteins that are transported into the plant cells are the two polypeptides of the T-complex, VirD2 and VirE2.
Integration is an "illegitimate or random recombination" process that does not require long stretches of homology (89). Sequence homology between the incoming T-DNA and the plant chromosomal DNA has an important function in T-DNA integration. Sequence analysis showed that the left-hand end of the T-DNA (3'-end of T-strand) has a short region of homology (at least 5 contiguous nucleotides) with the insertion site of the plant chromosome, while the right-hand end of the T-DNA (5'-end of T-strand) which is attached to the VirD2 protein is restricted to just one base homology (microhomology) (209). A small deletion usually occurs at the integration site of plant DNA (less than 100 bp) Analysis of these sites has shown that the right border of the T-DNA is preserved, possibly because of VirD2 attachment to the end of the right border that protects it from the plant DNA repair machinery. However, the left border end can lose up to 1000bp. It is therefore thought that VirD2 is involved in the integration process as a ligase enzyme (239).

It has been shown that the T-DNA is inserted into the plant genome in a single-stranded form and then second-strand synthesis, is performed by the plant cell DNA repair machinery. However there is a second study of T-DNA suggesting that the T-strand is converted into a double-stranded form before integration (181).

Both models proposed that T-DNA integration starts with synapsis between plant DNA and T-DNA at the 3'-end of the T-DNA (left border), where there is a short region homology between the two DNA strands. This step is followed by the ligation of the 5'-end of the single-stranded T-DNA to the 3'-end of the plant cell DNA. The plant enzymes such as ligase perform ligation, possibly using the energy-rich phosphotyrosine bond that links VirD2 to the T-DNA. Attachment of the T-strand to one strand of the plant DNA results in a second nick on the opposite strand of the target site. Each end of the T-strand is then ligated to the plant DNA and cellular enzymes using the T-DNA as a template copy its homologous strand. A short sequence of five amino acids (DGRGG) located downstream of the VirD2 NLS, called the omega (ω) domain was found to be necessary for T-DNA integration (183).

Although the host cell enzymes mediate much of the integration of the T-DNA into the plant genome, a part of VirD2 and other virulence proteins may also play very important roles in this process. New evidence suggests the direct involvement of VirE2 in the recombination process. This protein is required for integration of the T-strand at the 3' end, but not at the 5' end of the T-strand molecule. In the presence of VirE2, the 3'-end is only subjected to very short deletions not exceeding 100

23
nucleotides. VirE2 probably physically protects the T-DNA from plant nucleases. Another possibility is that VirE2 is somehow involved in the conversion of the T-strand into a double-stranded form.

DNA rearrangements, which include duplication, deletion, inversion and base substitution have been found after integration of the T-DNA into plant DNA at the end of the insert T-DNA. This can also occur during repair and replication activities.

1.2.8 T-DNA expression

The integrated T-DNA genes encode enzymes that catalyse opine synthesis and its subsequent transport, and enzymes that alter the levels of plant growth regulators. Two genes, iaaH and iaaM, are responsible for enzymes that are involved in the early steps in the conversion of tryptophan to auxin, a hormone causing cell expansion. Another gene, ipt, provides a product that is involved in the first step of trans-zeatin and trans-ribosylzeatin production (a cell division factor) from adenine (236).

The integrated T-DNA also contains genes involved in biosynthesis of opine (104, 236). Octopine-type Ti-plasmids direct their host to produce at least eight opines (236).

The transferred genes are transcriptionally controlled by eukaryotic promoter and activator sequences and can be expressed in most, if not all, eukaryotic cells (224). The promoter activity of several T-DNA genes from Agrobacterium tumefaciens have been shown to be organ or tissue specific, developmentally regulated and/or influenced by plant hormones (65, 224).

The expression level of integrated T-DNA genes can be affected by different plant factors. As the T-DNA that enters the nucleus lacks a chromatin structure, besides DNA modification pattern, random methylation, mutation, rearrangements or interactions with host cell factors, occurring prior to, during or after integration, might greatly influence the expression (33).

1.3 Chemotaxis
As mentioned earlier (section 1.2.3), it is important for microorganisms to monitor the environment surrounding them as they face a high competition for limited resources. They use different sensitive sensory-response systems to detect and respond to the changes in the chemical or physical environments. These informations are converted to signals, which allow cells to make appropriate adaptive responses essential for cell surviving and growth. Such sensory-response systems, usually called two-component regulatory system or His-Asp kinase systems, usually consist of receptor, histidine kinase and aspartate kinase elements. Over 50 different two-component systems are operating in \textit{E. coli} (29, 30, 38, 39, 69, 196). These systems generally control many physiological activities of bacteria including cell division, virulence, motility, adhesion, antibiotic resistance, metabolite fixation and utilization, response to environmental stress, sporulation, and different behavioural responses such as phototaxis, aerotaxis and taxis in response to cellular energy changes (69). This conserved monitoring mechanism has also been demonstrated properly in several other systems such as regulation of outer membrane porin synthesis in response to change in medium osmolarity, control of gene expression in response to the intracellular balance between carbon and nitrogen metabolism, and chemotaxis in response to attractant and repellent gradients. With such different two-component systems, specificity of kinase- response regulator interactions is crucial to avoid unwanted cross talk.

Up to present, one of the best-known two-component pathway in bacteria is the chemosensory system of \textit{E. coli} which has become one of the leading models to describe the mechanism of the such systems. As this can provide a bright molecular picture of the chemotaxis behaviour in microorganisms, and since the principles of chemotaxis are similar in most bacteria, the chemosensory pathway of \textit{E. coli} will be discussed in more detail.

There are at least three systems for chemoreception in \textit{E. coli} that must link together in the central signaling pathway of chemotaxis. They are the aerotactic response and taxis to other electron acceptors (9, 10), the methylation-independent behavioural response to carbohydrates transported by the phosphoenolpyruvate (PEP) dependent phosphotransferase system (PTS) (7, 9, 205) and methylation-dependent chemotaxis (155, 199, 206).
There are two chemotaxis pathways in *E. coli*, a primitive system linked to metabolism via the citric acid cycle and a second system which is metabolism-independent with dedicated, transmembrane receptors.

Chemotaxis is the directed movement of bacterium to a component of attractant (generally nutrients such as amino acids and peptides) or away from repellent gradients (generally toxic compounds) (2). These behaviours are called positive and negative chemotaxis, respectively. The movement is carried out by flagella which rotate in either clockwise direction (CW) making random tumbles or counter-clockwise direction (CCW) producing runs (123). In the liquid media and in the absence of a chemical gradient, bacteria appear to swim in a three-dimensional random walk consisting of runs in a smooth, straight line lasting 1-2 seconds followed by tumbles during which the cell stops and randomly reorients before swimming in a new direction (20, 201). A pause seems to result from CCW to CW futile switching attempt, and there is no signal for it. In the presence of attractants or repellents, this simple swimming pattern is transferred into complex cellular behaviour. When a gradient of attractants or repellents is present, the chemosensory pathway monitors the change of the chemical concentration with time as the cell swims through different regions of the gradient. If the system detects an increase in attractant or decrease in repellent concentration with time, a signal will be sent to the motor switch to suppress CW rotation of flagella and subsequent tumbles, thereby lengthening the run towards the highest attractant concentration. Conversely, when the pathway detects a decrease in the attractant concentration or an increase in the repellent concentration with time, tumbling event will be generated by CW rotation of flagella, so that runs in this direction becomes shorter. These processes change the random walk into a walk to or away from ligands. Although such simple chemotaxis pattern is studied under controlled condition and solution media, however in natural environments the taxis on the surface of wetted substrates, likely shares some of the same switching behaviour observed for swimming in solution (69).

*E. coli* has memory, which allows detection of changes in gradient concentration. As the cell swims, *E. coli* compares the current chemical concentration that monitored by the receptor, to that chemical of the recent past as remembered by the methylation level (26).

Some compounds recognized as attractants for *E. coli* are certain amino acids, sugars, carboxylic acids, oligopeptide (1), serine and aspartate that providing the
strongest response, whereas repellents include organic molecules such as phenol, benzoate, indole, ethanol, ethylene glycol and some inorganic ions such as $\text{Co}^{2+}$, $\text{Ni}^{2+}$, and $\text{S}^{2-}$ (125). Oxygen, fumarate and nitrates, which can act as electron acceptors in the respiratory chain, are chemoattractants (205). Chemotaxis also occurs to variation in pH and temperature (125).

The chemosensory pathway between bacteria shares nearly the same set of components, however the mechanism that operates through the system shows a great deal of variation, both between and within different bacteria. Most pathway components can be swapped between species to generate functional chimeric receptors with the expected changes to ligand specificity and without reduction in function (69).

Chemosensory proteins that communicate with one another to relay the information coming from the environment in the form of chemical signals, and regulate cellular behaviour carry out activation of chemotaxis. Three classes of proteins are involved in chemotaxis: transmembrane receptors, cytoplasmic signaling components, and adaptive methylation enzymes. Communication between the receptors and the flagellar switch motor involves four proteins which are CheA, the histidine protein kinase, CheY the response regulator, CheW, the receptor-coupling protein; and CheZ, an enhancer of CheY-P dephosphorylation. This part of the chemotaxis system works by phosphorylation reactions during intracellular signaling.

The chemosensory pathway in *E. coli* begins when the receptors sense and bind to the small-molecule ligands which enter the periplasmic compartment. The receptors, then bind to and activate the cytoplasmic histidine kinase, CheA. The receptor is coupled to the CheA by CheW protein providing a stable receptor-kinase complex. The kinase is autophosphorylated at a specific histidine residue, then the phosphate is transferred to an aspartate residue of a response regulator CheY which activates the motor switch by binding to FliM, a motor switch component. Finally, the CheZ protein provides a specific phosphatase activity which facilitates the dephosphorylation of CheY. Two other proteins, CheB and CheR which are involved in adaptation phase of chemotaxis, controls the covalent modification of the receptors by an adaptive methylation pathway (38, 69, 141, 201)].
1.3.1 Transmembrane receptors

The chemosensory signaling pathway in *E. coli* which works through phosphorylation, begins with sensing specific compounds in the environment surrounding the cell. Most of these signals are carried out by a set of proteins (receptors), often called MCPs (for methyl-accepting chemotaxis proteins). Some of the chemicals directly bind to the receptors but some must bind to one of the binding proteins and then, to the receptors. There is a set of at least five water-soluble binding proteins located in the periplasmic compartment of the cell that recognize some of the chemoattractants and also act as ABC transport receptors. They are: ribose-binding protein (RBP), glucose/galactose binding-protein (GBP), maltose/maltodextrins (MBP), dipeptides-binding protein (DBP), and Nickel-binding protein (in repellent response, NiBP) (69). When a chemical is bound to a binding protein, the complex of the binding protein and chemical is linked to the surface of its cognate transmembrane receptors, thereby initiating a signal process or passes the ligand to a transporter.

So far, five transmembrane receptors have been found in *E. coli* and the related bacteria, *Salmonella typhomurium* which recognize different attractants. The five chemoreceptors and their respective chemoattractants are: (a) Tsr, serine; (b) Tar, aspartate and maltose; (c) Trg, ribose, galactose; and (d) Tap, dipeptides. *Salmonella* lacks Tap but possesses Tcp, a citrate sensor and Tip (201). Trg and Tap recognize only binding proteins, Tsr only small molecules, while *E. coli* Tar is uniquely capable of sensing both small molecules and binding protein ligands. Tar can respond to MBP and aspartate as attractant and the Ni$^{2+}$ and CO$^{2+}$ as repellents. All of the five MCPs in *E. coli* show that the proteins are closely related and have the same structural domains. Recently, a sixth receptor involved in aerotaxis has been described, which also appears to regulate the CheA/CheY two-component signaling pathway (22). This receptor possesses a different domain organization at its N-terminus in the cytoplasm that is a suitable location for sensing the redox potential of the electron transport chain or other components of the cell interior.

Maltose, ribose, galactose, and dipeptides bind via binding proteins to their corresponding transmembrane receptors, whereas serine, aspartate, and citrate bind directly to the receptors without the assistance of a binding protein. Response to temperature, pH, and various repellents are also mediated by the receptors. It is also noteworthy that each of the transducers is also responsible for movement away from
one or more repellents, however no receptor for repellents has so far been directly identified.

Each of the four soluble binding proteins is a monomer and they range in size between 30 to 51 kDa. A fifth binding proteins has also been found to detect Ni$^{2+}$ as a repellent through Tar. All binding proteins contain a conserved two-domain structure linked by a hinge of two or three strands. This hinge is a critical feature, allowing the proteins a dynamic equilibrium between open and closed forms that can be shifted by binding the appropriate small molecules. The ligand binds in the deep cleft between the two domains. The state of cleft, opened or closed, plays an important role in binding of ligand. The closed form of the protein is the active form in both chemotaxis and transport, whereas open forms predominate when the ligand is not bound and provide a way to reduce inappropriate interaction with the membrane components. Closed forms do occasionally occur in the absence of ligand. The open form of the cleft is essential for ligand binding and release, whilst the closed form of cleft by ligand induction seems to be important for successful binding to a transmembrane receptor. Cleft closure regulates a large receptor-docking surface.

The classical receptors which are also called "MCPs" for methyl-accepting chemotaxis protein, consist of three main parts. They are: a) sensory domain, b) transmembrane domain and, c) a large cytoplasmic signaling and adaptation domain. The latter domain binds to the cytoplasmic proteins CheW, CheA and CheR generating the signaling complex. CheY and CheB also dock competitively to the complex creating a large signaling complex containing most of the pathway components except CheZ.

Each cytoplasmic domain contains four or five methylatable glutamate residues. They are activated by methylesterification and the receptors are therefore called methyl-accepting chemotaxis proteins (MCPs). There is a C-terminal called "tail" can be easily removed.

Some of the MCPs of *E. coli* and *Salmonella typhimurium* are thought to be localized primarily at one end of the rod-shaped cytoplasmic membrane called a "nose", although the cell orients this nose either toward or against the direction of swimming (126, 156).

The sensory domain of the transmembrane receptors that recognize certain chemicals is located in the periplasm at the outside surface of the cytoplasm. The periplasmic domain of the aspartate receptors consists of two 19 kDa subunits and
each of them is an antiparallel four-helix bundle. Two symmetric aspartate-binding sites are located at the interface of the two subunits (63, 69, 135). The other transmembrane chemoreceptors have the same structure, although each periplasmic sensory domain is bound to a distinct ligand.

The membrane part of the receptors, called the transmembrane signaling domain, consists of four long helices, providing by two subunits. Several experiments have shown that the transmembrane signal is carried by the second transmembrane helix (TM2) across the bilayer (41). Recent genetic studies on the aspartate receptor have revealed that the signal is transmitted largely or completely within the subunit. In fact the cytoplasmic domain of the other subunit can be removed without blocking the signal (63, 69).

The cytoplasmic domains or methyl-accepting domains of the receptors are a dimer with 31 kDa periplasmic subunit and contain the covalent modification sites for receptor adaptation and a binding site that regulates the histidine kinase by binding to CheA. It has been proposed that the dimeric cytoplasmic domain consists of three distinct functional regions: a) the linker, b) the methylation region and, c) the kinase-signaling domain (53, 69, 198). The linker region couples the transmembrane signaling helix (TM2) to the cytoplasmic domain and it is important for signal transmission. The methylation region is composed of two methylation segments. Each monomer contains four to five glutamyl residues, which are specifically methylated by CheR and demethylated by a protein glutamate methyltransferase, CheB. The first segment contains the regulatory methylation sites targeted by the adaptation pathway. This segment is important for kinase regulation as it transfers the information from the periplasmic ligand-binding site to the kinase. The kinase-signaling domain forms a ternary complex with CheW and CheA and is responsible for regulation of the bound histidine kinase activity (63, 201).

1.3.2 The Histidine Kinase

CheA is a key member of the two-component chemotaxis system, integrating the signals of five transmembrane sensor proteins and controlling two response regulators. CheA is autophosphorylated at His48 (23) by signaling output of the receptor and then phosphorylates CheY (132).
The kinase is a dimer of identical 71 kDa subunits with 654 amino acids, containing two symmetric active sites. Each site utilizes Mg\(^{2+}\)-ATP to drive phosphorylation of His48 (195, 201). Each subunit can be divided into four functional regions (63, 157), the first two of which fold as independent structures:
(a) the phosphotransfer P1 domain, which possesses the reactive His48 residue that serves as the site of autophosphorylation (85),
(b) the P2 response-regulator docking domain, which provides the binding site for CheY and CheB;
(c) the catalytic domain which is thought to contain all of the motifs needed to bind Mg\(^{2+}\)-ATP, recognize the P1 domain, and catalyze its phosphorylation;
(d) the C-terminal receptor docking region which is essential for receptor-mediated regulation. Genetic studies suggest that distinct CheW- and receptor-binding sites exist within this region of CheA. The conserved C-terminal of CheA is not required for protein-protein interaction between the histidine kinase and response regulator components, nor is this region required to catalyze the transfer of phosphoryl groups from histidine to aspartate chain. Rather, the conserved kinase domain probably defines the active site for ATP binding and histidine phosphorylation.

The activity of the kinase can be increased when it interacts with the receptor and CheW.

There are two forms of CheA; the full-length molecule having 97 additional amino acids in the N-terminal domain (P1 domain) and a short form called CheAs that lacks the P1 domain and cannot be phosphorylated (157, 201).

1.3.3 The aspartate kinase, CheY

In the two-component signaling pathway, the histidine kinase acts as a signal transmitter and the aspartate kinase serves as a receiver (157, 201). The receivers, including CheY, have an approximately 110-amino acid domain, referred to as a "receiver module". This domain contains the Asp-phosphorylation site.

The CheY is a 14 kDa monomer protein with 128 residues. It has two functions: as a receiver for signals from CheA by phosphorylation, and as a response regulator for motor switching. The phosphorylation site in CheY is Asp57 (170). CheY shows a great affinity for CheA when it is unphosphorylated, whereas
phosphorylated CheY exhibits a decreased affinity for CheA but a significantly increased affinity for the motor component FliM and the phosphatase CheZ (221). Interaction between CheY and FliM of the motor switch leads to a change of flagellar rotation from CCW to CW. Activation of CheY by phosphorylation generates a global conformational change that regulates the multiple docking site of CheY (15, 16, 66). These multiple docking sites interact with each of three effector proteins; P2 domain of CheA, FliM (222) and CheZ (28). The binding of Mg$^{2+}$ carries out significant conformational changes in CheY leading to modification of the surface domain which interacts with the switch. Mg$^{2+}$ is essential for CheY phosphorylation, dephosphorylation (CheZ dependent) (28, 195), and CheZ binding to CheY, however the CheY-P-FliM is Mg$^{2+}$ independent. Phosphorylation of CheY leads to its dissociation from the quaternary complex, receptor-CheA-CheW-CheY (201).

1.3.4 Other proteins involved in chemotaxis

There are other proteins involved in the chemotaxis response besides the two-component regulatory proteins. They act at different levels of the system and include CheW, CheR, CheB and CheZ.

1.3.4.1 CheW coupling protein

CheW is a 18 kDa coupling protein which has no regulatory or catalytic function, binds the CheA to the receptor enhancing the activation of the histidine kinase, CheA (50, 76). CheW serves, as a simple scaffolding protein and it is important for CheA function. However the specific site or residues, which are involved in receptor and kinase docking have not yet been identified. Kinase activity requires the coupling CheW, in contrast, kinase inhibition is CheW independent.

1.3.4.2 CheR methyltransferase protein

The CheR methyltransferase a 30 kDa soluble enzyme, which exist free in the cytoplasm, modifies the receptor cytoplasmic domain. It is a key factor in the adaptation phase of chemotaxis. In the cell, CheR is always found associated with the
cytoplasmic portions of the receptor complex. CheR binds to aspartate, serine (229), and citrate receptors and generates the receptor-kinase signaling complex. The receptor site which binds to CheR consists of four to five-residue sequences and located at the extreme C-terminal end of the receptor cytoplasmic domain, however the ribose/ galactose and dipeptide receptors lack this docking sequence and are hardly methylated in the absence of the aspartate or serine receptors (78).

CheR, which utilizes S-adenosyl-methionine as the methyl donor, consists of two domains. The active site cleft located between the two domains.

### 1.3.4.3 CheB methylesterase protein

The methyl esterase role in chemotaxis adaptation is played by CheB. This protein has two different activities (amidase and response-regulatory function), which are carried out by its different termini. The C-terminal methylesterase domain of CheB functions as an esterase to hydrolyse glutamylmethylester groups in the membrane receptor proteins (194). In some cases, CheB can hydrolyse the receptor methylation sites which are expressed as glutamines, and produces a bare glutamate side chain that is then available for methylation (69).

A linker to the C-terminal domain joins the N-terminal domain of CheB. The N-terminal domain is homologous to CheY, containing the site of phosphorylation that can be phosphorylated by transferring the phospho-His48 of CheA to its own active site similar to those of CheY (63). It seems that N-terminal domain regulates C-terminal methylesterase activity by blocking its active site physically. However, when CheB phosphorylated by CheA, generating a conformational change, this inhibition is removed, allowing CheB-P to interact with MCPs. Phospho-CheB dephosphorylates much faster than phospho-CheY by auto-hydrolysis reaction. This is because, unlike CheY-P, the phospho-CheB molecule is not dephosphorylated by CheZ (85, 194). Phosphorylation of the CheB receiver domain increases the methylesterase activity of the CheB.

### 1.3.4.4 The CheZ protein

CheZ protein is a component of the chemosensory pathway that is less understood. Homologues of CheZ have not been found in species outside the α-
subgroup of proteobacteria (8). It serves as a phosphatase to speed up dephosphorylation of CheY and thus inactivating the tumble signals (27). CheZ can dephosphorylate CheY-P with a high specificity, either directly using a CheZ site residue or indirectly via a conformational change which stimulates autophosphatase activity of the CheY active site (69, 85). The phosphatase activity of CheZ is essential for rapid response to attractant and suppressing tumbling of bacterial cell. However, little is known about how CheZ, itself is activated.

CheZ has been isolated as a hydrophilic dimer of 24 kDa subunits. Recently a distinct functional domain within CheZ has been found. This domain contains 19 residues of the C-terminus which form part of the CheY-docking site (27). There is another possibility that the N-terminal region of CheZ may also contribute to the CheY-docking surface (173).

1.3.5 Signal transduction in bacterial chemotaxis

Most chemical stimuli are detected by specific receptors. Some chemicals firstly interact with a binding protein, and the complex subsequently interacts with a transmembrane receptor, however other stimuli are directly detected by receptors themselves.

There are two kind of chemotaxis in *E. coli*. In metabolism-independent chemotaxis, the signal is generated from a change in the concentration of molecules that binds to the receptors rather than their absolute levels (112). In metabolism-dependent behaviour, the binding molecules must be metabolised to produce a signal. The term "chemotaxis" usually refers to metabolism-independent behaviour, whereas metabolism-dependent system is used for "energy taxis" which is responsible for monitoring the change in cellular energy resulting from metabolism of the signaling molecules. The metabolism-dependent system include basic requirement for bacterial growth, such as oxygen and other acceptors, light, most sugars and a range of electron donors (7, 8).

The chemotaxis can be divided into two parts: a) the excitation phase and, b) the adaptation phase. The excitation process includes all activities which transduce the initial signal from the receptor to the motor reversing the direction of flagellar rotation. This process operates through a phosphorylation cascade. The adaptation process which follows the excitation phase, restores the flagellar rotation to
prestimulus levels, even in the continued presence of the stimulus. This process operates via a methylation mechanism.

a) Excitation phase

The nature of the signals generated by receptors and subsequently transduced was discovered approximately 10 years ago. It was shown that the signals were not electrical, but rather chemical. Chemotactic responses begins when membrane receptors, which are clustered at one of the poles, bind directly, (or indirectly with periplasmic binding proteins) to specific compounds. Binding sites for attractants or repellents and binding proteins are distinct. A single receptor sends an additive signal if more than one compound is bound. Thus maltose can block tumbling even if the aspartate binding sites on the Tar receptor are occupied. Similarly the repellent response to phenol mediated by another receptor, is abolished with addition of ribose (69).

Binding proteins contain two domains. Ligand binds in the cleft between the two domains. The cleft is open in an unoccupied state but closed when bound to a compound. Binding of a ligand to the cleft generates a contiguous docking surface and conformational changes that activate the receptor binding sites.

The sensory domains of the receptors are located in the periplasm. They recognize binding proteins or small molecule ligands. It is thought that the chemotaxis receptors transmit the signal as an intra-dimer conformational change. The second transmembrane helix (TM2), of the transmembrane domain of the receptor transduces the signal across the bilayer via the linker of the cytoplasmic region domain (41). This domain binds and regulates CheA. Recent genetic studies showed that transmembrane signals are generated completely within one subunit.

The homodimeric C-terminal domains of the receptors contain the covalent modification sites responsible for receptor adaptation, and the docking site that regulates the CheA histidine kinase. The latter site of the cytoplasmic domain of the receptor forms a ternary complex with CheW and CheA. This region is an independent functional unit containing the minimal receptor sequences required for ternary complex and activation of the kinase.

CheA is an autophosphorylation histidine kinase. It exists as a dimer, containing two symmetric active sites, which utilize Mg^{2+}-ATP to drive phosphorylation of His48. The monomeric form of CheA is not active in the
laboratory; dimerization of CheA is therefore thought to enhance kinase activity, in which one monomer phosphorylates the other (203). Phosphorylation of CheA is increased when coupling protein CheW joins the CheA-receptor complex forming a large complex comprising six subunits. Genetic studies suggest that CheW directly binds receptor during the excitatory phase (116), however CheW can form a complex with CheA which increases the rate of phosphorylation and affinity for CheY (76, 134).

The kinase uses ATP to phosphorylate itself at a histidine residue. The autokinase activity is reduced when attractant is bound to the ligand-binding receptor and increased when no attractant is present. Repellents stimulate the histidine kinase activity and speed the production of CheY-P, whereas attractants inhibit the active ternary complex and slow phospho-CheY formation, thereby raising or lowering the steady state tumble signal, respectively.

CheA provides the docking sites for CheY and CheB. These binding sites, which are located in the P2 domain of CheA, allow transfer of the phosphoryl group from the histidine kinase to the aspartate kinase protein CheY or CheB. Transmembrane signaling occurs solely within the complex is composed of a receptor dimmer, CheA dimmer and two molecules of CheW.

CheY acts as a phosphorylation-activated switch that controls the direction of flagellar rotation. As a result it is known as the tumble signal (176). Its action is modulated by two opposing reactions: creation of CheY-P by the receptor-kinase complex, and destruction of CheY-P by hydrolysis of its phosphate. CheY binds the receptor-CheA-CheW complex by docking to the P2 domain of CheA. Transfer of a phosphoryl group then occurs, from CheA to Asp57 of CheY. Mg$^{2+}$ is required for phosphoryl transfer to CheY (113, 195). Phosphorylation of CheY results in a significant conformational change (18) that affects its interaction with other proteins, dissociating it from CheA, while facilitating binding with CheZ. CheZ is an auxiliary protein which accelerates the dephosphorylation of CheY (27). CheY-P binds to and regulates the flagellar motor switch (15, 16), altering flagellar rotation from the default direction of CCW to CW rotation, thereby favoring tumbling. CheY-P is therefore called "tumble factor". Mutation studies suggest that tumbling is modulated by a specific conformational change at the motor-docking surface.
After dissociation of CheY-P from the complex, the rest of the complex, containing phospho-CheA, can also phosphorylate the CheB methylesterase, thus controlling the adaptation phase of the system.

Dephosphorylation of CheY and termination of the CheY-switch interaction is enhanced by CheZ. Interaction between CheZ and CheY-P is higher than CheY itself and the binding of CheZ to both CheY and Mg\(^{2+}\) enhances CheY-P as well. Phosphorylation and dephosphorylation of CheY requires the presence of the cation Mg\(^{2+}\). It has been proposed that CheZ, rather than catalyzing the hydrolysis of the CheY-P, acts allosterically to stimulate the auto-dephosphorylation activity of CheY-P. Some studies suggest that CheZ is not always necessary for chemotaxis to occur, and the requirement for CheZ depends on the attractant gradient background of the bacteria. Although phosphorylation increases binding of CheY to the switch, some observations have indicated that CheY phosphorylation alone is neither sufficient nor essential for binding to the motor switch. All of the current evidence is consistent with the view that CheY phosphorylation induces a conformational change that causes CheY to bind to flagellar motor proteins and thereby induce a tumble response. It has been proposed that for the generation of CW rotation an additional cytoplasmic component, which is not a chemotaxis protein, is required. Bourret et al. suggest that switching from CCW to CW involves at least two steps which CheY phosphorylation is the first step (184). The second step is possibly carried out by lys-109 of CheY in an event following phosphorylation.

The chemosensory pathway is extraordinarily sensitive. As little as a single attractant molecule can trigger a detectable motor response. Each *E. coli* cell contains approximately 25000 receptor monomers, 2500 molecules of CheA and equivalent amount of CheW, and 12000 molecules of CheY (200). Thus cells can exhibit a significant response when attractant ligands bind to only a few receptors. The motor is extremely sensitive to very small fluctuations in CheY-P.

b) Adaptation phase

In most signal transduction pathways each reaction has a counter reaction. These activities that lead to prestimulus behaviour, have termed adaptation, habituation, etc.

The adaptation part of the chemotaxis system adapts the cell to a stimulus concentration so that the system can react even in a large amount of attractant or
repellent. When an attractant binds to a receptor, it generates negative signals as a conformational change that cross the membrane and block CheA activity, thereby decreasing the CheY-P level. This process causes the cells to tumble less and increasing their run lengths as they enter areas of higher attractant concentration. Two enzymes modulate responding to increasing attractant concentration, the methyltransferase CheR and the methylesterase CheB. The first level of adaptation occurs at the receptor via a methylation process. The receptors’ glutamyl residues in the cytoplasmic domain, are methylated by CheR and demethylated by CheB (229, 230).

Figure 1.3.5.1 Diagram showing main steps of the chemotactic response in *E. coli*. Figure adapted from (207)
Methylation of the receptors counterbalances the effects of ligand binding that diminishes the response in the presence of a constant stimulus.

CheR binds tightly to the C-terminus of some receptors (aspartate, serine, and citrate receptors), and the glutamate residues of MCPs are methylesterified via S-adenosylmethionine as the methyl donor (43). The resulting methylesters, which increase the kinase signal activation of the receptor, are hydrolyzed by the action of CheB (164). CheB is a target for phosphorylation from phospho-His48 of CheA and CheB-P removes methyl groups from the methylated receptors. In steady state, methyl addition by CheR balances methyl removal by CheB-P causing run-tumble behaviour. The feedback control of the adaptation system is generated by the receptor-kinase complex, which phosphorylates and activates CheB. CheR appears to work at a constant rate, whereas the activity of CheB is regulated by phosphorylation (7). When an attractant binds to a receptor and inhibits CheA activity, the level of CheB-P drops due to autohydrolytic dephosphorylation, resulting in a gradual increase of CheR activity. Eventually the rising level of methylation will stimulate CheA activity, thereby restoring the ability of the receptor to stimulate CheA and bring the signal to its basal level. Thus the adaptation phase of chemotaxis, exactly balances the signals generated by multiple attractants and repellents.

Attractants increase the methylation levels of the MCPs. A highly methylated transducer signals the motor to bring about CW rotation. Thus following the excitatory signals, increased levels of methylation return the receptor to a null signaling state. Conversely, repellent adaptation causes loss of methyl groups, with unmethylated receptors sending signals for CCW rotation. The level of MCP methylation also provides a simple chemical memory to find out whether the current direction of swimming is favourable or not. This process works based on a time difference between fast MCP conformational changes and relatively slow methylation changes after binding of ligands. The methylation level depends on the attractant or repellent concentration. For example the methylation level is high if the concentration was high. When the cell swims, it compares the current chemical environment, as monitored by the ligand occupancy of the receptor population, to the chemical environment of the recent past, as remembered by the methylation level. Thus if a bacterium with transducers in the adapted state moves in a long run down an attractant gradient, the still methylated, but no longer ligand bound receptor biases the flagella rotation and produce tumbling. Only if no more attractant molecules bind, to
the proteins become demethylated and return to the unstimulated state. For more reviews see (3, 9, 63, 69)

1.3.5.2 The Phosphotransferase (PTS) system

Prokaryotes such as *E. coli* possess a variety of systems besides the basic chemotaxis pathway, which contribute to the control of swimming behaviour. One of such systems is the phosphoenolpyruvate (PEP)-dependent phosphotransferase system, responsible for detection and translocation of a large number of carbohydrates, such as glucose and fructose, that can also act as chemoattractants (160). This system is thought to be methylation-independent and not carried out by the MCPs. There are several enzymes, EI, EII, and EIIC involved in translocation and phosphorylation of carbohydrates. Phosphorylation of compounds depends on enzyme phosphorylation by a pathway involving transfer of a phosphoryl group from PEP to the enzymes. PTS phosphate moves from PEP to enzyme EI, to HPr (histidine protein) to an EII/ EIIC complex and finally to the sugar.

The system requires the two-component proteins CheA, CheW, and CheY (168). It has been proposed that this system is integrated with the MCP pathway at the point of CheA (120), by phosphorylation of EI enzyme. Phospho-EI inhibits CheA autophosphorylation and hence also CheY, leading to the chemoattractant swimming response.

In spite of the importance of this system for bacteria, the chemotaxis response to PTS attractants is much weaker than chemotaxis that is carried out by the MCPs (220).

1.3.5.3 Aerotaxis/ Energy taxis in *E. coli*

An additional receptor has been identified in *E. coli* that responds to oxygen and a variety of other stimuli which cause changes in the energy (redox) levels in the bacteria. This receptor, referred to as Aer (since it is responsible for the aerotactic response) (22), can also activate flagellar rotation. It has been shown that Aer can sense redox potential and internal energy in the cell, in addition to external oxygen (78, 207).
The Aer receptor is located at the cytoplasmic surface of the bacterial membrane. It acts as an intracellular sensor with two cytoplasmic domains but no periplasmic sensing domain. The second cytoplasmic domain is thought to be the sensing domain. The carboxyl-terminal domain of Aer is very similar to that of a classic MCP. In contrast the amino-terminal domain looks like the redox-sensing domain.

Aerotaxis has been shown to require CheA, CheW, and CheY to function (168). Aerotaxis can also integrate with chemotaxis through the Tsr receptor. This protein is able to detect the cytoplasmic pH and periplasmic pH. It is thought that Tsr can mediate aerotaxis by detecting changes in the proton motive force of the cell.

Unlike most chemotaxis system, which have been shown to be metabolism-independent, energy taxis is metabolism-dependent.

1.3.6 Other factors regulating chemotaxis

There are some factors that can activate the chemosensory system in addition to the attractants and repellents described earlier, without the need for interactions with the MCPs. It is likely that these alternative pathways connect to the MCP system at some point (95, 131). However the exact mechanism of their effects are still not clear.

Intracellular Ca\(^{2+}\) levels

Increased levels of free cytoplasmic Ca\(^{2+}\) cause increased tumbling frequencies in chemotactically wild-type *E. coli*. This effect requires CheY, CheA, and CheW (212). The precise mechanism of action is unknown. Ca\(^{2+}\) may stimulate CheY or CheA phosphorylation or it may stabilize CheY-P.

Intracellular fumarate level

Fumarate has also the ability to act as a prokaryotic switch factor. Recent studies have shown that fumarate restores the ability to spontaneously switch the direction of flagellar rotation (139). No cytoplasmic constituents other than CheY are required for the fumarate effect. Malate, maleate and succinate also have the fumarate effect but to a lesser extent. Fumarate might be the connecting point between the metabolic state of *E. coli* and the chemotaxis system.
The acetate effect

Acetate acts as a repellent that is able to cause tumbles in *E. coli*. This is called the "acetate effect". The acetate metabolism pathway uses phosphotransacetylase and acetate kinase. The intermediate molecules in this process are acetyl phosphate and acetyl adenylate, shown to phosphorylate or acetylate several response-regulators, including CheY, which then brings about CW rotation (163). This phosphorylation event could therefore explain the connection between acetate and chemotaxis.

1.3.7 Chemotaxis in other bacteria

Chemotaxis in most prokaryotes follows the same fundamental mechanism as that used in *E. coli*. However there are also some differences between the systems, some of these will be referred to as an example.

Chemotaxis in *Bacillus subtilis* differs from that in *E. coli* in several ways. *B. subtilis* has three additional che genes (*cheC, cheD, and cheV, a cheW-cheY fusion*) but lacks *cheZ* and free CheY (7, 152). *cheZ* has only been found in enteric species. In addition, in *B. subtilis*, attractant stimulates CheA kinase, and CheY-P causes CCW flagellar rotation to mediate smooth swimming (25), which is in contrast of *E. coli* response. Both attractant and repellents stimulate hydrolysis of methyl groups in *B. subtilis* (7, 143). The role of methylation in adaptation is also different from that in *E. coli*.

Non-enteric bacteria are thought to possess more complicated sensory systems than those of enteric bacteria. Such differences may reflect the more-complex environments in which non-enteric organism live and the greater flexibility of their metabolism.

The α-subgroup of Proteobacteria including *Sinorhizobium meliloti* and *Rhodobacter sphaeroides* have a motility and chemotaxis system that are different from those of enteric bacteria. In these bacteria, at least two distinct chemosensory regions are involved in chemotaxis. In *Rhodobacter sphaeroides* together, there are two copies of CheA, CheW (81, 82) and CheR, at least three CheY and one of CheB, as well as five orfs of unknown function and three Tlp (transducer-like protein). *R. meliloti* possesses three motility and chemotaxis regions, which contains one copy of *cheA, cheW, cheD, cheR and cheB*, two of *cheY*, one unknown orf and one *tlp*.
Interestingly, in both bacteria CheZ is absent but at least two copies of the response regulator CheY are present (11).

These bacteria can vary their swimming speed during chemotaxis. It has been suggested that the two CheY homologues in Sinorhizobium control the speed of flagellar rotation. The S. meliloti flagellum neither stops nor switches direction, but changes its rate of rotation. Mutation studies have been shown that CheY2 slows motor rotation in response to negative chemotactic stimulation, thereby changing the swimming direction of the cell. CheA phosphorylates the two CheYs (189). This new mechanism seems to apply to the other members of the α-subgroup of Proteobacteria.

1.4 Motility in E. coli and S. typhimurium

Motility and movement of bacteria is an essential complementary action for chemotaxis. Motility is more valuable outside the host and less so within. Just as with chemotaxis, the best understanding of bacterial flagella, their structure and function, is in the closely related enteric bacteria E. coli and S. typhimurium.

E. coli is a motile bacterium, capable of movement in a liquid medium by means of flagella. However, it should be recognized that flagellar motility is not the only form of movement in bacteria. E. coli has typically 6-8 flagella arising from random points on the surface of the cell. Flagellation in E. coli and S. typhimurium is random with respect to both number of flagella and sites of origin. Such cells are described as peritrichously flagellated. The pattern of flagellation also varies considerably among species. Movement of bacteria is carried out by the rotation of flagella, which can rotate in either direction, counterclockwise (CCW) causing a run or clockwise (CW) producing a tumble. Bacterial flagella are rotated by reversible motors, located at the base of each flagellum, in the cytoplasmic membrane. The transmembrane gradient of protons (protonmotive force, PMF), or in some species, sodium ions is used as the energy source for movement (21). The flagellar motor, in this aspect, is a device that converts chemical energy stored in the membrane ion gradient into the mechanical work of rotation. The mechanism of this energy conversion is not yet understood.

The flagellar motor continuously switches between CCW and CW rotation. The default direction of the flagellar motor is CCW. In counterclockwise rotation,
flagella aggregate together, forming a bundle that pushes the bacterium forward. When one or more flagella switch to CW rotation, the bundle falls apart and thus the bacteria tumble. The tumble is a result of transition of some flagella from one form to another. The motor has also been shown to pause, which is thought to be a result of incomplete switching events. For more detailed review of flagellar structure and organisation see (122).

1.4.1 Flagellar Structure (21, 122)

The *E. coli* flagellum morphologically consists of a thin helical tube (20 nm in diameter), called the filament, which is joined by accessory proteins to a short curved segment named the hook. This, in turn is connected to a complex structure at the base of flagella consisting of several rings mounted on a rod, termed together the basal body. They are thought to generate flagellar rotation and act as a motor.

The filament, which is rather rigid, comprises of several thousand copies (About 20000 monomers) of a single protein called flagellin. This protein is encoded by *fliC*. The length of each filament varies and can be up to ten times the cell body length (up to 20 μm). The long helical filament which acts as a propeller when rotate, is known to have a central channel (25-30 Å diameter) that flagellin monomers can be exported through for filament assembly (144). The tip of the filament is covered by the product of *fliD* (HAP, hook-associated protein).

The filament is connected to the cell by the hook, which is structurally similar to the filament but contains different subunits. The hook protein is composed only of *flgE* gene product. Between hook and filament are two junctions or hook accessory proteins, FlgK and FlgL (HAPs) (87). The hook is believed to act as a flexible, or universal joint between filament and cell. Hook length is controlled by a mechanism that involves FliK protein. Recently, it has been proposed that a structure in the basal body, C-ring is the element that may control the hook length, rather than FliK. There is a possibility of involvement of other proteins such as FlhB in controlling hook length. The hook also has a central channel, like the filament, allowing the export and assembly of hook and filament subunits.

The hook is connected to a structural complex known as the basal body, which is mainly located within the cell envelope. The basal body is seen to be of cylindrical symmetry, consisting of a rod joined to the hook and passing through a set of four
The rings are named according to their position in the cell. The two outer rings are called L- and P- rings, and the inner two rings named S- and M- ring. These two rings make a double ring, called together the MS-ring, which is made up of a single protein encoded by \textit{fliF}. The basal body is thought to function as a mechanical motor, which the MS-ring acts the rotor, the rod as the transmission shaft and the outer membrane rings are the bushing.

Recent studies showed that the rod has a helical symmetry, structure similar to that of the hook and filament, and it has the hollow central channel required for protein exportation. The channel is made from axial proteins, which form the rod, hook and filament. The rod is composed of four proteins, FlgB, C, F and G. The rod contains about 6 copies of each FlgB, C and F and about 26 copies of FlgG. The FliE protein, a component of the basal body, has also been found to be required for assembly of the rod. There are other proteins that are thought to involve in this process, causing the flagellum-specific export apparatus to function correctly. Interestingly, the proteins responsible for export and assembly of the external components of the flagella are closely related to proteins responsible for export of virulence factors (see section 1.4.3).

The L- and P- rings are made up of proteins FlgH and Flgl respectively. These two proteins form a large pore in the periplasmic space and outer membrane, through which the rod passes. The P-ring lies in the peptidoglycan layer and L-ring is embedded in the outer membrane.

The last part of the flagellum is the switch complex, which consists of a rotor and a stator. The rotor is a large ring- or bell-shaped structure, called the C-ring (C as cytoplasmic), which is attached to the cytoplasmic face of the basal body. The C-ring is composed of three switch-complex proteins, FliG, FliM and FliN, known to be important for flagellar assembly, rotation and direction control. It is thought that FliG is attached to the cytoplasmic face of the MS-ring, while FliM and FliN form the C-ring. It is still not clear whether they are a part of the rotor, or stator of the flagellar motor. FliM is exposed to the cytoplasm and is connected to both FliN and FliG. The latter connects FliM to the basal body of the flagella. The precise composition of the C-ring is unknown, but it is quite large and would contain up to 100 copies of FliM and FliN. There is a possibility that the C-ring also contains other proteins in combination with FliM and FliN. The precise roles of the three switch proteins are still unclear. FliM consists of four functional regions involving in interaction with
CheY-P, FliG, and regions used for switching and assembly of the flagella and a C-terminal required primarily for assembly (213). The amino terminus of FliM is the CheY docking site. The motor switches when CheY-P binds to FliM, generating CW

Figure 1.4.1 Diagram showing the general structure of the *E. coli* flagellum.
Figure adapted from (123). The position of the C-ring is indicated by the switch complex.

HAPs= hook-associated proteins
rotation. The processes occurring within the switch after CheY binding are not known. Some observations suggest that the switch protein recognizes the conformation of CheY, rather than its phosphoryl group. Recent evidence has revealed an interaction between FliM and FliG indicating that the binding of these two proteins might be critical for switching.

The motor consists of two parts, rotor and stator. The FliG, FliM and FliN form the rotor, whereas MotA and MotB constitute stator. FliG directly participates in rotation as it rotates along with MS-ring. FliG can be fused to FliF, which forms MS-ring. This ring is attached to the filament and believed to rotate with it, so FliG is likely to rotate also.

There are two other motor subunits involved in power production. They are MotA and MotB that seem to be important for torque generation. They form a particular ring that surrounds the MS-ring. These two proteins are membrane bound, and function in transmembrane proton conduction. MotA is a proton channel, while MotB serves as an anchor, which connects the stator or non-rotating part of the motor to the cell wall. Mutation studies strongly imply that MotA and MotB interact with each other forming a complex and the presence of MotB is necessary for MotA function. The MotA and MotB complex forms a proton-conducting channel through the bacterial cytoplasmic membrane. Each motor can contain up to 8 independent MotA-MotB complexes (26).

In most species the periplasmic domain of MotB contains a sequence motif, which could be involved in binding to peptidoglycan. Interestingly, MotB of *R. sphaeroides* lacks this sequence, exhibiting instead a heptad repeat of histidine residues (153). These sequence differences might be related to the fact that the motors of *R. sphaeroides* rotate only in one direction, stopping occasionally but never reversing. In other species, the motors alternate between CW and CCW rotation. Recent evidence suggests that the electrostatic interaction between charged residues in the stator (MotA) and the rotor (FliG) elements of the motor might play a crucial role in torque generation.

In brief, there are currently 44 known flagellar genes in *S. typhimurium* and *E. coli*. Twenty three of these encode structural components of the flagellum. Five genes are needed for torque generation. The remaining components are the filament (propellor), the hook (universal joint), and the basal body. Another five flagellar genes have regulatory roles. The remaining genes (about 11 genes) encode the
component of the flagellum-specific export that are responsible for delivery of flagellar proteins to their destination (see section 1.4.3).

1.4.2 Chromosomal organization of the flagellar and chemotactic genes of *E. coli* and *S. typhimurium* (21, 26, 71, 122)

The products of over 60 genes are required for chemotaxis, and the assembly and operation of the flagella of *E. coli* and *S. typhimurium*. There are 15-17 operons coding for flagellum proteins, whereas a few operons coding the chemosensory components. Most of them are needed for flagellar assembly, several are involved in controlling the direction of motor rotation in response to chemotactic or other sensory stimuli, and a few are involved in torque generation. The majority of these genes are highly clustered on the chromosome.

The motility and chemotaxis genes of *E. coli* are arranged in three regions and in *S. typhimurium*, they are clustered in four regions. The regions are designated as I, II, III, and IV. Region III is further subdivided into two sections, IIIa and IIIb. Each region consists of up to 14 operons, which in turn contain between 1-9 genes. The genes are clustered in operons based on their similar function, or requirement at similar step in the overall process. The genes are designated according to the region of the chromosome where they are found, however some genes are also named based on their function in the chemotaxis and motility systems. The genes in region I are designated as *flgA-M* genes, region II as *flh, che* and *mot*, region III as *fliA-T* and in *S. typhimurium* region IV as *flj*. The genes that are named *fli, flg* and *flh* are involved in the flagellar structure or assembly. The *che* genes code for proteins in the central intracellular signaling pathway of chemotaxis, while *mot* genes are part of the motor complex of the flagella.

Expression of the operons is controlled by secondary sigma factors for RNA polymerase and takes place in a hierarchical manner. One operon to be expressed requiring the expression of the lower class operon. The operons are divided into four classes: class 1, 2, 3a and 3b. The lowest class in this regulatory hierarchy is class 1, which contains only 1 operon, called master operon coding FlhD and FlhC. The expression of this class is required before that of any other classes. The class 1 operon is under the control of cyclic-AMP (cAMP) via the regulatory protein CAP. The seven class 2 operons are directly under the control of the class 1 operon. Class II
operons contain a flagellum-specific consensus at \(-10\) position of their promoters (GGATAA). \textit{fliA}, a gene from class 2 encodes a sigma factor for classes 3a and 3b. The class 3a is controlled by the master operon (class 1) and \textit{FliA}, but the class 3b is solely controlled by \textit{fliA}. Class III promoters contain a specific consensus at the \(-35\) region (TAAA) in addition to the \(-10\) consensus sequence common to class II promoters.

1.4.3 Flagellar assembly and flagellum-specific export apparatus

The flagellum of \textit{E. coli} and \textit{S. typhimurium} is composed of four distinct parts, the motor complex, the basal body, which are embedded in the cell surface, the hook and the helical filament, both of which are external to the cell. The basal body consists of several structural subunits; called MS ring (the inner ring), P and L rings (the outer rings), the rod, and the flagellum-specific export apparatus. Biogenesis of the bacterial flagellum is thought to start from the most cell-proximal structure, the basal body, to the most cell-distal structures, the filament (122). It begins with the basal body formation. Once the MS ring is built, the rod, the P and L ring, the hook and the filament are constructed. It has been shown that flagellin monomers are added to growing filaments at their distal end (122). Modification of the flagellin subunits, or some other flagellar proteins, by glycosylation, acetylation, and methylation is required for correct assembly of the flagellin subunits into the filament. There are some experimental evidences indicating the export and usage of so-called flagellum-specific muramidase. Recent study has been shown that FlgJ protein from \textit{S. typhimurium} has such activity. This protein is the flagellum muramidase which hydrolyses the peptidoglycan layer to assemble the rod structure in the preplasmic space (145). The function of this enzyme is to punch a hole in the peptidoglycan layer in the early stage of flagellar morphogenesis, in order to allow penetration of the nascent rod (137). Flagellum assembly also requires the function of an anti-sigma factor, FlgM and a protein that controls the length of the flagellar hook, \textit{flgK} (71, 122, 204). FlgM binds to FliA, a specialised \(\delta\) factor, acting as an anti-\(\delta\) factor that prevents the binding of RNA polymerase \(\delta^{28}\) to class III promoter. This will inhibit the expression of class III operon that contains genes necessary for filament assembly before of the hook completion. Once the hook has been assembled, the FlgM will disassociated from FliA and secreted by flagellum-specific export apparatus to the outside of the cell, resulting of FliA activity and filament assembly.
The flagellar proteins, which are synthesised in the cytoplasm, have to cross the outer membrane to reach their destinations. There are at least nine structures that are located beyond the cytoplasmic membrane, and their proteins should be delivered to their locations. It has been proposed that the central channel existing within the growing flagellar structure is the physical pathway of protein transportation (107, 121, 122). It is thought that the proteins are exported to the periplasm, the outer membrane, or the extracellular space, where their assembly finally occurs by two distinct export mechanisms. One, which is more common, called type II sec-dependent pathway or GSP, is used for export of proteins with cleavable signal sequences (121). Only the P and L rings (FlgI and FlgH, respectively) have been found to be synthesised as proteins with cleavable sequences (161), and hence are transported across the cytoplasmic membrane, to the periplasm by the protein export system. The other mechanism, which is unusual, is called Sec-independent export pathway or typeIII flagellum-specific export pathway/apparatus (71). The distinct feature of this export pathway is that it transports proteins, which lack a cleavable signal sequence necessary for transportation with the general pathway. Twelve proteins of the filament, rod, hook, hook-associated proteins, basal body (FlgB, FlgC, FlgE, flgG, flgE, FlgK, FlgL, FliD, FliE and FliC), and FlgM, the regulatory protein (anti-sigma factor) do not carry amino-terminal leader peptides signal and are thus exported from the cytoplasm to their final destination via the flagellum-specific export pathway. Since most components of the flagellum are outside the cytoplasm and do not have signal sequences, the function of the export apparatus plays an important role in flagellar assembly. It has been shown that this system contains at least eight general components, which could rise to 13 if the genes with specific functions such as chaperones are included. These eight general components, identified by analysis of various conditional and knockout mutants, are flhA, flhB, fliH, fliI, fliO, fliP, fliQ and fliR without which no export of any kind can occur. All components of the flagellum-specific export pathway are conserved among different bacteria. Two proteins, FliJ, a cytoplasmic protein and FliE, a basal body component, were found to be necessary for export of at least some substrates such as FlgD (hook-capping protein) and FlgE (hook protein) (122, 137). FliJ seems to act as a general component of the typeIII flagellar export apparatus, rather as a specific Chaperone (138). It is believed that in a short period of time, the L ring (FlgI protein) alone or as a complex L ring-FlgH is also required for export of proteins, since a temperature-sensitive fliN mutant cannot
export flagellar proteins (138). FlhA, FlhB, FliO, FliP, FliQ and FliR are all integral membrane proteins, three of which are physically associated with the flagellar basal body. The proposed location of the membrane-associated components is in a central pore within the MS ring of the flagellar basal body. There is physical evidence that such a pore exists (138) and it has been shown that FlhA, FliP and FliR are associated with the basal body (137). The MS ring and C rings are also required for export of all proteins as they provide a physical support structure for the export apparatus, but they are not directly involved in export. The biochemical functions of most of these genes are not yet known. One member of this system, FliJ, was found to have sequence similarity to subunits of the proton-translocating FoF1 ATPase, and is believed to supply the energy necessary for export.

It has been suggested that export proteins, with the aid of cytosolic chaperones such as FliJ or FliS, are brought into contact with two soluble components, FliJ (an ATPase) and FliH (function unknown) to form FliH-FliJ-FliJ-protein complex (138). They deliver the proteins in an energy-dependent process to the soluble domains of FlhA and FlhB and release their substrates. FlhA and FlhB have large soluble domains that protrude into the cytoplasm within the cavity that lies at the centre of the cytoplasmic C ring of the motor. The substrates are then translocated across the plane of the membrane through a complex of FliO, FliP, FliQ, FliR and the transmembrane domains of FlhA and FlhB. At the end the translocated substrates cross the channel and assemble at its destination. The export apparatus will not export filament-type proteins until it has received a signal to switch its specificity, and this signal is generated only upon completion of the hook. The switch in export specificity is mediated by FlhB, by blocking the gate for the hook protein, but opening it for the filament protein, along with FliK that controls the hook-length.

One regulatory feature, FlgM, an anti-sigma factor which binds to FliA to prevent its association with RNA polymerase enzyme (150), is exported from the cell by the typeIII flagellum export apparatus.

Members of the flagellum-specific export pathway have similarity to components of a newly identified protein export system called the typeIII secretion pathway or contact-dependent system, which is responsible of the export of a variety of virulence factors, by pathogenic bacteria, into plant and animal host cells. For instance, FliJ was found to have homology to Spa47 of Shigella flexneri and HrB6 of X. campestris, FlhA to MxiA of S. flexneri, InvA of S. typhimurium and LcrD of
Yersinia pestis, FliR to MopE of Erwinia cartovora and SpaR (Spa29) of S. flexneri. The typeIII secretion system is a protein delivery machinery that enables bacteria to translocate pathogenicity proteins from the cytoplasm across the inner and outer membrane, into the plant and animal cells. It is composed of approximately 20 proteins, most of which are located in the inner membrane and build a large structure that spans both bacterial membrane and possibly the host cell membrane as well. The homology between members of two systems should not be surprising, since both systems are responsible for transportation and localization of proteins across the cell.

Figure 1.4.3.1 Diagram showing Hypothetical structure of typeIII Flagellum-specific export apparatus. Figure adapted from (138)
membrane. It has become clear in recent years that the flagellum-specific export apparatus and the typeIII secretion system have common characteristics (92, 138). The two systems constitute a superfamily of proteins that share similar overall structural features, but are involved in various cellular processes including virulence in Yersinia pestis, host invasion in S. typhimurium and Shigella flexneri, and flagellar assembly in E. coli and C. crescentus. The proteins exhibit several putative transmembrane domains in the amino-terminals, suggesting that they are membrane integral proteins. The proteins are exported by these specific pathways also share several characteristics. For example, harpin (Hrp), Yop, Ipa, Inv and flagellar proteins are all cell surface-associated and extracellular components. In addition, they do not contain the conserved N-terminal signal sequence characteristic of proteins secreted by the Sec-dependent pathway of Gram-negative bacteria. It is thought that a typeIII secretion signals is located at the amino terminals of the both flagellar and virulence proteins (19). It has been proposed that such a signal resides in the 5’ end of the protein mRNA (6). A second secretion signal has been described in YopE which is dependent on interaction with its chaperones (19). The conservation between genes of typeIII secretion mechines and flagellar-specific export complex compelling and speculate that latter may also recognize mRNA signal to export flagellar units across the bacterial membrane (5).

Some lines of evidence suggest that the virulence export pathway may also be used for export of flagellar components, since a mopE mutant, a fiIR homologue in the typeIII secretion system which is responsible for export of virulence factors in Erwinia carotovora, failed to assemble the flagella, in addition to a reduction in virulence level in plants. In the typeIII secretion system, a large homomultimeric complex, made of molecules called secretins, is found in the outer membrane. It is thought that the function of this complex is to act as a passage for export of substrates across the outer membrane. A needle-like structure with a hollow projection extending out from the bacterial surface, that contains a known component of the typeIII export system, and closely resembles a flagellar basal body, was recently found in Salmonella (106). This needle complex structure may play the same role as the basal body, in the typeIII secretion system. It has been postulated that the L-ring of the flagellar basal body may transiently correspond to a secretin-like structure that links the two systems of export of flagella and virulence proteins (137). It has recently been revealed that flagellar export machinery, like typeIII system has chaperones. The role
of chaperones is directly interaction with secreted proteins and is responsible for preventing folding or aggregation. Chaperones might have dual roles in the cytoplasm. They may preventing premature association of their target substrates, and also involved in substrate recognition by the export systems in the membranes (19, 119). FlgN, FliT and FliS have been found to facilitate export of flagellar proteins such as FliC and have chaperone-like functions. These evidences are strengthening the idea that typeIII flagellar export system and typeIII virulence secretion system closely related. In fact, the flagellar export apparatus is a sort of typeIII secretion pathway, differing only in the nature of its export proteins. It seems possible that the origin of the typeIII secretion system may have come from the flagellar protein export system, since flagella are very ancient organelles. Perhaps in evolution the flagella-specific export pathway has been changed for the elaboration of some pathogenicity factors in bacteria. For more review about relation between flagellar- specific export pathway and typeIII see (19, 92, 119). Interestingly, some function and structural similarities have also been found between the typeIV DNA-transfer system which is responsible to deliver Agrobactrium T-DNA into host cells, and typeIII secretion system (46). VirB10 and VirB11 are homologues to members of various bacterial systems for transport of virulence proteins (89, 239, 240). Homologies were also reported between VirB4 and FliI proteins from flagellar export system as both are ATPase. These evidences suggest that T-DNA transport system may have evolved from a protein transport system.

Many components of the flagellum-specific export pathway have also been found in A. tumefaciens and the closely related bacterium Sinorhizobium meloliti. These genes have sequence similarity to members of the system in many other gram-negative and Gram-positive bacteria, and different genes of typeIII secretion systems of Gram-negative bacteria. However, the order of the export genes in A. tumefaciens and S. meloliti is different from those in E. coli, C. crescentus and B. subtilis. Surprisingly, genes controlling cultivar-specific nodulation of soybean by S. fredii have been found to have sequence similarity to the genes encoding the typeIII secretion system in other bacteria. Homology between symbiotic genes and typeIII secretion proteins (73), and similarity between the flagellum-specific export pathway and the typeIII system, reinforces the idea that these three systems, symbiosis, pathogenesis and flagellar assembly, share common molecular mechanisms.
1.4.4 Motility and Chemotaxis in *Agrobacterium tumefaciens*

*Agrobacterium tumefaciens* is a motile bacterium. It can recognise some compounds released from wounded cells, and move to the source of these chemicals (13, 191, 192). Motility of the bacterium is carried out by bacterial flagella. Electron microscopy has revealed that *A. tumefaciens* has an unusual flagellation pattern. The bacterium typically has a polar tuft of two flagella, plus two to four lateral flagella with curved filaments, which are composed of up to four different flagellins. The flagella, which are generally longer than the cell itself, only rotate in one direction, clockwise (CW), producing a run. The length of run is relatively long (frequently in excess of 500μm in C58C1 strain), and is rarely interrupted by changes of direction such as those seen in *E. coli* (117, 118, 180). The mechanism used by *A. tumefaciens* to change direction is unknown. These behavioural features of *A. tumefaciens* are similar to *Sinorhizobium meliloti*, but are different from those of *E. coli*. Motility and chemosensory system of the α-subgroup of proteobacteria, such as *Agrobacterium*, *Azosperillum*, *Caulobacter* and *Rhizobium* are distinctly different from the enterobacterial paradigm (8). Some differences has been described in section 1.3.7. One distinguishing features between two groups is existence of two different CheY acting with new MotC and MotD motor proteins to control flagellar rotatory speed (79, 189). *Agrobacterium* flagella structure are also different from *E. coli*. Filaments of *E. coli* possesses only one protein (FliC) whereas *A. tumefaciens* filaments is composed of several types of flagellin subunits. Four flagellins contribute to *Agrobacterium* filament structure: FlaA is the major protein, FlaB and FlaC are present in lesser amounts, and FlaD is minor component.

The strain C58C<sup>1</sup> used in this work, has been shown to be an active swimmer, with long straight or curved runs. Other *A. tumefaciens* genotypes such as LBA4301 and A136 are relatively poorly motile in comparison to C58C<sup>1</sup> (14, 118).

It has been suggested that most, if not all, motile bacteria use the MCP chemotactic signaling system found in enteric bacteria. A number of experiments indicated that *A. tumefaciens* also possesses MCP-like proteins and employ a similar signal transduction pathway to *E. coli*. When *A. tumefaciens* is exposed to a gradient of attractant, it will migrate in a favourable direction (14). The involvement of methylation for chemotaxis of *A. tumefaciens* to sucrose was demonstrated by Loake
Experiment carried out by Brown et al. showed that Agrobacterium requires methionine for chemotaxis to occur (34). An open reading frame has also been identified by Farrand and his co-workers which shows significant identity to a C. crescentus MCP (70). Another experiment showed that some Agrobacterium DNA fragments hybridised with a synthetic oligonucleotide complimentary to the conserved signaling domain of E. coli MCPs (149). VirA, a member of the two-component virulence system, structurally resembles MCPs and has sequence conservation with CheA. VirG, another member of the same two-component system also possesses similarity to CheB and CheY. A report showing that a protein with significant homology to a known MCP has also been found in the Ti plasmid pTiC58. Taken together with other results described earlier, this evidence indicates the existence of methylation-dependent chemotaxis in A. tumefaciens.

Agrobacterium has been shown to possess a highly sensitive chemotaxis system, with threshold responses occurring below $10^{-8}$ M (181). However, in most cases the A. tumefaciens response is more sensitive than E. coli (118). This chemotactic interaction with possible host plants of Agrobacterium appears to involve two distinct signaling pathways. A general chromosomally encoded chemotaxis pathway, which responds to nutrients such as amino acids and sugars, is responsible for maintaining the bacteria in sufficiently large number in the rhizosphere for infection to take place (118). Another system is a plasmid encoded pathway, which is responsible for movement of the bacterium to susceptible plant cells (12).

1.5. Analysis of chemotaxis and Motility genes in Agrobacterium tumefaciens

In order to study Agrobacterium motility and chemotaxis genes in more detail, a number of chemotactic mutants were generated using a Tn5 suicide vector, followed by selection for non-chemotactic mutants (117, 180). Further experiments associated with complementation and hybridization showed that similar to E. coli and other motile bacteria, the motility and chemotaxis genes in A. tumefaciens were thought to clustered together on the chromosome. A number of genes involved in flagellar structure, assembly, function and switching have been found on cosmid pDUB1900 from an A. tumefaciens genomic library. Analysis of 22 kb of sequence from this cosmid have identified an operon encoding homologues to the flagellar switch
proteins FliM, FliN and FliG (56), a group of flagellar genes consisting of \( \text{flgB}, \text{flgC}, \text{flgG}, \text{fliE}, \text{flgI} \) and \( \text{flgH} \) whose protein products are involved in basal body formation, a possible flagellum export gene \( \text{fliP} \) and two genes of unknown function (57), \( \text{flgA} \) and \( \text{fliL} \). FlgBCG and E are rod proteins, whereas FlgI and H are L and P ring proteins. Recently four flagellin genes, \( \text{flaA}, \text{flaB}, \text{flaC}, \text{flaD} \) which are concerned with flagellar filament structure have been found in this cluster (55-57). FlaA is the major protein, FlaB and FlaC are present in lesser amount, and FlaD is a minor component. An additional putative operon including the homologues to the \( \text{fliI}, \text{flhB} \) and \( \text{flgF} \) were also found in this region.

A cluster of chemotaxis genes has been identified on cosmid pDUB1911 from a representative genomic library of C58C (228). The cluster consists of \( \text{orf1} \), whose predicted protein shows strong homology to an MCP, followed by \( \text{orf2}, \text{cheY1}, \text{cheA}, \text{cheR}, \text{cheB}, \text{cheY2}, \text{orf9} \) and \( \text{orf10} \). A homologue to the flagellar gene \( \text{fliF} \) was found directly downstream of this cluster.

### 1.6 Aims of this project

The main goal of this work was to further characterise the motility and chemotaxis genes in \( \text{A. tumefaciens} \) using cosmid pDUB1905. This cosmid had previously been shown to contain \( \text{che-2} \) sequence and sequences spanning the site of two other mutations, \( \text{fla-8} \) and \( \text{mot-6} \) (34, 180). These mutants had been created by insertion of the transposon Tn5. Electron micrographs and swarming experiments of these three mutants showed that the mutant \( \text{che-2} \) is a non-chemotactic flagellated motile mutation, which tumbles excessively. The non-motile mutants \( \text{fla-8} \) and \( \text{mot-6} \) were originally classified separately as by electron microscopy, \( \text{fla-8} \) did not possess flagella, whereas \( \text{mot-6} \) possessed normal flagella. The two latter mutants were shown to share a common site in the cosmid pDUB1905, their apparent phenotypes probably resulting from an unexpected error. Finding the real phenotype of \( \text{fla-8} \) and \( \text{mot-6} \) mutants was another goal of this project.

The region of pDUB1905, spanning the mutant sequences was initially to be sequenced and analysed. Depending upon the result, any flagellar or chemotaxis gene homologues identified, were to be further characterised. Functional studies of each gene were to be achieved using site-direct mutagenesis by insertion of a neomycin
cassette. This was to be followed by phenotypical analysis of the resulting mutant strains with light and electron microscopy, and in the swarm plate assays.

Preliminary evidence showed that the chemotaxis and motility genes are clustered together, in separate chromosomal regions (34, 180). The result of an early experiment suggested that the two cosmids, pDUB1900 and pDUB1905, which contain several chemotaxis and motility genes might be close to each other (55). Finding how close these two fragments was another goal of this project.
2.1 Materials.

All inorganic chemicals were of AnalaR quality and purchased from BDH Chemicals Ltd., Poole, Dorset, U.K. unless otherwise specified.

All organic chemicals and enzymes were from Sigma Chemicals Plc., Poole, Dorset, U.K. unless otherwise specified.

Lab M Nutrient Broth (no.2), Lab M Nutrient Agar, nylon hybridisation transfer membranes and radiochemicals were from Amersham Ltd., Bury, U.K.

Agar bacteriological (no.1) and yeast extract were from Oxoid Ltd., Basingstoke, Hants., U.K.

Restriction endonucleases, T4 DNA ligase, Klenow enzyme, Taq polymerase, corresponding buffers, DNA markers, X-gal, IPTG and wild type λ DNA were from NBL, Cramlington, Northumberland, U.K., Boehringer Mannheim (UK) Ltd., Lewes, U.K., or Helena Biosciences, Tyne & Wear, U.K.

Agarose was from BRL, Gaithersburg, U.S.A.

Ficoll 400 and Sephadex G-50 were from Pharmacia Fine Chemicals, Uppsala, Sweden.

Fuji RX-100 X-ray film was from Fuji Photo Film Co., Ltd., Japan.

Polaroid film was from Polaroid (UK) Ltd., St. Albans, Hertfordshire, U.K.

Filter paper (3MM) and laboratory sealing film were from Whatman International Ltd., Maidstone, U.K.

Minisart filters were from Sartorius GmbH, Postfach 3243, D-3400 Göttingen, Germany.
Nitrocellulose discs (25 mm, 0.22 μm pore size) for tri-parental matings were from Schleicher and Schuell, Postfach 4, D-3354, Dassel, Germany.

Oligonucleotides for use in sequencing and PCR analysis were from MWG-Biotech, Germany and PE-Applied Biosystems UK, Cheshire, U.K.

### 2.2 Bacterial Strains and Plasmids.

#### 2.2.1 *E. coli* Strains.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Characteristics</th>
<th>Resistance</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>DH5α</td>
<td>F- <em>supE</em>44 ΔlacU169 (f80 lacZΔM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1.</td>
<td>Lab. Stock (227)</td>
<td></td>
</tr>
</tbody>
</table>

#### 2.2.2 *A. tumefaciens* Strains.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Characteristics</th>
<th>Resistance</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>C58C'</td>
<td>Ti-cured strain of C58.</td>
<td>RifR</td>
<td>Lab. Stock (217)</td>
</tr>
<tr>
<td>C58C'/fliR</td>
<td>Behavioural mutant - created by insertion of a neomycin resistance cassette into <em>fliR</em>.</td>
<td>RifR, NeoR</td>
<td>This study</td>
</tr>
</tbody>
</table>

#### 2.2.3 Plasmids.

#### 2.2.3.1 Plasmid Vectors.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Characteristics</th>
<th>Resistance</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasmid</td>
<td>Characteristics</td>
<td>Resistance</td>
<td>Source</td>
</tr>
<tr>
<td>--------------</td>
<td>------------------------------------------------------</td>
<td>------------</td>
<td>-----------</td>
</tr>
<tr>
<td>pRK2013</td>
<td>Helper plasmid for conjugation in <em>A. tumefaciens</em></td>
<td>KanR</td>
<td>Lab. stock</td>
</tr>
<tr>
<td>pDUB2033</td>
<td>Neomycin resistance cassette.</td>
<td>AmpR, NeoR</td>
<td>Lab. stock</td>
</tr>
</tbody>
</table>

### 2.2.3.3 Recombinant Plasmid Containing Cloned Behavioural Genes.

<table>
<thead>
<tr>
<th>Cosmid</th>
<th>Characteristics</th>
<th>Resistance</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>pDUB1905</td>
<td>A C58C1 cosmid library clone containing a <em>BamHI</em> chromosomal fragment in the vector pLAFR-3 (carries behavioural genes).</td>
<td>TcR</td>
<td>Lab. Stock (180)</td>
</tr>
</tbody>
</table>
### 2.2.3.4 Recombinant SK+ Plasmids Containing Subcloned *Agrobacterium* Behavioural Genes.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Characteristics</th>
<th>Resistance</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pAB1</strong></td>
<td>pBR322 containing 12kb <em>BamHI</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>Lab. Stock (34)</td>
</tr>
<tr>
<td><strong>ph1</strong></td>
<td>pBluescriptII SK+ containing 5.0kb <em>EcoRI</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph2</strong></td>
<td>pBluescriptII SK+ containing 1.7kb <em>EcoRI-BamHI</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph3</strong></td>
<td>pBluescriptII SK+ containing ~3kb <em>EcoRI</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph4</strong></td>
<td>pBluescriptII SK+ containing 0.3kb PCR-synthesised fragment from chromosomal DNA, adjacent to pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph1.1</strong></td>
<td>pBluescriptII SK+ containing 3.7kb <em>EcoRI-HapI</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph1.11</strong></td>
<td>pBluescriptII SK+ containing 1.7kb <em>EcoRI-EcoRV</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph1.12</strong></td>
<td>pBluescriptII SK+ containing 1.5kb <em>EcoRV</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
<tr>
<td><strong>ph1.13</strong></td>
<td>pBluescriptII SK+ containing 0.8kb <em>EcoRV</em> fragment from pDUB1905.</td>
<td>AmpR</td>
<td>This study</td>
</tr>
</tbody>
</table>

### 2.3 Bacterial Growth Media, Conditions and Procedures.

The following media were used in this work:

**Lab M nutrient broth no.2 (LM-broth):**

25g made up to 1 litre with distilled water gives final concentrations of 10g.l⁻¹ beef extract, 10g.l⁻¹ balanced peptone no.1, 5g.l⁻¹ NaCl, pH 7.5 ± 0.2.
For swarm plates bacteriological agar was added at a concentration of 0.15 - 0.2%.

Lab M nutrient agar (LM-agar): 
28g made up to 1 litre with distilled water, gives final concentrations of 5g.l⁻¹ peptone, 3g.l⁻¹ beef extract, 8g.l⁻¹ NaCl, 12g.l⁻¹ agar no.2, pH 7.3 ± 0.2.

Antibiotics were added to media after autoclaving, to the following concentrations:
For *A. tumefaciens*; chloramphenicol 25μg.ml⁻¹, kanamycin 50μgml⁻¹, neomycin 100μg.ml⁻¹, rifampicin 100μg.ml⁻¹, streptomycin 300μg.ml⁻¹ and tetracycline 15μg.ml⁻¹.

For *E. coli*; ampicillin 50μg.ml⁻¹, chloramphenicol 34μg.ml⁻¹, gentamycin 10μg.ml⁻¹, kanamycin 50μg.ml⁻¹, neomycin 100μg.ml⁻¹, streptomycin 20μg.ml⁻¹, and tetracycline 15μg.ml⁻¹.

When selecting for the inactivation of the β-galactosidase gene, by insertion of DNA fragments into the multiple cloning sites of pBluescriptII and other plasmids, 40μl of 20mg.ml⁻¹ X-gal (in DMF) were spread over the surface of agar plates.

Liquid bacterial cultures were incubated on an orbital shaker at 200rpm at temperatures of 37°C for *E. coli* and 28°C for *Agrobacterium* strain. Short-term (1-2 months) stocks of cultures were kept at 4°C on solid agar plates. Long-term stocks were kept in 40% glycerol at -80°C.

Liquid cultures were inoculated with a flamed loop or a sterile cocktail stick. Solutions and bacterial cultures were spread onto agar plates using a glass spreader which had been sterilised in 70% ethanol. Bacterial colonies were inoculated into the centre of swarm plates using a sterile needle.
Aseptic technique was used throughout with bacterial cultures. All glassware, plasticware and other equipment were sterilised by autoclaving at 121°C, 15 psi. for 15 minutes. Generally solutions were prepared according to Sambrook et al. (169) and autoclaved, as above, if possible. Otherwise solutions were filter-sterilised through a 0.22 μm nitrocellulose filter into a sterile container.

2.4 Isolation of DNA.

2.4.1 Alkaline Lysis Plasmid Minipreps.

This method, used to prepare small amounts of relatively pure plasmid DNA, was according to Sambrook et al. (169)

A single colony of bacteria was grown overnight in 5ml of LM-broth containing the appropriate antibiotic selection. 1.5ml of this culture was pipetted into a sterile eppendorf tube, and the cells harvested by centrifugation for 1 minute in a microfuge. The supernatant was discarded, and a further 1.5ml of the culture pelleted as before. The supernatant was again discarded. The pellet was resuspended in 200μl of ice-cold solution 1 (1% glucose, 10mM EDTA pH8.0, 25mM Tris.HCl pH8.0), 200μl of solution 2 (0.2M NaOH, 1% SDS) was then added and the contents of the tube mixed by gentle inversion. 200μl of ice-cold solution 3 was added to the mixture and the tube inverted ten times to mix the contents. (Solution 3 was prepared by adding 1.15ml of glacial acetic acid to 2.85ml of distilled water and then adding 6ml of 5M potassium acetate). The solution has an overall pH of 4.8 and is 3M wrt potassium and 5M wrt acetate.) The tube was then microfuged for 5 minutes to remove bacterial debris. The supernatant was transferred to a fresh tube, spun down for a further 2 minutes and then pipetted into another fresh tube. The DNA was precipitated by the addition of 1ml of 100% ethanol. The tube was left for 5-15 minutes at -20°C then the DNA collected by centrifugation for 5 minutes. The DNA pellet was washed in 1ml 70% ethanol and air dried for 20-30 minutes. The final pellet was resuspended in 50μl of TE buffer and RNAase A added to a concentration of 20μg.ml⁻¹.
When plasmid DNA was to be prepared for sequencing, a QIAGEN QIAprep spin plasmid miniprep kit was used according to the manufacturers instructions.

2.4.2 Larger Scale Plasmid Preparation.

Single colonies were inoculated into two 50ml of LM-broth with antibiotic selection and grown overnight with shaking. The cultures were transferred to two 50ml centrifuge bottles and the bacterial cells harvested by centrifugation at 7000g/4°C for 7 minutes in a Beckman J2-HS centrifuge, using a JA-20 rotor. The supernatants were removed and the pellets resuspended in 3.335ml of ice-cold solution 1. 3.335ml of solution 2 was added to each tube and the tubes gently mixed. 3.335ml of ice-cold solution 3 was then added to each tube, and the tubes mixed well. The tubes were centrifuged for 5 minutes, at 4°C and 15000g in a Beckman J2-HS centrifuge, to remove cell debris. The supernatants were transferred to sterile 50ml Falcon tubes, 20ml of phenol:chloroform:isoamyl alcohol (25:24:1) were added and the tubes mixed well. The tubes were spun at 3500g for 3 minutes and the aqueous layer transferred to sterile centrifuge tubes. 20ml of ice-cold 100% ethanol was added to each tube and the tubes left on ice for 25 minutes. The DNA was pelleted by centrifugation for 15 minutes, at 4°C and 15000g in a Beckman J2-HS centrifuge. The pellet was then washed in 70% ethanol, dried and resuspended in 500μl of TE.

Solutions 1, 2 and 3 were made as in section 2.4.1

2.4.3 Small Scale Preparation of Bacterial Chromosomal DNA.

A 5ml LM-broth culture of bacteria was grown to stationary phase and 1.5ml of the culture transferred to an eppendorf tube. The cells were spun down for 1 minute in a microfuge, the supernatant removed and the pellet left at -20°C for 30 minutes. The pellet was then resuspended in 200μl of TE, and the cell suspension incubated with 8μl of lysozyme (10mg.ml⁻¹ stock) for 30 minutes at 37°C. The cells were then lysed by the addition of 40μl of 4M sodium perchlorate, 24μl of 10% SDS and 8μl of Proteinase K (20mg.ml⁻¹ stock), and incubated at 45°C for 2 hours. The DNA was precipitated by the addition of 2 volumes of ethanol and incubation at -20°C for 30 minutes. The DNA was pelleted by centrifugation in a microfuge for 5 minutes. The pellet was washed with 70% ethanol, dried and resuspended in 500μl of TE buffer.
The DNA solution was extracted three times with an equal volume of phenol:chloroform:isoamyl alcohol (25:24:1), the aqueous layer being transferred to a new eppendorf tube each time. The DNA was then extracted once with chloroform:isoamyl alcohol (24:1). The DNA in the final aqueous layer was precipitated by the addition of 2 volumes of ethanol and 0.1 volumes of 3M sodium acetate (pH 4.8), followed by incubation overnight at -20°C. The DNA was pelleted by centrifugation in a microfuge for 5 minutes. The pellet was then washed with 1ml of 70% ethanol, dried and resuspended in 50μl of TE buffer with RNAase A (10mg.ml⁻¹ stock) added to a final concentration of 20μg.ml⁻¹.

2.4.4 Small Scale Preparation of Bacterial Chromosomal DNA - 2.

This method was used as a quick preparation of Agrobacterium DNA when screening for possible mutant strains.

A 5ml LM-broth culture of bacteria was grown to stationary phase and 1.5ml of this culture transferred to an eppendorf tube. The cells were spun down for 1 minute in a microfuge, the supernatant removed and the pellet resuspended in 200μl of lysis buffer (lysis buffer consists of 40mM Tris-acetate pH7.8, 20mM Sodium acetate, 1mM EDTA and 1% SDS). 82.5μl of 4M NaCl was then added and the tube contents mixed by inversion. The tubes were centrifuged for 10 minutes in a microfuge. The supernatant was then pipetted into a clean eppendorf tube and an equal volume (approximately 250μl) of chloroform:isoamyl alcohol (24:1) added. The tube was inverted 50 times, then centrifuged for 3 minutes in a microfuge. The upper aqueous layer was removed to a clean eppendorf and the DNA precipitated by the addition of two volumes of 100% EtOH and incubation at -20°C for 10 minutes. The DNA was pelleted by centrifugation in a microfuge for 5 minutes. The pellet was then washed with 1ml 70% ethanol, dried and resuspended in 30μl of TE buffer with RNAase A (10mg.ml⁻¹ stock) added to a final concentration of 20μg.ml⁻¹.

2.5 DNA Manipulations.

2.5.1 Phenol:Chloroform Extraction of DNA.
A 25:24:1 solution of phenol:chloroform:isoamyl alcohol was equilibrated 3
times with TE buffer and stored, under TE, in a light-proof bottle at 4°C. To remove
proteins from DNA solutions an equal volume of phenol:chloroform:isoamyl alcohol
was added, the solutions mixed by vortexing for 30 seconds and the phases separated
by centrifugation for 2 minutes in a microfuge. The aqueous phase was transferred to
a fresh tube. This was repeated until no further protein was visible (as a white
precipitate) at the boundary of the two phases.

2.5.2 Restriction Endonuclease Digestions.

Digestions were carried out according to the enzyme manufacturer's
instructions. Generally plasmid DNA was digested in a total volume of 10-40μl, with
5 units of restriction endonuclease, 0.1 volumes of the supplied 10x concentrated
enzyme buffer, and sterile distilled water to make up the volume. The reaction was
incubated at the recommended temperature (usually 37°C) for 1-3 hours. If more than
one restriction enzyme was to be used in the same reaction and the buffers supplied
differed, the reaction was buffered using one-phor-all buffer PLUS (Pharmacia).
Chromosomal DNA was digested in a larger volume, 100-200μl, with 10 units of
restriction enzyme added for every microgram of DNA, as well as the appropriate
amounts of buffer and sterile distilled water. The reaction mixture was covered with a
layer of mineral oil to prevent evaporation and maintain the buffering conditions,
before being incubated overnight at the required temperature.

If the digestions were to be analysed by gel electrophoresis, 0.2 volumes of 6x
gel-loading buffer were added (6x gel-loading buffer contains 0.25% bromophenol
blue, 0.25% xylene cyanol FF and 40% sucrose in distilled water - this was filter
sterilised and stored at 4°C).

If the digested DNA was to be used in further subcloning steps, the required
fragment was cut out of the gel and purified using silica fines (see section 2.5.4).

2.5.3 Agarose Gel Electrophoresis.

Gel electrophoresis of DNA samples was carried out with large 180x200mm
maxigels (volume 400ml), 100x80mm minigels (volume 50ml) or 77x55mm
minigels (volume 35ml). Maxigels were run in a Pharmacia gel apparatus GNA-200
electrophoresis tank. Minigels were run in Pharmacia gel apparatus GNA-100 electrophoresis tanks. The concentration of agarose within a gel could be varied depending on the size of DNA to be separated. Usually a 0.7% agarose gel was used, which efficiently separated linear DNA between 10-0.8kb. The required amounts of agarose and 1x TAE buffer (50x stock - 242g Tris, 100ml EDTA pH8.0, 57.1ml glacial acetic acid per litre) were mixed and the agarose dissolved by microwaving the mixture. The solution was cooled to about 60°C, 10mg.ml⁻¹ ethidium bromide was added to a final concentration of 0.2μg.ml⁻¹, and the agarose was poured into a gel mould with a well comb in place. Once the agarose had set the gel was put in a tank and covered with 1x TAE buffer containing 0.2μg.ml⁻¹ ethidium bromide. The DNA samples (and size markers) were loaded and electrophoresis carried out at 0.1A (5-10 V.cm⁻¹) for the required amount of time, usually 1 - 1.5 hours. The size markers used were either λ-DNA digested with PstI and/or λ-DNA digested with HindIII.

When DNA samples less than 1kb in size were to be analysed by agarose gel electrophoresis, Metaphor agarose and TBE buffer (5x stock - 54g Tris, 27.5g Boric acid, 20ml 500mM Na₂EDTA (pH 8.0) made up to 1 litre with dH₂O) were used. Usually 2% Metaphor agarose gels were made with 1x TBE, using the same methodology as above.

**PstI digested λ-DNA produces DNA fragments of the following sizes (in kb):**

14.05, 11.49, 5.07, 4.75, 4.51, 2.84, [2.56, 2.46, 2.44], 2.14, 1.99, 1.70, 1.16, 1.09, 0.81, 0.52, 0.47, 0.45, 0.34...

The fragments enclosed in brackets run together on an agarose gel. Smaller fragments are also produced, but were rarely seen in this work.

**HindIII digested λ-DNA produces DNA fragments of the following sizes (in kb):**

23.13, 9.42, 6.56, 4.36, 2.32, 2.03, 0.56, 0.13.

DNA within the gel was visualised on a transilluminator (UVP Inc.), and photographed with a Polaroid RP4 Land camera (using a red filter) onto Polaroid 667 film.
2.5.4 Isolation of DNA Fragments from Agarose Gels Using Silica Fines.

The required band was cut out of the gel using a sterile scalpel blade, and placed in a labelled 1.5ml eppendorf tube. 800µl of sodium iodide solution were added to the tube and the tube placed at 70°C for 5 minutes to melt the agarose. Once the agarose had completely melted the tube was mixed by inversion and allowed to cool to room temperature for 5 minutes. 5µl of silica fines were then added, mixed and the solution left for 10 minutes at room temperature with continuous shaking. The fines were spun down for 30 seconds in a microfuge, the supernatant removed and the fines washed with 70% ethanol. The silica fines pellet was dried with a piece of tissue, then resuspended in 30µl of TE buffer and incubated at 37°C for 10 minutes with occasional shaking. The fines were spun down in a microfuge for 30 seconds and the supernatant, containing the DNA, collected. The fines were then resuspended in 20µl of TE and incubated as before. Further centrifugation and collection of the supernatant gave a total volume of 50µl of DNA.

Preparation of the silica fines:

250ml of silica 325 mesh powder was resuspended in distilled water to give a total volume of 500ml. The suspension was stirred for 1 hour and left to settle for a further hour. The suspension was then centrifuged at 5000g in a Beckman J2-HS centrifuge using a JA-14 rotor. The pellet was resuspended in 150ml of distilled water plus 150ml nitric acid. The suspension was then heated to 98°C and allowed to cool to room temperature. The silica fines were then repeatedly washed with sterile distilled water until the pH was greater than 5.5. Silica fines were stored at 4°C as a 50% slurry in sterile distilled water.

A QIAGEN QIAquick Gel Extraction Kit was also sometimes used, according to the manufacturers instructions, for isolating DNA from agarose gels.

2.5.5 Filling in 3'-Recessed Termini.

The DNA fragment (maximum of 500ng) was resuspended in 50µl of TE following isolation from an agarose gel. 7µl of a solution containing all 4 dNTPs
(each at 1mM) was added to the DNA. 7µl of Klenow buffer (10x) was added (10x buffer is 0.5M Tris.HCl pH 7.6, 0.1M MgCl₂) and the reaction buffer made up to 70µl with sterile distilled water plus 1µl (1 unit) of Klenow fragment. The reaction mixture was left at room temperature for 30 minutes and then the Klenow fragment either inactivated by incubating at 70°C for 5 minutes, or removed using a QIAGEN PCR Purification Kit, according to the manufacturers instructions.

2.5.6 Ligation of DNA.

T4 DNA ligase was used to ligate DNA fragments with compatible cohesive or blunt termini. The fragments of insert and vector DNA were usually mixed at a ratio of 3:1 (insert:vector) with a maximum of 300ng DNA. 0.1 volume of 10x ligase buffer (0.66M Tris.HCl pH7.5, 50mM MgCl₂, 50mM DTT, 10mM ATP) was added and 1 unit of DNA ligase. For cohesive termini the tubes were incubated overnight at 4°C. For blunt-ended termini the reactions were incubated at 16°C overnight. The ligation mix was then used immediately to transform competent E.coli cells.

2.5.7 Polymerase Chain Reaction (PCR).

All PCR reactions were performed in a Perkin-Elmer Thermal Cycler. Reactions were usually carried out in a total volume of 25µl, in thin-walled, 0.5ml, PCR tubes. Per reaction tube the following components were added:

- 10x reaction buffer 2.5µl
- dNTPs 2.0µl (each at 1mM)
- primer 1 1.0µl (at a concentration of 10pmol/µl)
- primer 2 1.0µl (as above)
- Taq polymerase 0.25µl (1 unit)
- template DNA 1.0µl (see below)
- 25mM Mg²⁺ 1-4µl (concentration used varied)
- dH₂O appropriate to make final volume 25µl

The amount of template DNA used varied, however it was usually approximately 100ng for plasmid DNA, and 500-1000ng for genomic DNA.
To avoid pipetting very small volumes (<1μl) the 10x reaction buffer, dNTPs, primers and Taq polymerase were made into a "Master mix". For n tubes, the master mix consisted of appropriate volumes of the different components to make up n+1 reaction tubes. The mixture was then dispensed into each of the reaction tubes.

**Primer design:** Primers used were generally designed so as to contain an equal number of G+C's and A+T's, usually 10 of each pair. Care was taken to ensure the primers would not self anneal or hybridise to each other, as this would reduce the efficiency of the PCR reactions.

**PCR Protocol:** The temperatures and times used in PCR reactions followed the same basic pattern:

- **94°C** 5 minutes
- **94°C** 1 minute → denature
- **55°C** 1 minute \(\times 30\) → anneal
- **72°C** 1 minute → extend
- **72°C** 5 minutes

*1 The annealing temperature used varied as it is set by the composition of the primers; The approximate Tm for each primer can be worked out using the following equation:

\[
T_m = \left\{\left[\text{(no. of G + C) x 4}\right] + \left[\text{(no. of A +T) x 2}\right]\right\} - 5 \, ^\circ C
\]

If the annealing temperatures of the two primers differed, the lower temperature was used in the reactions.
• The extension time is set by the length of the largest fragment expected from the reactions, e.g. if expect a 3kb fragment the extension time would be 3 minutes.

2.5.7.1 Hot Start PCR.

Occasionally "hot start" PCR was used, this reduces the amount of non-specific binding of the primers by compartmentalising the contents of the reaction tube until the denaturation temperature is reached. Half the total volume of the reaction (consisting of the 10x buffer, dNTPs, Mg$^{2+}$, primer 1 and dH$_2$O) was put in a tube, a wax bead was then added, using sterile forceps, and the tube heated to 70°C for 5 minutes to melt the wax. After heating, the tube was held on ice for 2 minutes to solidify the wax and seal off the bottom half of the tube. The second half of the reaction volume (consisting of the Taq polymerase, template DNA, primer 2 and dH$_2$O) was then added, on top of the wax seal, and the tubes placed in the thermal cycler.

2.6 Transformation of E. coli.

2.6.1 Preparation of Competent Cells.

5ml of LM broth was inoculated with DH5α and grown overnight at 37°C, it was then subcultured 1:100 into fresh LM broth and the cells grown to an OD$_{550}$ of 0.3-0.35. The culture was chilled for 5 minutes on ice before being spun down in pre-chilled centrifuge tubes at 4000g, 4°C, for 7 minutes. The supernatants were poured off and the cell pellets resuspended in 2/5 of the original culture volume with solution A (solution A = 30mM potassium acetate, 100mM rubidium chloride, 10mM calcium chloride, 50mM manganese chloride and 15% glycerol. The solution was adjusted to pH 5.8 with 0.2M acetic acid and filter sterilised.) The tubes were held on ice for 5 minutes and spun down as before. The supernatants were poured off and the pellets resuspended in 1/25 of the original culture volume with solution B (solution B = 10mM MOPS, 75mM CaCl$_2$, 10mM RbCl$_2$ and 15% glycerol. The solution was adjusted to pH 6.5 with KOH and filter sterilised.) The tubes were left on ice for 15
minutes, then 150μl aliquots were added to pre-chilled eppendorfs, frozen in liquid nitrogen and stored at -80°C.

2.6.2 Transformation Procedure.

The cells were thawed by hand and placed on ice for 10 minutes. The DNA to be transformed was added, and the tube held on ice for 30 minutes. The cells were heat shocked at 42°C for 90 seconds, held on ice for 3 minutes and then 800μl of pre-warmed (to 37°C) LM broth were added. The tube was incubated for 1 hour at 37°C, with continuous shaking. Finally appropriate aliquots, usually 1/10th and 9/10ths of the tube, were spread onto selective agar plates and incubated overnight at 37°C.

2.7 DNA Hybridisation Procedures.

2.7.1 Radio-Labelling of DNA Fragments.

DNA fragments were labelled with [α-32P] dCTP by the random primer labelling method using an Amersham Multiprime kit. 30-50ng of the DNA to be labelled, in a total volume of 28μl, were boiled for 5 minutes and then held on ice for 2 minutes. 10μl of labelling buffer, 5μl of random hexanucleotide primers, 5μl 32P-dCTP (equivalent to 50μCi) were added, followed by 2μl of Klenow enzyme. The labelling reaction was left to proceed either at room temperature overnight, or for 2-3 hours at 37°C, the probe was then purified through a 10cm Sephadex G-50 column using STE as the eluant. The labelled DNA was collected and stored at -20°C until required. Immediately before use the probe DNA was boiled for 5 minutes.

2.7.2 Southern Blotting.

DNA was transferred to Hybond-N (Amersham) nylon membranes, according to the manufacturer's instructions. Firstly the agarose gel containing the DNA samples was photographed with a ruler down its side. Gels known to have DNA fragments greater than 10kb in size were soaked in 0.25M HCl for 15 minutes, to partially depurinate the DNA, and rinsed twice with distilled water. For blotting the gel was soaked in denaturation buffer (1.5M NaCl, 0.5M NaOH) with occasional
shaking for 30 minutes. The gel was then rinsed twice with distilled water and soaked in neutralisation buffer (1.5M NaCl, 0.5M Tris.HCl pH7.2, 0.001M EDTA). After rinsing the gel twice with distilled water, the blot was set up.

For single sided (one-way) blots a reservoir of 20x SSC was set up (20x SSC - 3.0M NaCl, 0.3M Na.citrate pH7.0). A platform was placed over this reservoir and a long piece of Whatman 3MM paper (presoaked in 20x SSC) put on the platform with its ends dipping into the reservoir. The gel was placed, wells uppermost, on the 3MM paper, and a piece of Hybond-N nylon membrane cut to the same size as the gel placed on top of it. Any air bubbles were carefully removed before 3 sheets of Whatman 3MM paper, cut to the gel size and presoaked in 20x SSC, were placed on top. Finally 2 layers of disposable nappies (also cut to gel size) were placed on top, the stack covered with a glass plate and a 1kg weight placed on top.

Double sided (two-way) blots were created by sandwiching the agarose gel between two equivalent stacks of a glassplate, 2 layers of disposable nappies, 3 pieces of Whatman 3MM paper (presoaked in 20x SSC) and a piece of Hybond-N nylon membrane. A 1kg weight was placed on the top glass plate. Liquid retained in the gel transferred the DNA onto both membranes.

Both types of blot were left for at least 16 hours to allow DNA transfer. After which time the apparatus was dismantled, and the positions of the wells marked on the nylon. The nylon filter was washed carefully in 2xSSC, then allowed to air dry for up to 1 hour before being wrapped in clingfilm. It was then exposed to UV light for two minutes each side to fix the DNA to the membrane.

2.7.3 Hybridisation of Radio-Labelled Probes to Blots.

Hybridisation reactions were carried out using Techne Hybridisation tubes in a Techne Hybridiser HB-1 oven. The nylon filter was put inside the hybridisation tube and 200μl of pre-hybridisation solution (5x SSC, 5x Denhardt's solution [50x Denhardt's solution is 1% ficoll, 1% polyvinylpyrrolidone, 1% BSA fraction V], 0.5% SDS, 0.1% pyrophosphate and 100mg.ml⁻¹ of denatured salmon sperm DNA) added per cm² of filter. Any air bubbles were carefully removed and the tube incubated at 65°C, whilst being rotated in the oven. After two hours the labelled probe was denatured, by boiling for 5 minutes, and 100-200 μl added to the tube contents. The tube was replaced in the oven and incubated at 65°C for at least 12
hours. After incubation the pre-hybridisation solution (containing the probe) was poured off and the blots washed.

2.7.4 Washing of Probed Blots.

The nylon filter was washed within the hybridisation tube and was never allowed to dry out. The filter was washed twice in 2x SSC, 0.1% SDS for 10 minutes at room temperature, followed by one wash in 0.1x SSC, 0.1% SDS for 15 minutes at 65°C. After each washing solution was removed, the filter was checked with a Geiger counter and the washing continued until sufficient (apparent) non-specific radio-labelled probe was removed (until a reading of <20cpm was obtained). Finally the filters were wrapped in clingfilm.

2.7.5 Detection of Hybridising Probes.

The wrapped filter was taped onto a larger piece of Whatman 3MM paper in a lead cassette. Radioactive bands could be detected on the filter by exposing Fuji RX-100 X-ray film to the filter. The film sheets were pre-flashed once to sensitise them, and exposure carried out at -80°C, for varying amounts of time. Exposed films were developed with Ilford Phenisol developer for up to 4 minutes and fixed with Kodak Unifix fixer for 2 minutes. The positions of the wells on the filter were marked on the film and the size of any hybridising fragments calculated using the original gel photograph (with ruler included).

2.7.6 Stripping of Probed Blots.

Blots were stripped by pouring a boiling solution of 0.1% SDS over the membrane and allowing the solution to cool to room temperature. Stripping was usually allowed to proceed for a period of at least two hours, after which time the membrane was wrapped in clingfilm and autoradiography carried out for the normal exposure time to check that the probe had been completely removed. The membrane was then pre-hybridised and hybridised with a new probe.
2.8 DNA Sequencing.

DNA sequencing was carried out with an Applied Biosystems 373A DNA Stretch Sequencer and a 377 DNA sequencer, using double stranded DNA templates and dye terminator chemistries. Usually, the universal M13 forward and reverse primers were used, but occasionally custom-synthesised oligonucleotides were also used.

Nucleotide sequence searches were carried out against the GenBank and EMBL databases using the Fasta programme at SEQNET. Putative protein sequences were searched against the Owl database, using the programmes Sooty and Sweep, and BLASTP.

2.9 Conjugation of Plasmids into Agrobacterium.

Triparental matings based upon the method of Ditta et al. (62) were used to mobilise plasmids into Agrobacterium with pRK2013 as a helper plasmid. Cultures of the recipient Agrobacterium strain, and the E.coli plasmid donor and helper strains were grown to mid log phase. 100μl of each culture were pipetted onto a 0.22μm nitrocellulose filter disc on the surface of an LM-agar plate. This was incubated at 28°C overnight, and the disc then transferred to a universal bottle containing 10ml of 10mM MgSO4. This was vortexed vigorously to wash the bacteria off the disc. Dilutions were made from the resulting cell suspension and plated onto agar plates, with appropriate antibiotic selection.

2.10 Mutagenesis.

2.10.1 Gene Replacement Mutagenesis.

Gene replacement mutagenesis was carried out according to the protocol of J. Quandt and M. F. Hynes (162). The technique involves replacement of a functional gene with a copy of the same gene interrupted by the insertion of an antibiotic resistance cassette, thereby creating a non-functional copy of the gene in question.
Two vectors were used in this work, pJQ200mp18 and pJQ200uc1, which only differ in the restriction enzymes available in their respective multiple cloning sites. The pJQ200 vectors have gentamycin resistance and a β-galactosidase gene for selection procedures. In addition they have a functional mob site which allows conjugation into *A. tumefaciens* strains. They also possess the sacB gene from *B. subtilis*, which when activated by sucrose, produces levansucrase, an enzyme whose expression causes the production of toxic compounds that are lethal in Gram negative bacteria. Also crucial to their functioning in this work, is the fact that the pJQ200 vectors are unable to replicate outside of the enterobacteria, they therefore act as suicide vectors in *A. tumefaciens*.

Genes to be mutated were ligated into a pJQ200 vector and correct insertion confirmed by inactivation of the β-galactosidase gene. The neomycin resistance cassette from pDUB2033 was then ligated into the gene and positive constructs obtained by selection on plates containing both gentamycin and neomycin. The final plasmid construct was then conjugated into *A. tumefaciens* C58C1 by tri-parental mating (see section 2.9).

The pJQ200 constructs are only able to survive in *A. tumefaciens* by integration into the genome via homologous recombination. For correct construction of the mutant strain, a rare second recombination event must occur between the mutated gene and the wild type copy, thus excising the wild type copy of the gene and the plasmid vector from the cells.

After the first recombination and subsequent integration step the entire mutant construct was incorporated into the genome, giving the bacteria both gentamycin and neomycin resistance. Since both intermediate strains and the final mutants possess resistance to neomycin it was not possible to select for correct construction of the mutant by simply growing on plates with neomycin selection alone. The presence of the sacB gene on the pJQ200 vector however acts as an additional selectable marker, allowing direct selection of only those strains that have undergone the second recombination event, and contain only the mutated copy of the gene. Selection on sucrose induces the sacB gene, thereby killing those strains that still contain the vector sequence.

Tri-parental matings were therefore plated on LM-agar plates containing rifampicin, neomycin and 5% sucrose.
To confirm that gene replacement had occurred, genomic digests and subsequently Southern blots were made of the putative mutant strains and the wild type strain C58C1. The blots were probed with a radioactively labelled copy of both the wild type gene, and the neomycin cassette. Due to the insertion of the neomycin cassette, when digested with the same enzyme, different sized hybridising bands were expected in the mutant strains, as compared to the wild type strain, when probed with the gene in question. The neomycin probe should only hybridise with known sized fragments in the mutants. The differences in band patterns between the wild type strain and the mutant strain therefore confirms gene replacement has occurred.

2.11 Bacterial Growth Measurement.

The optical density of bacteria was measured at 600nm on a Beckman DU7500 spectrophotometer to produce growth curves of the changes in optical density over time.

2.12 Microscopy.

2.12.1 Light Microscopy.

A loopful of bacteria, taken from a fresh swarm plate, was resuspended in 40μl of chemotaxis media (1ml 0.1M EDTA and 10ml 1M KH$_2$PO$_4$ (pH7.0) made to1 litre with distilled water.) and observed under phase contrast optics (using a Nikon Optiphot microscope).

2.12.2 Electron Microscopy.

A loopful of bacteria, taken from a fresh swarm plate, was washed onto a Formvar-coated grid with 100μl of 10mM MgSO$_4$. The bacteria was allowed to settle on the grid for 5 minutes before the excess solution was blotted off. Grids were not allowed to dry out completely at this stage. A drop of saturated aqueous uranyl acetate (pH 4.8) was placed on the grid for 30 seconds, the excess stain was then partially blotted off and the grid allowed to air dry. The grids were viewed and photographed using a Philips EM400.
3. Gene Identification
3.1 Background of this project:

In order to study motility and chemotaxis genes in *Agrobacterium tumefaciens* in more detail, a number of mutants were generated using a Tn5 suicide vector, followed by selection for non-motile and non-chemotactic mutants. Tn5 is a 5.8 kb transposon carrying a kanamycin resistance gene and has specific properties that facilitate genetic analysis of resulting mutants (185). Tn5 has transposase activity allowing it to insert itself randomly, with a high frequency, into the chromosome or plasmid of a wide range of Gram-negative bacteria, producing a mutation at the insertion site. It exhibits a low probability of genome rearrangement during insertion and a high stability when inserted. The DNA fragment, containing Tn5 can be easily isolated using standard methods.

A plasmid with a high transfer frequency was constructed to deliver the insertion mutagen Tn5 (117). The *E. coli* strain carrying the Tn5 "suicide vector" was mated directly to the rifampicin resistant *A. tumefaciens* strain C58C1. Transconjugants were screened for motility defects in L-agar swarm plates containing Rif and Km antibiotics. Putative motility and chemotaxis mutants were isolated by selecting strains that produced smaller swarm morphologies compared to the wild type (figure 3.1.1) (180).

Twenty independent behavioural mutants, showing impaired swarming ability, were isolated (180). They were further characterised using flagella staining, electron microscopy and motility studies with a phase contrast microscope. Preliminary phenotype analysis showed that 14 of the non-swarming mutants were non-motile. One of the mutants however proved to be motile but tumbled continuously, and was thus identified as a possible non-chemotactic mutant and assigned as che-2. Although che- mutants are fully motile, they are thought to form smaller swarms than wild type due to a defect in the intracellular signaling pathway.

Electron microscopic observation divided the 14 non-motile mutants into two subclasses: those possessing normal flagella, which are presumably non-functional in some flagellar components (mot- mutants) and those lacking flagella, which were designated fla-. The other mutants, ssw and tpc have flagellar structures different from the wild-type, however they show poor motility and small swarms (180).
Figure 3.1.1 Tn5 mutagenesis. (see text for details). Adapted from (180).

Further experiments were carried out to clone the DNA fragments surrounding the Tn5 insertion sites. Chromosomal DNA from each mutant was isolated, digested with EcoRI, then probed with a radioactively labelled Tn5 fragment. The bands containing the Tn5 insertion site plus flanking sequences from each *A. tumefaciens* behavioural mutant were cloned into the plasmids pUC18/19. The new plasmids were designated pDUB1801 to 19. The mutant flanking sequences were isolated by cleavage
Behavioural mutant che-2

Cut with EcoRI, transform E. coli, ligate pUC18; select km

Cut with EcoRI/Hpal; purify flanking sequences

Use as probes

Screen A. tumefaciens cosmid wild-type pLAFR3 library

che-2 25 kb cosmid insert

Figure 3.1.2 Isolation of behavioural mutants. che-2 is shown as an example (see text for details). Adapted from (180).
with EcoRI and HpaI. The latter enzyme has two sites, 185 bp from each end of Tn5, while EcoRI has no site in Tn5.

To isolate DNA that codes for chemotaxis and motility genes in wild-type A. tumefaciens, a representative genomic library of C58C1 genomic DNA was constructed (117). A. tumefaciens C58C1 DNA was partially digested with BamHI and 20-25 kb fragments ligated into the cosmid vector pLAFR3. The ligation mixture was transformed into E. coli and 1200 separate recombinant clones were selected and maintained separately. The motility flanking sequences were then radiolabelled and used as probes to screen the genomic library cosmids (see figure 3.1.2). The identified cosmid clones were then used in complementation tests on the mutants, and any others found to hybridise to the same cosmid. This method showed that the mutated loci mapped to two separate cosmids carrying a total of 12 loci. Cosmid clone pDUB1900 was found to either complement or hybridise to seven of mutants: mot-1, mot-4, mot-9, mot-12, fla-3, fla-11 and fla-15. Cosmid clone pDUB1905 was found to complement three mutants, che-2, mot-6, and fla-8. The two latter mutants share the same position in the cosmid pDUB1905. E. L. Wright has recently constructed a cosmid, pDUB1911, using heterologous probing with a DNA fragment from S. meliloti. pDUB1911 contains a che operon which consists of a number of chemotaxis and motility genes (228).

3.2 Sequencing plasmid pH1.

It has previously been shown that the behavioural genes in the cosmid pDUB1905 are located on a 12 kb BamHI fragment. This fragment was isolated to produce the plasmid pAB1. The initial work was to sequence a 5 kb EcoRI fragment from pAB1. This region was known to contain DNA complementing three phenotypically distinct motility and chemotactic mutants, mot-6, fla-8 and che-2 (see figure 3.2.1a, and 3.2.2)(180).

Generally, DNA fragments to be sequenced were subcloned into pBluescript SK+ and the universal M13 forward and reverse primers were used as described in section 2.8. SK+ was chosen because of its wide range of unique restriction enzymes that make analysis of double digestion easier, and the lacZ region, which is used for blue/white selection of recombinant plasmids.
**Figure 3.2.1 Cosmid pDUB1905 and major plasmids derived from the cosmid.**

a) Cosmid pDUB1905. The arrows underneath the cosmid showing plasmid pABI.
b) Plasmid pABI, arrows above the plasmid indicating the position of the genes and ORFs found in this work. X, Y, W and Z representing *orfX*, *Y*, *W*, and *Z*. Arrows underneath pABI are showing three other plasmids generated from pABI.
c) Plasmid pH1 and three major plasmids, which generated in this work. Positions corresponding to the mutations and restriction enzyme also have been shown.

B = *BamHI*   E or RI = *EcoRI*   R5 = *EcoRV*   H = *HindIII*
Figure 3.2.2 Double digestion of pAB1 (a) with BamHI-EcoRI, and (b) pH1 with EcoRI-EcoRV.

The fragments isolated and used for subcloning are shown by arrows.

The 5 kb EcoRI fragment was subcloned into pBluescript SK⁺, producing the plasmid pH1. Sequencing of this plasmid was carried out by further subcloning of fragments isolated from the plasmid pH1 into pBluescript SK⁺. When there were no convenient sites for restriction digesting, primers were designed for sequencing. The plasmid pH1 has been fully sequenced in both directions (see figure 3.2.1c).

Raw sequence data was initially manipulated and aligned using the Macintosh computer programs DNA Strider™ 1.2 and Sequencher. Restriction enzyme sites were found by computer analysis, and confirmed by experimental digests of the DNA and comparing the size of the fragments produced with those predicted. Possible open reading frames (orf's) of the pH1 sequence were identified using DNA Strider and the Seqnet program TESTCODE (this program calculates the probability of the sequence to contain coding regions.).
The whole sequence was compared against nucleic acid databases to identify homology for chemotaxis or motility genes. The University of Wisconsin genetics computer group (UWGCG) sequence analysis software package mounted on the SEQNET VAX 3600 at SERC, Daresbury, UK, was used for DNA homology searches. The sequence was compared against the GenBank and EMBL databases using the FASTA program. Alternatively the orfs were first translated into their protein sequence and compared against the Owl database using the SEQNET program SWEEP and BLASTP.

Preliminary TESTCODE analysis of pHi showed that the sequence contained several coding regions. A number of possible open reading frames, in both the sense and antisense directions were also identified. Subsequent DNA and protein searches using the open reading frames of coding regions showed significant sequence homology to known motility and chemotaxis genes of many bacteria. The 5' end of the sequenced region was a partial orf that when translated, showed homology with FlhA from many bacteria. Other ORFs displayed substantial sequence similarity to FliR, and CheL in the databases (see figure 3.4.2).

3.3 Sequencing pH2 plasmid.

The open reading frame which showed homology to FlhA contained only 120 amino acids, whilst this protein in other bacteria consists of about 680 to 707 amino acids. This result suggests that the open reading frame extend further in an adjacent fragment. To find the beginning of this open reading frame, a 1.7 kb BamHI-EcoRI fragment, upstream of the pHi fragment from plasmid pAB1, was subcloned into pBluescript SK+ (see figure 3.2.2b). This created a plasmid named pH2. The plasmid was fully sequenced in both direction and aligned to the pHi sequences (see figure 3.2.1b). TESTCODE analysis of the new alignment showed that the whole sequence of pH2 with about 350 bp of 5' end of pHi contained a large coding region. This result was further confirmed when open reading frame maps of the alignment, showed a large orf in this region. Homology analysis also showed a potential similarity between this ORF and FlhA of many bacteria (see table 3.5.1), however the beginning of the *Agrobacterium flhA* sequence was still missing. Next step of project, therefore was construction of missing part of *flhA*.
3.4 Construction and sequencing of plasmid pH3.

The 1.7 kb *BamHI-EcoRI* fragment pH2 (as can be seen in figure 3.2.1) includes the 5' end of the cosmid pDUB1905. This, together with the results obtained from homology searches, suggested that the beginning of the open reading frame showing homology to *flhA* gene was in the DNA fragment adjacent to the cosmid pDUB1905. Previous work conducted in this laboratory showed that cosmid pDUB1905 does not overlap with cosmid pDUB1900, which contains several flagellar genes. Therefore, reverse PCR was used to construct the beginning of *flhA* (see figure 3.4.1). A Southern blot experiment was carried out to find the appropriate restriction enzyme for digesting the *Agrobacterium* chromosomal DNA.

*Agrobacterium* genomic DNA was digested with different restriction enzymes include *HindIII, EcoRI, Hpal, HincII, BclI*, and *Sacl*. The digests were probed with a 0.8 kb pH2/*HindIII* fragment from pDUB1905. The result showed that *HindIII* is the most suitable site to use for reverse PCR.

Reverse PCR is used when the sequence of only one end of a fragment is known. In this method specific primers are designed the opposite direction. Two specific primers, which were used in the PCR, were designed from pH2. They were:

- **HSNI**: 5'-ITC AGC GAC AGA CGG ATC AT-3'  
  57 76
- **HSNII**: 5'-GCG GCA TCA TCA TCG GTT AT-3'  
  476 495

Using these oligonucleotides in the PCR reaction allowed synthesis of chromosomal DNA directly upstream of the 5' end of the cosmid pDUB1905.

The diagram of reverse PCR is shown in figure 3.4.1. *Agrobacterium* genomic DNA was digested with *HindIII*. The digest was religated and then cut with *HincII* to open the circle. Two specific primers (HSNI and HSNII) were applied to the PCR reaction samples containing the digest. The PCR program, described in section 2.5.7, was used. An approximately 0.3 kb fragment was synthesised. The PCR product was subcloned in SK$^+$ and called pH3. The complete length of the plasmid was sequenced in both directions. The plasmid pH3 sequence overlapped with the 5' end of pDUB1905, meaning that it was indeed a continuation of pDUB1905. Therefore, The pH3 sequence was aligned to the other sequences, creating a new alignment about 5.8 kb in length (see figure 3.2.1b). The complete sequence of the sense strand of 5.8 kb
Figure 3.4.1 Diagram outlining the main steps involved in reverse PCR to synthesise the begining of *flhA*.

The box with thick borders shows the fragment to be synthesised. Arrows within the circle and underneath the main figure indicate the primers and the direction that they act to synthesise the begining of *flhA*.

Dark shaded regions $\equiv$ pDUB1905.
Table 3.4.1 Some characteristic of potential genes and orf found in this work

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<th>Protein length (aa)</th>
<th>Molecular weight (kDa)</th>
<th>Isoelecteric point (Ip)</th>
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Figure 3.4.2 The complete sequence of the sense strand of 5.8 kb region of the cosmid pDUB1905.

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93
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GTCTGTGATC TCGACTGGAA TGGCGCGCTG GAACTGGTCC GCGGCAAGGC
TTCGGCCTGT GAAGGGCACA CGATGCTCTC CTTCTGACC ATCGAGAAGA
CAAAGTTGTC ACTCAAGGAT AATGCCTATC GGCAAAATTCT GGAAGCTGC
CTGCTTCTGC CGCAGGGCAA GGGCATGAAA GCCGCCGCGG GGGCGAACA
TGCCGAGCCT CTGCCCGCCA CCATCGATGG CGTGCCCCAC ACCATGGCGC
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GCAAGACTTC GTCATGCTGA ACCGGGACGC GCCCGCCGCTG AAGGTGGATG
TGCTCCTGGC AGGAAATGCTC AAGGAAATTC AGGAGACATG GCTGACCCGC
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CAAGGTCCTG TCCTCAGAAG TGGCGGGGTAT GCCGGACATT TTCCGCAAGC
TGCGATGAG CTGGCTCTAC AGCCTTTGTG CCGAACCCTG GCACATCGCG
CTGGCAAGA GCTGATATC
Figure 3.4.3 DNA Strider open reading frame map of the sequenced region of pDUB1905 showing the location and name given to each identified open reading frame.

Numbers on the vertical axis indicate the three different sense (1, 2 & 3) and antisense (-1, -2 & -3) reading frames. For each reading frame, short vertical lines indicate possible start sites (ATG or GTG), full length vertical lines indicate possible stop sites (TGA, TAA or TAG).
**Figure 3.4.4 Diagram of Testcode result of the sequenced region of pDUB1905.**

Regions above the upper horizontal line (9.4) are considered as coding regions with likelihood greater than 95%. Conversely, the regions below the lower horizontal lines (7.4) indicate with 95% confidence as non-coding regions of DNA by Testcode. Area between these two values (9.4 and 7.40, are considered as "window of vulnerability" which Testcode can not predict significantly. The horizontal arrows above the diagram indicate the genes and the *orf* found in this project.
Figure 3.4.5 Strategy for the partial sequencing of pDUB1905

The dark box in this figure represents only the sequenced region of pDUB1905 and a DNA fragment (H-B in the diagram) synthesised by PCR. The letters in bold above the box is showing the position of selected restriction enzyme sites. The position corresponding to the mutants, fla-8, mot-6 and che-2, have also shown above the box. The horizontal arrows below the box represent plasmids generated to sequence the region.
region of the cosmid 1905 is shown in figure 3.4.2. The open reading frame map and TESTCODE of this alignment are shown in figure 3.4.3 and 3.4.4. The strategy applied to fully sequence the alignment is shown in figure 3.4.5. Homology searches for possible motility or chemotaxis genes revealed that the alignment had similarity to different genes including $\textit{flhA}$, $\textit{flIR}$, and $\textit{cheL}$ in various bacteria. Some characteristics of the potential genes and orfs found in this work have been shown in table 3.4.1.

3.5 The $\textit{flhA}$ homologue:

Approximately 100 bp downstream of the 5' end of the alignment an open reading frame starts which shows to have a significant homology at the DNA and protein levels, to $\textit{flhA}$ gene of many bacteria containing both Gram-negative and Gram-positive bacteria. This open reading frame that finishes at position 2214 bp, is 2108 bp long and its protein product contains 703 amino acids, with predicted molecular weight of 76.86 kDa. Figure 3.5.1 shows the sequence of the $\textit{flhA}$ homologue along with the predicted gene product. The range of identity and similarity of homology between $\textit{Agrobacterium flhA}$ and the corresponding homologues in other genes were 40-60% and the similarity was 54-69% over at least 679 amino acids. FlhA is a member of the protein delivery system called the flagellum-specific export pathway/apparatus (see section 1.4.3).

The $\textit{Agrobacterium flhA}$ product was also found to have sequence similarity to the virulence proteins that are involved in the typeIII secretion protein apparatus in several bacteria, including $\textit{Yersinia pestis}$ and $\textit{Y. enterocolitica LcrD}$, $\textit{Shigella flexneri MxiA (VirH)}$, $\textit{Salmonella typhimurium InvA}$ and $\textit{SsaV}$, $\textit{Pseudomonas aeruginosa PcrD}$, and $\textit{Y4yR}$ a probable translocation protein from $\textit{Sinorhizobium sp. NGR234}$. Table 3.5.1 shows the proteins with sequence homology to FlhA. A multiple alignment was carried out using CLUSTALV at HGMP, the results of which are shown in figure 3.5.2. As can be seen, the sequence similarity between the ORF and the homologues is throughout their entire lengths, however it does not usually cover the first 32 amino acids.

A potential ribosome-binding site (rbs), that bears a weak similarity to the Shine-Dalgarno sequence, was found 19 bp upstream of the $\textit{flhA}$. There are also nucleotides that have similarity to the consensus ribosome binding site sequence 5 bp
after the start codon of the orf and 7 bp upstream of the second methionine codon (M). The size of the FlhA protein varies in different bacteria and is between 673 amino acids to 733 amino acids. In most cases the FlhA in other bacteria is shorter than the protein product of this supposed flhA. Therefore it is possible to speculate that the gene may start from the second M codon.

A possible transcription terminator, with a stem-loop motif, was identified upstream of flhA using the Macintosh program DNA Strider and the terminator program at SEQNET. The putative terminator consists of a stem with 8 bp and a loop of 3 bp. Finding a terminator in this region suggests that this sequence could be the first gene of an operon. flhA is often the middle gene of an operon starting with flhB as in C. crescentus, S. typhimurium, Y. entercolitica or che/fla in Bacillus subtilis (123, 172). Attempts to identify a promoter upstream of flhA were unsuccessful. Therefore, the translated sequence upstream of flhA was searched for significant homology to any other flagellar genes. No similarity was found in this region. This might be because the sequence, which contains only106 bp is too short to reveal any homology with relevant genes, or there is a gap between flhA and the previous gene, as can be seen in B. subtilis (37) where a gap of 32 bp exists between flhB and flhA.

The hydropathy profile of the A. tumefaciens FlhA homologue, using the Kyte and Doolittle algorithm, revealed a similar pattern to that of other known homologues. The resemblance of hydropathy pattern between A. tumefaciens FlhA and E. coli FlhA is less than FlhA from C. crescentus. They have seven membrane-spanning segments, while E. coli has only four transmembrane domains. The result showed that the amino acid sequence could be divided into two distinct domains. The amino-terminal half of the FlhA protein contains seven strongly hydrophobic regions, predicted to be membrane-spanning segments, suggesting that this protein could be an integral membrane protein. There was no cleavable signal sequence. In contrast, the carboxyl-terminal is quite hydrophilic and is likely to represent a water-soluble domain. Figure 3.5.3 shows the hydropathy profile of FlhA and four of its homologues from flagellum-specific export and typeIII export systems. The predicted transmembrane domains of the FlhA, using SOSUI program at HGMP is shown in table 3.5.2.
Figure 3.5.1 The entire coding and untranslated regions of flhA homologue and the predicted gene product.

The flhA coding regions begins at nucleotide position 106 and finishes at 2214. The possible start and stop codons are overlined and labelled flhA and "*" respectively. The start codon of next homologue, which overlaps flhA stop codon, giving an "ATGA" configuration, is also overlined and marked fliR.

HindIII (at position 1 bp) and BamHI (at position 308 bp) denote the beginning of the sequence (and also PCR product) and beginning of cosmid pDUB1905 respectively. Two possible ribosome-binding sites, 19 bp before and 10 bp after the start codon, are overlined and labelled "rbs". The potential stem-loop structure, which may be a transcription termination signal is also shown, by staggered arrows and marked "TM".

HindIII
1 aag ctt cag cgg cta ggc tga agg cat cgg ccc gaa ttc cgt ctt ttc gaa 60

rbs
61 aac cga tgc ccg gaa ccc tgc gac ggc ttc gtc gaa

flhA
121 GAA TTC ATG GCG CAG CCA CCA GTC ATC TCC TGG CCC AAG GTC AGT CCG AGC ATG GAT 180

BamHI
301 GCG CTC TGG ATC CAG CCG CCG CTG CGC TAC TGG TGA TGG GCG ACC GTC ATG ATC GCC 360

421 GAG GGG CCG AGC GGG GCA GTG GAC GTG TTC TCC AGC CTC GTC ATG TCC GGT 480

481 GAC TTC GTC ATC GGT CTC ATC GTC TTC CTG ATC CTG ATC ACG GTC AAC TTC ATC GTC ATC 540

102
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Table 3.5.2 Transmembrane domain of *A. tumefaciens* FlhA.

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Figure 3.52 Multiple alignment (pretty box) of the potential A. tumefaciens FlhA to those of flagellum-specific export component and type III export proteins

Abbreviation used for bacteria are as follow: AGRO = A. tumefaciens, ECOL = E. coli, BACSU = Bacillus subtilis, YERPE = Yersinia pestis, YEREN = Y. enterocolitica, SHIF = Shigella flexneri.
Figure 3.5.3 Hydropathy profiles of the *A. tumefaciens* FlhA homologue and FlhA from other bacteria, LcrD and MxiA from type III export system.

FlhA from *A. tumefaciens*

FlhA from *E. coli*

FlhA from *C. crescentus*

LcrD from *Yersinia pestis*

MxiA from *Shigella flexneri*
3.6 The fliR homologue:

There is an open reading frame starting immediately downstream of flhA, at position 2211 and finishing at position 2966. The open reading frame spans a region of chromosomal DNA which contains two behavioural mutants fla-8 and mot-6. These two mutants, which were generated by insertion of Tn5 transposon, share an identical site. The putative start codon of this orf, which has 756 bp (252 amino acids), with predicted a molecular weight of 26.87 kDa, overlaps with the stop codon of flhA by 3 bp, to give an "ATGA" configuration.

Two putative ribosome-binding sequences (TGGG and AGGG) resembling the Shine-Dalgarno sequence were identified 9 and 30 bp upstream of the fliR homologue respectively. The latter is unlikely to be a ribosome-binding site as it is too far from the start codon. There was no terminator-like sequence downstream of the fliR stop codon. The DNA sequence of the fliR homologue along with the predicted gene product and potential ribosome-binding sites are shown in figure 3.6.1.

The product of the open reading frame, which is transcribed in the same direction as flhA, showed sequence similarity with FliR of several bacteria, at the protein level. It was, therefore given the name fliR. The FliR protein is a flagellar biosynthetic protein, and like FlhA is a member of the flagellum-specific export proteins that are involved in transport of extracellular flagellar proteins. A search of the GenBank database showed the A. tumefaciens FliR to have significant homology to the Erwinia carotovora MopE, Erwinia herbicola and Erwinia amyloyora HrcT, Shigella flexneri and S. sonnei SpaR (Spa29), Chlamydia pneumonia YopT, Yersinia enterocolitica YscT, S. typhimurium SsaT, Sinorhizobium fredii Y4yN, Pseudomonas syringae pv. syringae HrpX, all of which are involved in secretion of virulence factors into plant and animal cells. The range of identity between A. tumefaciens FliR and other homologue proteins was 20-29% and similarity was 38-51%, over at least 213 amino acids. The size of FliR proteins varies between 252 to 277 amino acids in different bacteria. Table 3.6.1 shows the proteins with sequence homology to the Agrobacterium FliR. A multiple alignment of the predicted A. tumefaciens FliR protein with their homologues was also carried out, which is shown in figure 3.6.2.

The fliR gene from Caulobacter crescentus is a member of fliQR operon which consists of only fliQ and fliR (238). The E. coli fliR protein is located in the fliL operon, and it is required for flagellar biogenesis (127).
Figure 3.6.1 The untranslated and coding regions of fliR homologue with its predicted translated product.

The fliR homologue starts immediately after flhA at position 2211 bp and finishes at position 2966 bp. The start codon of the gene is underlined and labelled fliR, while the stop codon shown by "*".

Two potential ribosome binding sites 9 bp and 30 bp upstream of the start codon are underlined and labelled "rbs".

```
2124 tat cga ggc act ctt tgc gac cct gcc ggt tct ttc aca tgt gga act ggc caa ggg tct 2183

rbs

2184 cga gat cca gat tct ggg cgg cct ttc ATG ATA ACC GAC CCG GGA ACA ATC ATT GCA 2243

fliR

M I T D P Q G T I I A 11

2244 TTG TTC CTT GCC ATC TGC CGC ATA GCC CCC ATC GCC GCA ATG TAT GGG CTG ATC GCC CGT TTC TAT 2303

12 L F L A I C R I G A C F M T M P G F S S 31

2304 TCG CGC ATC TCG CCG CAG ATT CGC ATT TCG GTG GTA GGA GTA TCC ATG GCG CTT CTG 2363

32 S R I S P Q I R M L L C V A V S M A L L 51

2364 CCC GTC CTC TGC GAC ACC ATC TAC CCA AAG GTC TCC GGT GCG AGC CAG GGC CGC GTC GGT 2423

52 P V L W D T I Y P K V S G A S Q G A V V 71

2424 GGC CTC ATC TTC TCC GAG GTT CTC ATC GGC GCA ATG TAT GGG CTG ATC GCC CGT TTC TAT 2483

72 G L I F S E V L I G A M Y G L I A R F Y 91

2484 AGC CTC GGT TTC CAG TCC ACC GGT GCG CTC ATC GCC GCT TCC ATC GGC CTC ATG GCG CCC 2543

92 T L G F Q F T G A L I G A S I G L S A P 101

2544 GGC GGT GCC GAT CCC ATC GAA GAC GTG CAG GAA AAC CAG ATC GCC AAT TTC ATC ACC TTC 2603

102 G G A D P I E D V Q E N Q I A N F I T F 123

2604 GGT GCC CTC CGT CTG GTG TTC ATG ATG GTT TTC CAC ATC GTC CTG AAG GCG CTG GTC 2663

122 G G L L V L F M M D F H H I V L K A L V 143

2664 GAT TCC TAT AGC GCC ACA CCG GTG GTG CTC ATC GGA CAG AAG ATG CTG ATC ACG 2623

144 D S Y S A T P V G A L I S G Q K M L I T 163

2724 CTG AGC GAT AGC CTC AGG GCC TTC TCG ATC ATG CTG CCG CTG GCG ACG ACC CCC TTT GGT 2683

164 L T D T L R A S F S I M L R L A S P F V 183
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**NdeI**

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184 I Y G H M F N V A V G L I N K L A P Q I 203

2844 CCG GTC TTC ATA TCG ACA CCC TTC GTG GCG GGT GGT CTT TTC ATG CTT TAT TTG 2803
204 P V F F I S T P F V L A G G L F M L Y L 223

2904 TCG GTC GCG GCC CTC ATC CGG CAA TTC GTG GAT GGG TTC GGC CCG GTC TTT ATC GCC TTT 2863
224 S V A G L I R Q F V D G P V F I G F 243

2964 TGA
244 *

2966
244
Table 3.6.1 Proteins with homology to potential *A. tumefaciens* FliR

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<td>51/231</td>
<td>251</td>
<td>Flagellar biosynthetic protein</td>
</tr>
<tr>
<td><em>Pseudomonas putida</em></td>
<td>FliR</td>
<td>25</td>
<td>49/244</td>
<td>258</td>
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</tr>
<tr>
<td><em>Escherichia coli</em></td>
<td>FliR</td>
<td>26</td>
<td>44/217</td>
<td>261</td>
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</tr>
<tr>
<td><em>Erwinia carotovora</em></td>
<td>FliR</td>
<td>27</td>
<td>43/219</td>
<td>261</td>
<td>Flagellar biosynthetic protein</td>
</tr>
<tr>
<td><em>Rhodobacter sphaeroides</em></td>
<td>FliR</td>
<td>25</td>
<td>40/228</td>
<td>269</td>
<td>Flagellar biosynthetic protein</td>
</tr>
<tr>
<td><em>Salmonella typhimurium</em></td>
<td>FliR</td>
<td>26</td>
<td>44/230</td>
<td>264</td>
<td>Flagellar biosynthetic protein</td>
</tr>
<tr>
<td><em>Bacillus subtilis</em></td>
<td>FliR</td>
<td>25</td>
<td>42/227</td>
<td>259</td>
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</tr>
<tr>
<td><em>Aquifex aeolicus</em></td>
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<td>26</td>
<td>43/250</td>
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<tr>
<td><em>Borrelia burgdorferi</em></td>
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<td>25</td>
<td>40/178</td>
<td>269</td>
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</tr>
<tr>
<td><em>Vibrio parahaemlyticus</em></td>
<td>FliR</td>
<td>23</td>
<td>42/231</td>
<td>260</td>
<td>Flagellar biosynthetic protein</td>
</tr>
<tr>
<td><em>Helobacter pylori</em></td>
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<td>20</td>
<td>38/236</td>
<td>255</td>
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<tr>
<td><em>Erwinia amylovora</em></td>
<td>HrcT</td>
<td>26</td>
<td>43/230</td>
<td>265</td>
<td>Type III secretion protein</td>
</tr>
<tr>
<td><em>Shigella flexneri</em></td>
<td>SpaR*</td>
<td>22</td>
<td>44/214</td>
<td>256</td>
<td>Surface presentation of antigen</td>
</tr>
<tr>
<td><em>Chlamydia trachomatis</em></td>
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<td>22</td>
<td>44/174</td>
<td>289</td>
<td>Translocation protein</td>
</tr>
<tr>
<td><em>Salmonella typhimurium</em></td>
<td>SpaR</td>
<td>23</td>
<td>42/205</td>
<td>263</td>
<td>Surface presentation of antigen</td>
</tr>
<tr>
<td><em>Erwina carotovora</em></td>
<td>FliR**</td>
<td>27</td>
<td>42/172</td>
<td>180</td>
<td>Type III secretion protein</td>
</tr>
<tr>
<td><em>Sinorhizobium sp. NGR234</em></td>
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<td>23</td>
<td>27/217</td>
<td>272</td>
<td>Translocation protein</td>
</tr>
<tr>
<td><em>Yersinia pestis</em></td>
<td>YopT</td>
<td>20</td>
<td>42/230</td>
<td>261</td>
<td>Type III secretion protein</td>
</tr>
</tbody>
</table>

* formerly Spa29
** formerly MopE (as mentioned in the text)
Table 3.6.2 Transmembrane domains of *A. tumefaciens* FliR.

<table>
<thead>
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<th>Transmembrane NO.</th>
<th>N terminal</th>
<th>Transmembrane region</th>
<th>C terminal</th>
<th>length</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6</td>
<td>QGTIIALFLAICRIGACFMTMPG</td>
<td>28</td>
<td>23</td>
</tr>
<tr>
<td>2</td>
<td>36</td>
<td>PQIRMLLCVAVSMALLPVLWDTI</td>
<td>58</td>
<td>23</td>
</tr>
<tr>
<td>3</td>
<td>64</td>
<td>GASQGAVVGLIFSEVLAGAMYGL</td>
<td>86</td>
<td>23</td>
</tr>
<tr>
<td>4</td>
<td>120</td>
<td>FITFGGLLVLFMMDFHHVILKAL</td>
<td>142</td>
<td>23</td>
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<tr>
<td>5</td>
<td>161</td>
<td>LITLTDTLRASFSIMLRLASPFV</td>
<td>183</td>
<td>23</td>
</tr>
<tr>
<td>6</td>
<td>211</td>
<td>PFVLAGGLFMLLYLSVAGLIRQVF</td>
<td>233</td>
<td>23</td>
</tr>
</tbody>
</table>
Figure 3.6.2 Multiple alignment (pretty box) of the potential *Atumefaciens* Flir to those of flagellum-specific export system and type III secretion proteins.

Abbreviation used for bacteria are as follow: AGRO = *A. tumefaciens*, ECOLI = *E. coli*, BACSU = *Bacillus subtilis*, ERWCA = *Erwinia carotovora*, YEREN = *Yersinia enterocolitica*, SALTY = *S. typhimurium*.
Figur 3.6.3 Hydropathy profiles of the A. tumefaciens FliR homologue and FliR of E. coli, C. crescentus, S. typhimurium and MopE of Erwinia carotovora from type III export system.
*fliR* is a part of a large flagellar gene cluster containing seven additional genes encoding FliL, O, P, Q, R and B (75). The *Erwinia carotovora mopE* gene is in the *mop* operon (142). Mutations in this operon lead to a reduction of virulence in plants. In addition, a mutation in the *mopE* gene results in a non-flagellated phenotype. The *Shigella flexneri spaR* (*spa29*) gene resides in an operon (174), which encodes several genes that are essential for invasion of host cell and surface presentation of the virulence proteins. The function of *yscT* in *Yersinia pestis* has not been reported, however it is known that homologues of this gene in *Y. pseudotuberculosis* is essential for export of Yop proteins (92).

The hydropathy profile of the *A. tumefaciens* FliR homologue and the two other homologues, using the Kyte and Doolittle algorithm (shown in figure 3.6.3) revealed that they have similar patterns. It showed FliR to be very hydrophobic throughout almost the entire polypeptide sequence, and contain at least six membrane-spanning domains, suggesting that it is a membrane protein. The predicted transmembrane domains of the FliR, using SOSUI program at HGMP is shown in table 3.6.2.

**3.7 orfX**

About 16 bp downstream of the *fliR* homologue an *orf* 756 bp long, starting at position 2982 bp and finishing at position 3401 bp was identified. The gene product of this *orf* (*orfX*) contains 139 amino acids, with molecular mass of 16.053 kDa. TESTCODE analysis predicted, at the 95% confidence level, that this *orf* is a coding region. The guanine and cytosine content of the *orf* is 54.7% and this is significantly higher than those of non-coding regions (see table 3.4.1). In addition, a potential ribosome-binding site was found 5 bp upstream of the start codon of *orfX*. However, the GenBank database search showed no significant homology to any previously identified proteins. This might be because no relevant protein has yet been found in other bacteria. The whole DNA sequence of *orfX* together with its possible protein sequence is shown in figure 3.7.1. The hydropathy diagram of ORFX, which has been shown in figure 3.7.2, has a very hydrophilic profile.
Figure 3.7.1 The DNA sequence of orfX with its putative translated protein and DNA sequence upstream of orfX.

orfX starts at position 2983 bp and finishes at position 3401 bp. The initiation codon of the orf is underlined and labelled orfX, while the stop codons shown by "*".

A potential ribosome binding site "GAGGA" 7 bp upstream of the start codon is underlined and marked "rbs".

```
2964 tga tgc gga gga aac gcc
   ATG GCT TCG GAC AAG CGC TCG GAA AAG CTC AAG CGT CTG GTA
   M A S D K R S E K L K R L V

3024 ACG GTC CAG CGG CAT ATG GAA AAA ATG GCC GAG GTG GAA CTG GCC GAT ACC ACT CGC GTG
   15 T V Q R H M E K M A E V E L A D T T R V

3084 CGC AGC GAA GTG GCCCAA TCC ATG GAG ACG TTG GAG GCC ATG AGT TCC ATG GAA CCG
   35 R S E V A Q S M E S T F E A M S S M E P

3144 GTG CAT CAG ACC TTC ACC AAT TCG GAC CGC TAC AGC CGC CTG GTC GTC CAG GAC
   55 V H Q T F S K H Y S D R Y S R L V V Q D

3204 CGC CAG CTC GAA GGC GTT CAG CAG TTT CAG GAA AAG GTC CTG AAG GAA AAG ACC AAG
   75 R Q L E G V Q F Q E N K V L K E K T K

3264 GCT GAC AGG CTG GAG GAC AGG ATG CAT ATT GCC CGC GAT GTC GAG GAT CGC GAG GCC GTT
   95 A D R L E D R M H I A R D L E D R E A G

3324 GAT AAT GCA ATT TAC GAT TGT TTA GAA ATT ACG AAT GTT TCC CAC ACA CCA GCC TCC AGC
   115 D N A I Y D L L E I T N V S H T P A S S

3384 AAG GTT GCC GAT CCA TAG
   135 K V G D P *

3401
```

118
3.8 The cheL homologue:

The cheL homologue is a 532 bp open reading frame that in the TESTCODE analysis was shown to be a coding region. The orf, which spans the che-2 mutant, starts at nucleotide 3644 (about 240 bp downstream of orfX) and finishes at nucleotide 4073 bp. The gene product of the orf is 144 amino acids long, with a predicted molecular weight of 15.66 kDa. It has sequence homology to the CheL protein of C. crescentus (28% identity and 53% similarity, over 75 amino acids). The Caulobacter CheL protein is the only CheL protein which has been found in the bacteria so far therefore this ORF is the second protein of this kind to be reported. The whole DNA sequence of cheL together with its possible protein sequence is shown in figure 3.8.1, while the GAP alignment of the possible Agrobacterium CheL, with the Caulobacter CheL, is shown in figure 3.8.2. Pairwise alignment between the two CheLs has revealed that there are four major conserved areas within the two proteins. The conserved regions are between residues 34 and 96 of the 111 amino acids. The hydropathy profile, according to the Kyte and Doolittle algorithm, of the two proteins is shown in figure 3.8.3. The CheL of A. tumefaciens is significantly longer than its counterpart in C. crescentus (about 60 amino acids).

The DNA sequence upstream and downstream of cheL was analysed to try to find any promoter or terminator. No sequence indicating the presence of a terminator downstream of orf was found. A stretch of sequence, which overlaps the initiation codon of the cheL showed similarity to the A. tumefaciens putative flagellar promoter (GCCAATG) and -10 class II flagellar promoter from E.coli (GCCGATAA) (55), but no typical -35 consensus sequence preceded this sequence. A sequence (ACCAG) weakly resembling a Shine-Dalgarno motif was also found, 8 bp upstream of the start codon of cheL, this may therefore be a weak ribosome-binding site.
Figure 3.8.1 The DNA sequence of *cheL* with its putative protein and DNA region upstream of it.

*cheL* begins at position 3644 bp and finishes at position 4073 bp. The start codon of the gene is underlined and labelled *cheL*, while the stop codon shown by "*". Tn5 insertion site is also underlined and labelled Tn5.

A potential ribosome binding site 8 bp upstream of the start codon is underlined and marked "rbs". The putative promoter, which overlaps the start codon of the gene is also double underlined and labelled "PR".

```plaintext
3547 gtg gtt cgt cgc cgga ccc gaa gta cgt gtt gcc cag gaa cta gca 3606
  rbs PR  cheL
3607 aac aag ggg gct ttt gcc gaa cca gta tgg cgg ATG CCC GCA AGG GCT TTG GTG CTG CCG 3668
1
3669 TCG ATA TGC TTT GAC GGT TGG TCG AAG GCC GCC GCT TTT GGC GTT ATT GCC TCG 3728
10 SC IRC TGA S T N I R S 29
3729 GCA CGT ACC GAG ATT CCC GAA GAC GAC GGA GGA GTT TAC GCC AAG GGT AAT GCC GCC GAA 3788
30 A R T E I P E T Y R K Y H A L Q V T F 49
3789 GTT GCC AAC ATG CTG CCG AAA GAC AGG GAG GAA GTT TAC GGC AAG GGT AAT GCC GCC GAA 3828
50 V A N M L P K D S E E V Y G K G N A G E 69
3829 ATC TGG AAG AGC ATG ATG GCG GAA CAG TTT GCC GAC ACT ATT TCC AGA AAT GCC GCC GTC 3888
70 W K S M A E Q F A D T I S R N G V 89
71 G I A E Q A Y K D A L R K A E S K G I T 109
3889 GGC ATT GCG GAG CAG GCT TAC AAG GAT GCG CTG CCG AAA GCC GAA AGC AAA GCC ATT ACC 3988
90 G I A E Q A Y K D A L R K A E S K G I T 109
3929 GAC GTG TCG ATG AAT GAC AAA GAC CAC AAT GCT GCA ATT CCAG ATG GTG GCG GAG TTC GAG 3988
110 D V S N D K D H N A A I R M V A E F E 129
3989 CCG CAG GTC CTC GGC GGT TCC AAT GAT AAA AAC GAC GAG GCT TGA
130 R Q V L G V S N D K T D E A * 4073
144
```
Figure 3.8.2 Pairwise alignment (pretty box) of potential CheL of *A. tumefaciens* and *C. crescentus*.

Abbreviation used for bacteria are as follow: AGRO = *A. tumefaciens* and CAUC = *C. crescentus*.
3.8.3 Hydropathy profiles of the *A. tumefaciens* CheL homologue and its counterparts in *C. crescentus*.

![Hydropathy profiles](image)

3.9. *OrfY, orfW and orfZ*:

The sequence downstream of *cheL* contained three open reading frames, *orfY*, *orfW* and *orfZ*. All open reading frames were predicted to be coding regions by TESTCODE analysis, at the 95% confidence level. Moreover they had guanine and cytosine content of 54.86%, 57.4% and 60.89% respectively, significantly higher than
that of non-coding regions (see table 3.4.1). However, the orfs have no significant sequence identity to any previously identified proteins in the GeneBank database.

**OrfY** begins just 2 bp downstream of *cheL* at position 4078 bp and finishes at position 4473 bp. It has 396 base pairs and contains 132 amino acids, with a predicted molecular mass of 14.72 kDa. A sequence (GAGG) found 7 bp upstream of the start codon could be a ribosome-binding site. No possible promoter or terminator was found upstream or downstream of *orfY*. The hydropathy profile shows that *orfY* is extremely hydrophilic.

The second open reading frame, **orfW** starts at position 4471 and ends at position 4923. This open reading frame contains 453 bp and 151 amino acids. The start codon of open reading frame W overlaps the stop codon of *orfY*, giving a "TGATG" configuration. Two motifs that resemble Shine-Dalgarno sequence are found 12 bp and 26 bp upstream of start codon of this orf; these may therefore be possible ribosome-binding sites.

There is a gap of 168 bp between *orfW* and the third open reading frame, **orfZ**, which was predicted as a non-coding region by TESTCODE analysis. A sequence-like-terminator, which consists of a 9 bp stem and a 3 bp loop was identified 47 bp upstream of *orfZ* in this region. If this sequence is a real terminator, it is therefore possible to postulate that *cheL* along with *orfY* and *orfW* constitute an operon as they overlap or one close to each other. Moreover, the existence of a possible -10 classII promoter-like sequence upstream of *cheL*, increases the probability of these being an operon in this region.

**OrfZ** starts at nucleotide 5191 and extends to the end of the sequence. It is 675 bp long and contains 225 amino acids. No typical ribosome-binding sequence was found upstream of the orf, however a TGGG that could be a ribosome-binding site, was identified 9 bp downstream of start codon. A -10 box classII promoter sequence, GGCGAT, with one bp mismatch and a -35 classII sequence, TCATGA, were also found 24 and 63 bp upstream of *orfZ*.

The DNA sequence of the three orfs and their deduced amino acids is shown in figure 3.9.1 to 3.9.3. The hydropathy profiles of these orfs, according to the Kyte and Doolittle algorithm, are shown in figure 3.9.4.
Figure 3.9.1 The DNA nucleotide and deduced amino acid sequence of *orfY* with untranslated region upstream of the *orfY*

*OrfY* begins at nucleotide 4077, two bp downstream of *cheL* homologue, and finishing at nucleotide 4472. Start codon is underlined and labelled "*orfY*", while stop codon indicated by asterisk. Stop codon of previous gene is also underlined and shown by double asterisks.

A potential ribosome binding site is underlined and labelled "rbs"
Figure 3.9.2 The DNA nucleotide and deduced amino acid sequence of orfW with untranslated region upstream of the orf.

OrfW starts at position 4471, which overlaps the stop codon of orfY, and finishing at position 4472 bp. The start codon is underlined and labelled "orfW", while stop codon indicated by asterisk. Previous orf stop codon is underlined and shown by double asterisk. Two potential ribosome-binding sites 12 (ACGG) and 26 (AGGA) bp upstream of the orf start codon are underlined and labelled "rbs".

```
orfW
4400 gaa cgc tat tca gga agc cga tac gga cgg cac cta ttc cca gga aca att cct tta cgg 4459
4460 cgc agc ctc ctc AGG TTT AGG AGT CTG CCT ATG ATG ATG ATG ATG ATG ATG ATG ATG ATG 4519
   1 ** M L K L L L T G V W V C A V T L 16

4520 GCC GCG GTG TAT TTC TCC GTC TAT ATG ATG ATG ATG ATG ATG ATG ATG ATG ATG ATG 4579
   17 G A V Y F S V Q M A T A P A P D E A G A 36

4580 AAA AAG GCT GAT CTG CAA GTG GTC AAG GGC GAA AAG ATC GAG ATG CCC ATG AAT GAT 4639
   37 K K A D L Q L V K G E S I T I P V I N D 56

4640 GGT GGA GTG AAC GGC TAT TTC GTC CTC GTC GTC GTC GTC GTC GTC GTC GTC GTC GTC 4699
   57 G G V N G Y F L S R I S L R V D K A K M 76

4700 GCG AAG ATC GAA CTG CCG GCA ACA CAG CTG ATG ACC GAT GAA CTG TTA GTC GTC GTC 4759
   77 A K I E L P A T Q L M T D E L F T L L A 96

4760 GGT TCT TCC ATG ATG TAC ATC ACC ATT GCC AAT ATC TCC ACC ATC GCC GAA GCC TTC AAG CAG 4819
   97 G S S M V N I A N I S T P D P E A F K Q 116

4820 GCC ATC CGC GAA GGG TTT AAG AAG AGG CTC GAT GAC GAG GTC GTC GTC GTC GTC GTC GTC 4879
   117 R I R E G L N K R L D D E V V E D V L I 136

4880 GAG CAG CTC GAT TAC CTG TCC GAA AGC CGA CAT CCC GCA GAA GAA CAA TGG TTC GCC 4939
   137 E Q L D Y L F E S R H P A E G Q W F A 156

4940 CCA CTC CTG TGT CAA ACT CTG CTA GAA AGC CAT CCC GCA GAA GAA CAA TGG TTC GCC 4999
   157 P L C Q T R R G K S R S R G C G T Q 176

5000 CCA CTG AGG AGG GAG CGG CGG CGA AAA AGC AGC AGG AGA GGC TGG GCC ACC CAG 5059
   177 P L T E R R R N G T L V N H L L L H 196

5060 TCA CGA GCA TTC GAG AGA TTT CTT CCA AGT TAA 5093
   197 S R A F E R L L S * 208
```
Figure 3.9.3 The DNA nucleotide and deduced amino acid sequence of orfZ with untranslated region upstream of the orf.

OrfZ starts at position 5188 and finishing at position 5731 bp. The start codon is underlined and labelled "orfZ", while stop codon indicated by asterisk.

A potential ribosome binding sites (TGGG), 8 bp upstream of the orf start codon are underlined and labelled "rbs". Approximately 24 and 63 bp upstream of start codon are a putative -10 and -35 promoter sequences underlined and labelled "-10 and -35".

The potential stem-loop structure, which may be a transcription termination signal, is also shown between nucleotides 5140-5160 by staggered arrows and labelled “TM”.

```
5092 aat tca acc tgc acc tgt at a aca aac tca tga acc acc gcc ccg cag gcc atc CGC CGG 5152
-35
5152 gcc ggt ggc gat ggg ttt ggc att tgg ggg tgc a ATG AAT TTC GCG TCA GGT GTT ACG 5211
-10
5212 GCC CGG GAG TTT CAA CGG AAT ATC CAC GGG CTG CGC GTC GTC GCT GTG GAT GTC GAG GAA CAT GCG 5271
9 R R G L Q R N I H G L R V C D L D W N G 28
5272 GCC CTG GAA CTG GTC AGC GCC GCG AAG GCT TCG GCC TGT GAA GGG CAC AGC ATC TCC TCC TCC 5331
29 A L E L V S G K A S A C E G H T M L S F 48
5332 CTG ACC ATC GAG AAG ACA AAG TTT TCA TCA ATG GAT AAT GCC TAT CGGCAA ATT CTG GAA 5391
49 L T I E K T K L S L K D N A Y R Q I L E 68
5392 AGC TGC CTG CTT CTG CCG CAG GGC AAC GGC ATG AAA GCC GCC CGG GCC GAA CAT GCC 5451
69 S C L L L P Q G K G M K A A A R A E H G 88
5452 GAG CCT CTG CCC GCC ACC ATC GAT GCC GTT GCC TTC ACC ATG GCG CTT CTG ACC TAC ATG 5511
79 E P L P A T I D G V A F T M A L L T Y M 108
5512 GCT GTG CCG AAA GCC ATC GCC GTG GCG GCC GAC GAT GCG CGG CAG ATC GGT GAA GTG 5571
99 A V P K R I G V A D D A A Q I G E V L 128
5572 GCC CGG CTG CTT GCC CAT GCG CCA TGG CAA GAC TTC ATG GCG GTT ACC ATT GCA GTC GGG CCC CTG 5631
119 A R L R A H A P W Q D F V M L N R D A A 148
5632 GCC GTG AAG GTG GAT GAT GTC CTG GCA GGA ATG CTC AAG GAA AAT CAG GAG ACA TGG CTG 5692
139 G V K V D V L L A G M L K E N Q E T W L 168
5692 CAC CGC AGC GTC AGC GGT GAG GAT GCG GCT GCC TCC ATT GCA GTC GGG CCC CTG TCC AAG 5751
159 H R S V S R E D A R V T I A V G P L F K 188
5752 CTG CTG TCC TCA GAA GTG GCC GGT ATG CCG GAC ATT TTC CGC AAG CTG CAT ATG AGC TGG 5811
179 V L S S E V A G M P D I F R K L H M S W 208
```
Figure 3.9.4 Hydropathy profiles of the three unknown ORFs, ORFY, ORFW and ORFZ in *A. tumefaciens*.
3.10 Discussion

Previous work in this lab showed that the chemotaxis and motility genes are clustered together in *A. tumefaciens*. They showed three cosmids, pDUB1900, pDUB1905 and pDUB1911 to contain many of the behavioural genes (34, 180, 228). The genes on the cosmids pDUB1900 and pDUB1911 have already been reported (56, 57, 228). The data presented here describes the identification and sequencing of two flagellar homologues, one chemotactic gene and four open reading frames encoding potential proteins without significant sequence similarity to any other proteins in the GenBank database. All are located on the cosmid pDUB1905.

Based on data obtained from sequence analysis, it is possible to postulate that the flagellar and chemotaxis homologues along with the three unknown open reading frames, *orfX*, *orfY* and *orfW* are transcribed together. This hypothesis is supported by the existence of a transcription termination sequence at the start of the sequence and another terminator in the end of the *orfW* sequence. However no classical promoter was found upstream of the possible operon. It is also tempting to speculate that the sequence can be divided into at least two distinct operons. One operon consists of the *cheL* homologue and the two unidentified *orfs*, *orfY* and *orfW*. The *flhA* and *fliR* genes with or without *orfX* could constitute a second operon, or may be members of a larger operon. This speculation comes from the fact that a rather big gap exists between the two assumed operons and the genes in the operons overlap or are close to each other. The presence of a terminator upstream of *flhA*, a −10 element class II promoter-like sequence upstream of *cheL* and a terminator downstream of *orfW* are further indications of the existence of these operons. On the other hand, no promoter was found upstream of *flhA*, this might be because the sequence upstream was not enough to reveal any promoter, or relevant genes. In *C. crescentus*, it has been shown that a sigma 54 promoter controls transcription of *flhA* (171). Also no terminator was found downstream of *fliR* or *orfX*. The last *orf*, *orfZ*, appears to belong to another operon or is translated individually, since a possible terminator and a promoter was found upstream of this open reading frame. Sequencing downstream of this *orf* and upstream of *flhA* would be valuable to try to determine whether or not *orfZ* or *flhA* are part of an operon, by looking for possible genes or other information. It has been shown that the *Agrobacterium* motility and chemotaxis gene cluster has considerable
similarity with the closely related bacterium from the α-subgroup, *S. meliloti* (190). Most of the flagellar genes (*fla, flg, flh,* and *fli*) and motility genes are in the same order in the two clusters. However, there is a difference between the two clusters. The *che* operon in the *Sinorhizobium* cluster is linked downstream of *flhB*, while downstream of the *Agrobacterium flhB* are the fourth flagellin gene, *flaD* and the *motBCD* operon. Despite this difference between the two clusters, the gene order within each group (*fla, fli, flh* and *che*) appears identical. Therefore, it is possible to predict that the order of genes found in this work is also the same in both bacteria. Although, *flhA, fliR* and *cheL* have not yet been identified in *Sinorhizobium*, but it has been shown that a fragment from *Agrobacterium* containing the *che-2* mutant, hybridised to a 3.7 kb fragment of *S. leguminosarium biovar viciae* (55).

The protein product of the *A. tumefaciens flhA* homologue had significant sequence similarity to the FlhA proteins of several bacteria in GenBank (see section 3.5). The predicted *flhA* product has similar features with other FlhA proteins. It is slightly acidic (pI=6.679), and has 703 amino acids that is in the range of other FlhA proteins (Table 3.5.). The amino acid sequence is divided into two domains with the N-terminal half consisting of six strongly hydrophobic regions that are predicted to be a membrane-spanning segment, it is therefore predicted to be an integral membrane protein. On the other hand, the carboxy-terminus end of FlhA is hydrophilic. The high similarity between *Agrobacterium* FlhA and other FlhAs was found in the N-terminal half. *A. tumefaciens* is therefore predicted to have the same structure and function as FlhA in other bacteria. *flhA* is classified as a class II gene of the regulatory hierarchy genes in *E. coli* and *S. typhimurium* (122). This class of genes is required for flagellar assembly and function. FlhA along with at least eight other genes constitute the flagellum-specific export apparatus (see section 1.4.3), which are responsible for transport of the protein components of the extracellular structures, such as the hook and filament proteins, across the membrane. It has been shown that FlhA in *S. typhimurium* is a membrane-associated protein. There is evidence that FlhA is associated with the basal body. FlhA has large soluble domains that presumably protrude into the cytoplasm within the cavity that lies at the centre of the cytoplasmic C ring of the motor (138). It has been demonstrated that FlhA has a regulatory activity in addition to its role in flagellar transportation. In a nonmotile mutant of *Proteus* lacking FlhA, expression of classII and classIII flagellum genes is suppressed (80).
The protein product of the *A. tumefaciens* flhA homologue also showed significant homology to a family of typeIII proteins involved in virulence from a variety of animal and plant pathogens (see table 3.5.2). The similarity between proteins involved in the flagellum-specific export pathway in several gram-negative and gram-positive bacteria, and proteins from typeIII secretion systems is not surprising given, they share similar overall structural features and are involved in transport and localisation of the component proteins of the extracellular structure across the cell membrane by a mechanism that does not require signal cleavage. They have high similarity in their hydrophobic amino-terminus domain and less similarity in their C-terminus domain. Perhaps the carboxy-terminal of these proteins functions in different manners in the diverse environment that the bacteria live. InvA from *S. typhimurium* is required for the ability of *Salmonella* to invade cultured epithelial cells. Secondary-structure analysis of InvA (74) predicts that there are seven transmembrane domains located in the amino-terminal half of the protein, and a largely hydrophilic carboxy-terminus that is located in the cytoplasm (74). LcrD, which exhibits a virulence property called low-Ca\(^{2+}\) response (lcr), is necessary for expression of genes encoding a number of Yop proteins that are required for virulence and host invasion. It has been shown that LcrD contains eight possible membrane-spanning segments in its hydrophobic amino-terminal region (159). Other proteins such as MxiA of *S. flexneri* (former VirH), which is necessary for secretion of the Ipa invasion proteins, and HrpO, HrpI and HrpC2 that are all involved in virulence, have the same structure as LcrD (92). Although, members of this family appear to have similar structures, they are involved in diverse cellular processes, including virulence in *Y. pestis*, host invasion in *S. typhimurium*, and flagellar biogenesis in *C. crescentus*, *E. coli* and *B. subtilis*. Taken together, the size, the sequence homology, and protein structural features between the possible *Agrobacterium* FlhA, other FlhA and virulence proteins in the other bacteria, leads to the conclusion that this region of pDUB1905 contains the *A. tumefaciens* flhA gene.

The predicted protein product of *Agrobacterium* fliR, which is located immediately after flhA, showed similar features to FliRs of several other bacteria (in addition to its strong homology to FliRs). FliR of *E. coli* is a 261 amino acid protein and is extremely hydrophobic, containing at least five membrane-spanning domains. The possible *Agrobacterium* FliR also contains 261 amino acids that are within the range of FliRs (251 aa *C. crescentus* and 269 aa in *R. sphaeroides*). The hydropathy
profile of the predicted FliR is similar to those of other FliRs. It is very hydrophobic throughout almost the entire polypeptide sequence. Hydropathy analysis revealed the Agrobacterium FliR to have at least 5 membrane-spanning domains. The similarities between A. tumefaciens and other homologues suggest that this polypeptide likely folds into a similar protein structure and serves similar functions. It has been suggested that the location of the N-terminal domain of the FliR protein is probably cytoplasmic. This protein is known as a class II flagellar gene residing near the top of the regulatory hierarchy that determines the order of flagellar gene transcription (122). It has been shown that FliR is a membrane protein, involved in early step of flagellar biogenesis and like FlhA, belongs to a family of proteins that constitute the flagellum-specific export pathway.

The predicted Agrobacterium FliR protein like FlhA also has homology with a family of proteins (type III secretion proteins) that are involved in transport of virulence factors (see table 3.6.1). This is again not surprising, because the members of both families are involved in the same activities of exporting the flagellar or virulence proteins that do not contain cleavable peptide signals. For example, SpaR (Spa29), which is essential for invasion of host cells and surface presentation of the virulence factor, or proteins like Ipa, harpin, Yops, InvE, and some flagellum that are exported by these two specific pathways do not contain the conserved N-terminal signal sequence. It seems that this superfamily of proteins is used for export or assembly, or both, of proteins that lie beyond the cell membrane.

Many components of the flagellum-specific export pathway have already been identified in A. tumefaciens in this laboratory (fliI, fliP, flhB and fliN), and are located in cosmid pDUB1900 (56). Now, two more possible members, flhA and fliR, of this system are being reported, located on a different fragment of the chromosome, which is located close to the cosmid pDUB1900. More components of the export pathway in A. tumefaciens are still required to be identified. fliO, fliH and fliQ have not yet been found in Agrobacterium. Finding these members would give a full perspective of function of the flagellum-specific export pathway and relation between the members.

Many bacteria have both protein delivery systems for pathogenic factors and flagella units, however this is not the case for Agrobacterium. It is that A. tumefaciens does not possesses a type III secretion protein system in despite finding such system in S. meliloti. Since the mechanism of pathogenicity that is used by Agrobacterium is different from those bacteria which possess type III secretion pathways involved in

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delivery of virulence proteins. *Agrobacterium* pathogenicity is brought about by insertion of a piece of plasmid DNA (T-DNA) instead of virulence factors. This piece of DNA contains the genes required for producing tumours and opines. Transformation of plasmid T-DNA from *Agrobacterium* into host cells, mediated by a mechanism that is referred to as typeIV system.

The protein product of the sequence that spans the *che*-2 mutant in *A. tumefaciens* showed similarity to CheL of *C. crescentus*. However the size of *A. tumefaciens* CheL is longer than that of *C. crescentus* (101). Unfortunately little is known about CheL and hence its precise function is unclear. It has been shown that *cheL* can complement a mutant possessing a functional flagellum, but unable to respond to chemotactic signals. The same phenotype was seen in *Agrobacterium che*-2 mutant. Since the characteristics of this mutant are similar to those of the *che*-2 mutant, it is possible to conclude that the two genes have similar functions, in addition to their homology in their structures. The middle part of the two proteins was found to have higher similarity; this may indicate the importance of the central region of the proteins for function. It has already been shown that at least the last 5 codon of *cheL* are not critical for function, as deletion of these does not affect the ability of the gene to complement a *cheL* mutant (101). The size difference between the two *cheLs* may reflect the evolutionary dirvegence between two bacteria. Another possibility is that this *orf* in *A. tumefaciens* might not be a *cheL* gene, instead is a unique gene, specific to *A. tumefaciens* with *cheL* nature and structure.

Unlike the flagellar and chemotaxis gene homologues, it is impossible to infer functional roles for the gene products of *orfX*, *orfY*, *orfW* and *orfZ* as a result of homologies to previously identified proteins. This may be because the gene homologues to these *orfs* have not yet been identified in any other species. Two open reading frames encoding proteins with unknown functions and no homologies to previously identified proteins have also been reported within a flagellar operon of *B. subtilis* (36). Such an open reading frame, *orfB*, has also found in a cluster of flagellar genes in *A. tumefaciens* (57). There is a possibility that these *orfs* encode proteins involved in functions or structural roles specific for *A. tumefaciens*, which are not found in other bacteria. However, their position within a region that contains two components of the flagellum-specific export pathway suggests that they may also be involved in this pathway. Considering the characteristics of *orfs*, *X*, *Y* and *W*, it is tempting to postulate that they may have chaperone functions (see section 1.4.3). All
of three ORFs share some of chaperone properties. They have molecular weight in the range of typeIII flagellum export proteins and typeIII secretion system (15-20 kDa), acidic isoelectric point (pI, see table 3.4.1). Since the activity of chaperones is specific to one or a few substrates, they do not usually show sequence similarity to other sequences. This could put more weight on speculation of chaperone-like activity of three ORFs, as they did not show to have homology to any other genes in database. On the other hand, they surround the only chemotactic gene in this region, as can also be seen in C. crescentus where other flagellar genes surround invA gene. Similarities in size, molecular mass, isoelectric point and guanine and cytosine content of all the orfs, except orfZ, with cheL should be considered as another possibility that they have a cheL-like nature and are possibly chemotactic genes. It is interesting to note that CheL has also the same chaperone properties as the ORFs (see table 3.4.1). This suggests that CheL might have chaperone-like activity as a flagellar or chemotactic protein too. This is unlikely, since evidence suggest that cheL is more likely to be a chemotactic gene rather a flagellar gene, and no chaperone has yet been reported for a chemotaxis system. Chaperones are specific to the delivery of proteins from inside the cell, across the membrane to the surface or beyond the cell, while the chemotaxis system transfer the extracellular signals from outside to inside the cell through a series of protein-protein interactions. It is therefore, unlikely to find a chaperone for chemotaxis system. More experiments should be carried out to determine the role of the orfs and cheL identified in this work. In the first place, mutation analysis experiments should be conducted, followed by phenotypic observation to find out whether this has any effect on motility or chemotaxis. Protein expression studies are also required to show that the gene products of the orfs expressed. Any information obtained from the overexpression of the orf gene products could also reveal the roles of these putative proteins. If the proteins are expressed, then immunocytochemistry could be carried out to localise their position in the cell and, thus, helping to find their possible functions.

The results obtained in this part of the project identified more behavioural genes in A. tumefaciens that are located on fragment of chromosomal DNA which is different from the fragments containing other motility and chemotaxis genes previously found in this laboratory. Since no flagellar or chemotaxis gene was found downstream of cheL and the fragments that contain the other behavioural genes (pDUB1900, pDUB1911 and pDUB1905) do not overlap each other, it may be useful.
to work on the proximity of these by finding cosmids from an *A. tumefaciens* library that overlap them. Resulting information can then explain the relation between the genes, whether they are clustered in one large fragment or there are intergenic gaps between the clusters. It would also give the location of the cosmids on the chromosome and would determine the exact distance between their *BamHI* sites.
4. Mutagenesis
4. Mutagenesis of fliR:

In an attempt to determine the effect on motility of one of the flagellum-specific export proteins of A. tumefaciens, strain C58C\(^1\), fliR was mutated by the gene replacement mutagenesis method using a neomycin cassette. The gene in question was knocked out by insertion of a neomycin cassette within the gene. The effect of the disrupted gene on motility was analysed phenotypically under light and electron microscopy and by swarm plate analysis. The gene replacement mutagenesis method was performed according to the protocol of J. Quandt and M. F. Hynes (162). The experimental procedure and basic theory are explained in section 2.10. A diagram of the procedure used to create the fliR mutation is shown in figure 4.1.1 This experiment involved two main phases. The first phase was to create a plasmid containing the gene disrupted by insertion of the neomycin cassette. The second phase involved introduction of the plasmid into Agrobacterium.

4.1 Creation of the fliR mutant plasmid:

Information obtained from sequencing showed that fliR lies between two XhoI sites of the plasmid ph1.11. The first step therefore was isolation of fliR from ph1.11. This plasmid was digested with XhoI and a 0.8 kb fragment was cut out. The 0.8 kb fragment contains the complete fliR sequence plus a few bases of SK\(^+\). The next step was to clone the fliR fragment into the suicide vector pJQ200SK (called pJQS for simplicity). The vector was cut with XhoI, purified and the 0.8 kb fragment (ph1.11/XhoI) was then ligated into the vector, forming the plasmid pJQfliR. Positive colonies containing the fliR fragment were selected based on inactivation of the β-galactosidase gene present within pJQSK, and confirmed by subsequent restriction enzyme digests of the possible correct colonies. Plasmid pJQfliR contains a unique NdeI site, located approximately in the middle of the second half of the fliR fragment (see figure 4.1.1). The plasmid was digested with NdeI and blunted.

The third step was disruption of fliR gene by inserting the neomycin resistance cassette within it. The 1.2 kb cassette was obtained from the plasmid pDUB2033 (see table 2.2.3.2) by digesting with EcoRI. The cassette was blunt ended and then
Figure 4.1.1 Diagram outlining the main sub-cloning steps involved in the construction of the $fliR$ mutant by insertion of neomycin cassette. (see text for details)

The empty box represents $fliR$, while the light shaded box indicates Neomycin cassette.
subcloned into blunt ended \textit{NdeI} cut, pJQfliR. Since pJQSK contains a gentamycin resistance gene, correct colonies were selected for by growth on LM-agar plates containing both gentamycin and Neomycin, and confirmed by subsequent restriction enzyme digests. The plasmid was named pJQfliR/Neo.

\textbf{4.2 Construction of an \textit{A. tumefaciens} fliR mutant strain:}

The second phase of the mutagenesis was to conjugate the plasmid pJQfliR/Neo into \textit{A. tumefaciens} C58C\textsuperscript{I} by tri-parental mating (see section 2.9). Nine colonies, which grew on neomycin, rifampicin and sucrose, were selected and subcultured in RifNeo5%sucrose plates.

Two Southern blots (see section 2.7) were carried out to find out whether the nine possible mutant colonies contained the neomycin cassette. Genomic DNA of the possible mutants was extracted and digested with \textit{HindIII}. Wild-type \textit{Agrobacterium} C58C\textsuperscript{I} was also cut with the same enzyme. The neomycin cassette and the \textit{fliR} fragment were radiolabelled and used as probes. The neomycin probe was expected to hybridise only with the mutated DNA, while the \textit{fliR} probe was expected to hybridise to both wild type and mutant DNA, but to different sized bands. The Southern blot hybridisation results can be seen in figure 4.2.1. When the blot was hybridised with the neomycin probe (see figure 4.2.1a), there was no band for wild-type C58C\textsuperscript{I}, and two bands, of the expected size for the two mutant colony lanes. The \textit{fliR} probe (see figure 4.2.1.b) also gave the predicted bands, and hybridised to both wild type and mutant DNA (see figure 4.2.2).

One mutant in the lane 3 appeared to have an extra band with the same size of wild type meaning that the mutant probably is not pure. The mutant phenotype was analysed in swarm plates and the result showed that the swarming pattern of mutant is as the wild type (5 cm after three days) indicating that this colony is a mixture of wild type and mutant. This mutant was discarded.

Because the Southern blot gave the expected results, one of the remaining colonies was selected for further analysis and renamed \textit{fliRNeo}. 

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Figure 4.2.1 Agarose gel of the genomic DNA cut with $Hind$III and subsequent Southern blot of nine putative $fliR$ Agrobacteria mutant strain hybridised with $fliR$ fragment (a) and neomycin cassette (b).

(a) Lane 1 is the wild type $A. \text{tumefaciens}$ (C58C1) genomic DNA digested with $Hind$III; lane 2 is $fliR$ fragment used as control; and lane 3 to 11 are mutants genomic DNA cut with the same restriction enzyme.

(b) Lane 1 is the wild type $A. \text{tumefaciens}$ (C58C1) genomic DNA cut with $Hind$III; lane 2 is neomycin cassette fragment used as a control; and lanes 3 to 11 are mutants genomic DNA cut with the same restriction enzyme.

$\lambda$ represents $\lambda$-$Hind$III marker.
Figure 4.2.2 Diagram showing the position and approximate size of the hybridising fragments of $fliR$ and neomycin cassette to C58C$^1$ and $fliR$ mutant strain.

a): Hybridising $fliR$ probe with C58C$^1$ wild type produces a 4.1kb fragment.
b): Hybridising $fliR$ or neomycin cassette to $fliR$ mutant produces two bands of 2.8 and 2.5 kb due to HindIII site in neomycin cassette.
Figure 4.3.2 Photograph showing the swarming behaviour of the flir mutant (right) and C58C1 (left) as a control.

Figure 4.3.3 Electron micrograph of the flir mutant strain.
4.3 Phenotypic analysis of *flir*Neo mutant:

The phenotype of the *flir* mutant was analysed under light and electron microscopy, and by swarm plates. The mutant strain (*flir*Neo) and wild type C58C<sup>1</sup> were each inoculated into the centre of 0.2% LB swarm plates. The plates were incubated at 28 °C typically for a period of 2-3 days. The swarms formed by the mutant strain were very small, compared to those of the wild type strain, (figure 4.3.2) indicating a possible defect in the motility machinery.

Further phenotypic analysis, using microscopic observation, was used to find out whether the difference seen in the swarming size of the mutant was due to a change in the construction or action of the flagellar machinery of the cells, compared to wild type *A. tumefaciens*. Visualisation by light microscopy showed that the *flir* mutant was non-motile, confirming the result obtained from swarm analysis. Further analysis of the mutant cells under the electron microscope showed that the cells were non-flagellated (see figure 4.3.3), therefore explaining the non-motile phenotype seen in previous tests.

4.4 Discussion:

Previous work producing mutations by insertion of the transposon Tn5 into chromosomal DNA of *A. tumefaciens* then created two non-motile mutants, *fla*-8 and *mot*-6 (180). The two mutants share an identical Tn5 insertion site. The *flir* mutant produced in this work showed the same phenotype as the non-motile mutants *fla*-8 and *mot*-6. This was not surprising as the two methods of mutation, disrupted a region of chromosomal DNA which spans *flir*, a member of the flagellum-specific export pathway. A similar phenotype has been seen in other bacteria, such as *C. crescentus* (238) and *R. sphaeroides* (75) when *flir* was mutated. It has been shown that defects in any of the flagellum-specific export genes causes delivery of flagellar proteins to be blocked and assembly of flagella to be affected as a result. For instance, mutations in *flhA* affect the translocation of a number of the *E. coli* flagellar proteins (219). Mutation in *flhA* created by insertion of aphA-3 cassette, produced a non-motile and non-flagellated phenotype in *Y. enterocolitica* (4). A similar phenotype was seen in *flhA* mutation in *B. subtilis* (36). *flIQ* mutant in *C. crescentus* produced a non-motile, non-flagellated phenotype (238). A site-specific mutant of *flil*, which was performed in this laboratory, resulted in non-motile phenotype with no visible flagella in *A. tumefaciens*. Interestingly, it has been observed that mutation on some components of
flagellum export apparatus, not only disrupt flagellar biogenesis but also normal cell
division as is seen in mutants of \textit{fliR} and \textit{fliQ} in \textit{C. crescentus} (238), and \textit{flhA} in
\textit{Proteus mirabilis} (80). These observation leads to suggestion that some classII
flagellum-specific export proteins apparatus genes may share, at least partially, their
regulatory pathway with other cell cycle events.

The phenotype seen in the \textit{fliRNeo} mutant was similar to that of \textit{fla-8} but not
\textit{mot-6}, in terms of having flagella (see figure 4.4.1). Considering the fact that the
phenotype of the most \textit{fliR} mutant is non-flagellated, and the two mutants share the
same Tn5 insertion site, raises a question about the source of variation between the
two mutants. One explanation for this at this stage of experiment, is that because the
transposon Tn5 insertion site was toward one end of the \textit{fliR} gene in \textit{A. tumefaciens},
sufficient transcription and translation may occur to produce a partially functional
FliR protein. This could allow exportation of flagellar proteins to proceed and
fragile

\textit{fla-8} \hspace{1cm} \textit{mot-6}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure.png}
\caption{Figure 4.4.1 showing \textit{fla-8}, which is unflagellated and \textit{mot-6} mutant, which is
flagellated}
\end{figure}

flagella to be produced under certain conditions. In this case both mutants should
have flagella, however it is possible that the fragile flagella present in the \textit{fla-8} mutant
were lost in the media while being prepared for observation under the electron
microscope, and the flagella in the *mot-6* mutant may remain untouched due to different conditions. In other words, the two mutants could in fact be only one mutant. This hypothesis could be true if Tn5 transposon was inserted toward the end of the gene, however Tn5 was found to locate at the beginning of the *fliR*. In this case the gene is not transcribed to produce FliR protein necessary for flagellar synthesis. It is possible that during the experiment and mutant process some uncontrolled error was made and *mot-6*, a wild-type bacteria with a normal flagella showed mutant phenotype and assigned as a mutant.
5. Conclusion
5. Conclusion:

The evidence obtained so far, suggests that *A. tumefaciens* like other bacteria, possess a flagellum-specific export apparatus which is used to deliver the extracellular protein units of the flagella that do not contain a cleavable leader signal peptide. Some components of this pathway were identified in the past, from work carried out in this laboratory (see section 1.4.3). Two more potential components of the flagellum-specific export system were found in this work. At present, it appears that *A. tumefaciens* contains six of the ten common components of the system, defined by Minamino *et al* (138). However, three other common components, *fliO, fliQ, fliH* and other members of the pathway, such as chaperones, which may be specific to *A. tumefaciens* or the α-subgroup of proteobacteria, have not yet been identified.

The protein products of two genes (*flhA and fliR*) show similarity to both the flagellum-specific export pathway and typeIII proteins. The homology between *Agrobacterium* FlhA and FliR, and their counterparts in the flagellum-specific export system in other bacteria was higher than similarity to typeIII proteins. This level of similarity may count as evidence indicating that the two genes are flagellar-related genes, rather than typeIII, which it would be unlikely to find in *Agrobacterium*. The given data cannot state whether the two genes belong to an operon upstream of *flhA* or whether they form an operon with genes found downstream of *fliR*. Further analysis of the region immediately upstream of *flhA* is required to determine whether there are more flagellar-related genes, a promoter or a terminator in this region. If any promoter-like sequence was found, transcriptional fusion experiments using *lacZ*, could be carried out to analyse the promoter function. In light of what is known, it would be possible to determine the boundary of the putative operon, as well as any other putative transcription signals within it, and find the relationship between them and the rest of the flagellum-specific export genes. Northern blot analysis could also be used to identify the organisation of the genes in the operon.

In order to study the function of *flhA* in *A. tumefaciens* in more detail, mutation analysis should be made on the *flhA*. This would reveal the effect of disrupting *flhA* in flagella biogenesis and then motility. It has been shown that inactivation of the *flhA* gene leads to non-motile phenotype in other bacteria. FlhA in addition to its protein export activity has a regulatory function too. *flhA* in *C. crescentus* is required for
expression of several flagellar gene, including flaN, flbG, flgL, and flgK. In a non-motile mutant of Proteus lacking the flhA gene, expression of class II and III flagella genes is suppressed. Surprisingly, the flhA mutant did not elongate beyond vegetative cell length (80) which has been seen in the fliR mutant (see section 4.3).

Mutation in A. tumefaciens fliR gene result in a non-flagellated cell as has been observed in many other bacteria. An attempt to complement the flhA and fliR mutants by other bacterial counterpart genes would show whether the proteins have functional homology as well as sequence similarities. It has been known that fliR is activated early in the cell cycle (238). Vegetative growth of both mutants could be measured under electron microscopy to find the effect of mutation in cell division in Agrobacterium.

At present we have no idea as to the location and function of the gene products (FlhA and FliR) in Agrobacterium. It has been suggested that in S. typhimurium the membrane-associated components of flagellum-specific export system (FlhB, FlhA, FliO, FliQ and FliR) reside in the central pore within the MS-ring of the flagellar basal body (137). Whether this suggestion is correct for Agrobacterium needs to be investigated. To find the location of FlhA and FliR, immunoblotting analysis (antibodies/western blotting) of purified inner and outer membrane fractions could be used. In this experiment the FlhA and FliR protein would be fused to a fragment of β-lactamase, which lacks its signal peptide for membrane translocation.

Besides the two components of the flagellum-specific export genes, a chemotaxis gene, cheL, along with four unknown orfs were also found in this work. The function of cheL remains to be determined, however mutation in the region containing cheL, resulted in a flagellated strain with a wild type motility phenotype (180). A cheL mutant in C. crescentus does not respond to chemotactic substrates, indicating a defect in the chemotaxis system (101). The same result was obtained in A. tumefaciens for the che-2 mutant (180) which was mutated in a region spanning cheL. However, methylation is reduced in the A. tumefaciens che-2 mutant (178), whereas the methylation in this type of mutation would be increased. It has been shown that mutations in other che genes in A. tumefaciens produces impaired chemotactic capabilities with a wild type pattern of motility (228). This leads to the suggestion that there may be an additional level of control of the flagellar motor, independent to that from the chemotaxis system. The function of protein product of cheL in the chemosensory system is not yet known. Mutation analysis, followed by chemotaxis assays using specific chemotractants, will
show the effect of cheL in the chemosensory network. The effect of this gene on motility of A. tumefaciens could be studied using a computer-linked tracking system that analyses statistically the motion of the bacteria, comparing it with wild type motion.

The four other orfs identified did not have homology to any previously known flagella or chemotaxis genes. It may be because the gene homologues to the orfs have not yet been identified in other bacteria, or because they are specific to Agrobacterium. It has been shown that bacteria belonging to α-subgroup of proteobacteria are equipped with a number of unique genes (57) along with known gene homologues that are different from the enterobacteria. Mutation in the orfs could be used, in the first place, to study the effect of the orfs in motility or chemotaxis. Primer extension experiments could be carried out to show whether they are transcribed together, along with the other genes found in this fragment. Protein expression studies would be then required to show that the protein products of the orfs are expressed. Any effects on motility from mutagenesis or over expression may also help to determine the roles of the putative proteins. If any of the proteins were expressed, the immunobloting could be carried out to localise their position in the cell, thus giving a possible insight into its function.

There is not enough information to explain how the flagellar export gene homologues, cheL and the four orfs are transcribed and regulated. A good transcription termination sequence was found downstream of orfW, it is therefore plausible to suggest that flhA, fliR, cheL and orfXYW, but not orfZ, form an operon or are part of an operon containing more genes upstream of flhA. Further experimental work will be necessary to resolve whether this speculation is correct.

The three cosmids, pDUB1900, pDUB1911 and pDUB1905 contain several flagellar and chemotaxis genes (57, 228). An experiment was carried out using hybridisation of fragments of these cosmids to cosmids from S. meliloti containing similar genes, (pRZ1, pRZ2 and pRZ4), which are known to overlap each other (55). The work showed that the three Agrobacterium cosmids to be close, however they do not overlap. It may be useful to identify a cosmid from an A. tumefaciens library that overlaps the cosmids pDUB1900, pDUB1911 and pDUB1905. Pulsed-field gel electrophoresis (PFGE), in combination with Southern hybridisation, could be used as a powerful tool to unify the sequences obtained from the three cosmids in A. tumefaciens. This would explain the relationship between the genes, whether they are clustered into one large fragment, or whether there are intergenic gaps between the clusters. It would
also give the location of the cosmids on the chromosome and would determine the exact distance between their BamHI sites.

The results obtained in this work provide a lot of scope for further work, particularly on the flagellum-specific export system, which is an important system without which, no flagella can be built. It is thought that this transport protein machinery has general components that are common in all bacteria, and members that are specific for each organism. It is known that \textit{A. tumefaciens} contains most of the general components, however the specific members should be identified. One possibility is that the ORFs found in this work could be the specific members of the typeIII flagellum-specific export system. The location, regulation and function of various components of the system also need to be investigated. The flagellar proteins that each of the components deliver could be a subject of research. The way that the system works and the interaction between the components to transport proteins also need to be found. Although, in recent years, much is known about function of this system in \textit{S. typhimurium}, however the flagellum export system in \textit{A. tumefaciens} could work in different way, since the structure and organisation of flagellar proteins of the two organisms are different.

Investigation of the flagellum-specific transport components could be extended to find out if the homologue components are functionally conserved among other bacteria.

Homologies between flagellum export system and typeIII secretion system suggests that two systems could be related functionally too. \textit{A. tumefaciens} is about only Gram-negative bacteria that does not possess the typeIII secretion system. Thus, it is interesting to investigate the interaction between flagellum export system and typeIII secretion system in the sense that proteins (effectors in typeIII, and flagellar proteins in flagellum export systems) are translocated by one system could be exported or even secreted intracellularly by another system. It has been shown that the corresponding components of the typeIII system in different bacteria are not interchangeable. However, the existence of interaction between different typeIII export/secretion systems has been demonstrated in many experiments. In one occasion (52) the \textit{pcrV} gene from \textit{P. aeruginosa} show to compete an \textit{lcrV} mutation in \textit{Y. pseudotuberculosis}, and ExoS protein from \textit{P. aeruginosa} and IpaB from \textit{S. flexneri} could be secreted by \textit{Y. pseudotuberculosis} typeIII system. \textit{S. typhimurium} can also translocate YopE (52). It has been found that the \textit{Y. enterocolitica} unable to synthesise flagellum could not also
produce several secreted proteins. In another experiment, mopE, a typeIII secretion system component, mutant did not produce flagella (142). A more exciting experiment was that which showed the YpIA, a virulence-associated phospholipase, which is required for Yersinia pathogenesis was exported by the flagellar export apparatus (232). This suggests that flagellum export machinery may have a role in the secretion of proteins involved in bacterial-host interaction. Therefore, it would be useful to determine the functional conservation between corresponding proteins of the flagellum-specific export system and the typeIII pathway. This would enable a more accurate look into the export machinery containing a superfamily of proteins that are used in a variety of ways to control and enable the process of export or assembly, or both, of proteins that lie beyond the cell membrane.

The interaction between the flagellum-specific export pathway and pathogenicity in Agrobacterium should also be considered. The motile mutant could be inoculated directly into test plants in order to assess their tumourgenicity. However, this method might not show the requiremnt of motality, as perior to attachmant for the bacteria to move toward their target. It has been shown that mutation in any of the flagellum export components leads to a non-motile phenotype (75, 127, 136, 137). The role of motility in virulence has been investigated in different bacteria. It has been reported that motility mutant, non-flagellated bacteria, are affected in their virulence because of aberrations in their host adherence, invasion mechanism or other unknown factors (42). The effect of motility in Agrobacterium pathogenicity is not clear. Results obtained from studies on fla- mutants in A. tumefaciens, including mutations in some of the flagellum-specific export genes, showed that there were no differences between the parent and mutant in the ability to bind suspension cells and cause tumours (180). However, Chesnokova, O. et al. (42), have reported that virulence in non-motile, bald strains, which were generated by deletion of flaABC, was significantly (in statistical terms) below that of the parental wild type. Some results suggest that in bacteria that contain both systems of protein export, (The flagellum-specific export pathway and the typeIII export system) pathogenicity could be affected by a motility disruption. There is evidence showing that mutation in a typeIII protein, MopE (a FliR homologue), in E. carotovora, affected assembly of the flagella and lead to a non-motile phenotype (142). These results may suggest that the virulence factor export pathway may also be used for export of flagellar components, and vice versa. It is possible that the two systems share a common point in their routes. However, in Agrobacterium a different mechanism of
pathogenicity, which involves transfer of T-DNA to host cells, is used, and a different pathway, typeIV, is responsible for transfer of the T-DNA. Therefore disruption of the flagellum-export proteins should not affect T-DNA transfer and then tumourigenicity. However much more work is required to be done to fully understand the biology, physiology and biochemistry of the flagellum-specific export system in A. tumefaciens.

It should now possible to study biogenesis of flagella, motility, and in particular, the role of flagellum-specific export system in more detail in A. tumefaciens. This should ultimately lead to a greater understanding of A. tumefaciens biology in general, increasing the knowledge of Agrobacterium as a genetic tool and a plant pathogen.
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