Genome size and macrorestriction map of *Xanthomonas campestris* pv. *glycines* YR32 chromosome

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Abstract

*Xanthomonas campestris* is an important plant pathogenic bacterium which causes severe diseases in a wide variety of plant species. We have generated a macrorestriction map of the *X. campestris* (axonopodis) pv. *glycines* chromosome employing pulsed-field gel electrophoresis (PFGE). Restriction endonucleases *PacI* (5'-TTAATTAAA), *PmeI* (5'-GTTTAAAC) and *SmaI* (5'-ATTAAAAAT) digested the chromosomal DNA into three, five, and five fragments, respectively. In addition, intron-encoded restriction endonuclease *I-CeuI* was employed to locate the position of the 23S rRNA genes (*rrlA* and *rrlB*). All of the generated restriction fragments were aligned along the chromosome using multiple restriction enzyme digestion and two-dimensional PFGE (2-D PFGE) in conjunction with Southern hybridization analysis. This physical map construction has revealed a single circular chromosome with a size of approximately 5 Mb. Two rRNA genes were localized on the chromosome map. Several genes involved in pathogenesis (*xpsD*, *opsX*, and *pat*) as well as genes involved in the biosynthesis of xanthan gum (*xanAB*, *rfhCDAB*) were also localized. © 1999 Federation of European Microbiological Societies. Published by Elsevier Science B.V. All rights reserved.

Keywords: *Xanthomonas*; Pulsed-field gel electrophoresis; Mapping; Circular chromosome

1. Introduction

*Xanthomonas campestris* (axonopodis) pv. *glycines* [28] is a plant pathogenic bacterium that causes bacterial pustules on soybean cultivars and is a member of the most diverse of the five species in the genus *Xanthomonas* [2]. *X. campestris* has become a model system for the study of pathogenesis because of its broad capacity for causing diseases in various species of plants. To date, our knowledge of the pathogenesis mechanisms of *X. campestris* pathovars is rather limited. It is believed that extracellular proteases [27], lipopolysaccharides [11], exopolysaccharides [17], and endoglucanases [7] directly or indirectly play a role in the pathogenesis process. However, the mechanisms of how these enzymes interact to cause disease in plants are poorly understood.

Despite numerous studies of genes involved in pathogenesis [5,7-11,16,17,27] and extracellular polysaccharide biosynthesis [12], little is known about the structure and organization of the *X. campestris*
genome. Therefore, we undertook the construction of a physical map of the *X. campestris* genome using pulsed-field gel electrophoresis (PFGE) [3,23]. We report here the chromosomal structure and genome size of *X. campestris* pv. *glycine* YR32. We have mapped several genes involved in pathogenesis, i.e. *upxX* [11], *pat* [10], *xpdA* [8], and xanthan gum biosynthesis, i.e. *xanAB, rfbCDAB* [12], and have determined the location of two rRNA genes in the chromosome. (Part of this work was presented at the 8th International Congress on Molecular Plant-Microbe Interactions, Knoxville, TN, USA, in 1996.)

2. Materials and methods

2.1. Bacterial strains and plasmid

Bacterial strains, plasmid and specific DNA fragments used in this study and their relevant characteristics are described in Table 1.

2.2. Growth conditions and media

Strains of *Escherichia coli* and *Rhodobacter sphaeroides* 2.4.1 were routinely grown in Luria-Bertani (LB) broth (yeast extract 5 g l⁻¹, tryptone 10 g l⁻¹, NaCl 10 g l⁻¹) pH 7.2 at 37°C [21], and *X. campestris* pv. *glycine* YR32 was routinely grown in LB+5% glucose pH 6.8 at 30°C. Plate cultures of *X. campestris* pv. *glycine* YR32 and the *E. coli* were grown on YDC (yeast extract 10 g l⁻¹, dextrose 5 g l⁻¹, CaCO₃ 20 g l⁻¹, Oxoid agar 15 g l⁻¹) [10] and LB with 15 g l⁻¹ Oxoid agar, respectively. Whenever appropriate *E. coli* cultures were supplemented with ampicillin 100 µg ml⁻¹, tetracycline 15 µg ml⁻¹, chloramphenicol 34 µg ml⁻¹, or kanamycin 50 µg ml⁻¹.

2.3. Preparations of intact genomic DNA and restriction digests

Bacterial suspensions were prepared as follows: one loopful of a single colony was inoculated into 10 ml LB pH 7.2 (*R. sphaeroides* 2.4.1) or 10 ml LB+5% glucose pH 6.8 (*X. campestris* pv. *glycine* YR32), and incubated at 30°C for 24 h and 48 h, respectively. Bacterial suspensions were harvested by centrifugation and resuspended in PIW buffer (10 ml Tris-HCl, pH 7.5; 1 M NaCl) to a final concentration of approximately 2×10⁹ cells ml⁻¹. Chloramphenicol (10 µg ml⁻¹) was added 15 min prior to harvest in order to synchronize chromosomal DNA replication.

The gel inserts (10×5×1 mm) were prepared as described by Smith and Cantor [24]. Restriction endonuclease digests were completed according to the protocol of Sawanto and Kaplan [25], with 8–15 U of enzyme for each digest, except for *I-CeuI*, for which we used 1 U of enzyme. Digestions with *PstI* or *PmeI* were performed at 4°C for 16 h, then at 37°C for 12 h, while digestions with *SsoI* were conducted at 4°C for 16 h, then at 25°C for 12 h (*SsoI*). Digestion with *I-CeuI* was performed at 37°C for 3 h. Multiple digestions were performed sequentially by washing the gel inserts with 1× TE buffer [21] for 1 h at 4°C, followed by incubation in the second or third restriction enzyme buffer for 15 min before subsequent digestion. All restriction buffers and conditions were carried out as recommended by the manufacturers (New England Biolabs or Boehringer Mannheim). *SsoI* was obtained from Boehringer-Mannheim Co. (Singapore), *AseI, SpeI, PstI, PmeI,* and *I-CeuI* were obtained from New England (Singapore). Low melting point agarose for gel inserts was obtained from Bio-Rad (P.T. Diastika Biotekindo, Jakarta).

2.4. DNA fragment separation and macrorestriction fragment analysis

CHEF DR-II (Bio-Rad, Richmond, CA) [3] was used to separate DNA fragments. For separations of *PstI, PmeI,* and *SsoI* fragments, electrophoresis was carried out at 3.6 V cm⁻¹ for 50 h with 50-650 s pulse times. For resolution of *I-CeuI* fragments, electrophoresis was conducted at 3.6 V cm⁻¹ for 45 h with 50-650 s ramping pulse times. Two different pulse times of 20–25 s and 25–70 s were employed to separate *AseI* and *SpeI* small to medium, and large fragments, respectively. PFGE was conducted at 5.4 V cm⁻¹ for 22–24 h. For all gels, we used 1% multi-purpose agarose (Boehringer-Mannheim) or ultra-pure agarose (Gibco BRL). *Saccharomyces cerevisiae* chromosomal DNA (New England Biolabs or Pharmacia) and *SpeI*-digested *R. sphaeroides* 2.4.1 genomic DNA [26] markers were used as molecular
size standards for the PucI, Pmel, Swal, and I-CeuI digests. For Asel and Spel digests, lambda concatemeric DNA (New England Biolabs) and Asel-digested R. sphaeroides 2.4.1 genomic DNA [26] were used as a molecular size standards.

2.5. Two-dimensional PFGE (2-D PFGE)

Initial separation of single digestions of X. campestris pv. glycines YR32 genomic DNA with either PucI, Pmel, or Swal was conducted as described in the DNA fragment separation, with the exception that 0.8% multi-purpose agarose (Boehringer-Mannheim) was used instead of 1% agarose. Each of the single digestions was performed in duplicate and run side by side in the same electrophoresis gel. Following separation, one lane of separated DNA fragments was cut out and stained with ethidium bromide (EtBr) for 15 min, and destained with sterilized water for another 30 min. The other lane consisting of the same DNA fragments was kept unstained. The DNA fragments in the stained gel were visualized using a UV transilluminator. The approximate location from the well of each DNA fragment was recorded. These data were then used to locate DNA fragments in the unstained gel. Each of the DNA fragments from the unstained gel was excised and washed by incubating in 1.5 ml 1×TE buffer for 1 h at 37°C. This treatment decreased the concentration of boric acid and EDTA in the gel, since the concentration of EDTA in 1×TE buffer is lower than that in the 0.5×TBE running buffer.

After three repeated washes, the gel slice was incubated with 150 µl restriction buffer for 1 h at 4°C. Incubation in the appropriate restriction buffer was also repeated three times. The restriction enzymes were added and digestions were performed as described previously [25]. The resulting DNA fragments were separated by PFGE using the conditions described in Section 2.4.

2.6. Southern hybridization analysis

Separated DNA fragments were transferred onto Photogene nylon membrane (Gibco BRL) using the capillary transfer method [21]. Because of the large size of many of the DNA fragments to be transferred, the DNA was first depurinated with 0.2 N HCl for 10 min, and washed in distilled water for 5 min prior to denaturation. Non-radioactive probes were prepared from specific DNA fragments derived from unique plasmids (Table 1) using the Bio-nick Translation kit (Gibco BRL). The probes were made by labelling each DNA fragment with biotin-14-dATP. Southern hybridization was performed at 42°C as described by Donohue et al. [6]. The washes were performed at 45°C, except for the RNA probe at 50°C. For detection, we used the chemiluminescent assay procedure described by the manufacturer (Photogene detection system, Gibco BRL). The X-ray film used in this study was the product of Agfa (Germany).

3. Results and discussion

3.1. Resolution of restriction fragments using rare cutting restriction endonucleases

To construct a physical map of the X. campestris pv. glycines genome, restriction endonucleases which cut its genomic DNA into a small number of fragments were sought. The mol% G+C content of the DNA was generally used to narrow the range of restriction endonuclease to be tested [26]. Based on information that X. campestris has 63–71 mol% G+C DNA [2], we looked for enzymes with A+T-rich recognition sequences. Among the enzymes that we have tested, we found that PucI (5'-TATAATTAA), Pmel (5'-GTGTTAACA), and Swal (5'-ATTTAATTAA) cut the DNA infrequently. PucI, Pmel, and Swal digested X. campestris pv. glycines YR32 genomic DNA into three, five, and five fragments, respectively, with manageable distributions of DNA fragments and no overlapping fragments (Fig. 1). Double digestion of genomic DNA with combinations of these enzymes yielded fewer than 10 restriction fragments. Each of the PucI-Swal, Pmel-Swal, and PucI-Pmel digests generated eight restriction fragments with one band shown to consist of two fragments. By using a low voltage electric field, a long separation time, and a wide range of pulse times, we were able to separate large DNA fragments of more than 2000 kb in size employing the PFGE apparatus. Digestion of genomic DNA with the above mentioned enzymes yielded numbers and distribution.
Fig. 1. Pulsed-field gel electrophoresis (PFGE) of X. campestris pv. glycines YR32 genomic DNA digested with Pael, Smel, and Pwel (50-650-s pulse; 58-h run). A: Pael, Smel and Pael-Smel digestion. Lanes: 1, yeast chromosomal standards; 2, X. campestris pv. glycines YR32 genomic DNA digested with Pael; 3, Pael-Smel digest; 4, Smel digest; 5, R. sphaeroides 2.4.1 genomic DNA digested with SpeI as molecular size markers. B: Pmel, Smel and Pmel-Smel digestion. Lanes: 1, yeast chromosomal standards; 2, Pmel digest; 3, Pmel-Smel digest; 4, Smel digest; 5, SpeI-digested R. sphaeroides 2.4.1 markers. C: Pael, Pmel and Pae-Pmel digestion. Lanes: 1, yeast chromosomal standards; 2, Pael digest; 3, Pael-Pael digest; 4, Pmel digest; 5, SpeI-digested R. sphaeroides 2.4.1 markers.

of DNA fragments suitable for physical mapping [27]. The DNA fragment sizes are summarized in Table 2. When undigested total genomic DNA was run concurrently with the digested DNA using the same condition, no extrachromosomal DNA was visible (data not shown). Therefore, all restriction fragments generated using either Pael, Pmel, or Smel were considered to be chromosomal DNA.

3.2. Genome size of X. campestris pv. glycines YR32

X. campestris pv. glycines YR32 genomic sizes were estimated from the summation of the individual fragment sizes represented by each restriction endonuclease digestion. Genomic size, from single or multiple digestion with Pael, Pmel, and Smel, was estimated at approximately 5020 kb (Table 2). This result was in agreement with the results from measurements of AseI and SpeI digests of genomic DNA (Fig. 2). Digestion of X. campestris pv. glycines YR32 genomic DNA with AseI and SpeI yielded approximately 37 and 47 restriction fragments, respectively, with several bands containing more than one DNA fragment. The genome size was estimated to be 4974 ± 123 using SpeI-digested genomic

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firmed this conclusion. The probe hybridized strongly to both *PmeI*-A (Fig. 4, lane 3), *SspI*-A (Fig. 4, lane 4), and the 1400-kb subfragment seen in *PmeI*-SspI digests of genomic DNA (Fig. 4, lane 6). The *PcoI* digest of the *SspI*-A fragment also contained two 275-kb subfragments (Fig. 4, lane 5), the order of which was unknown. Since the sequence of the *SspI*-A, -B, and -C fragments has been determined, the positions of *PmeI*-A, -C, and -B fragments relative to those three *SspI* fragments could be determined. The *PmeI*-A fragment (Fig. 4, lane 2) must overlap the *SspI*-B fragment (Fig. 4, lane 9) since both these fragments contained the same 53-kb subfragment. This ordering was in agreement with the result that the *PmeI*-A fragment was linked to the *PmeI*-C fragment, which contained a 1035-kb subfragment of the *SspI*-B fragment. These data indicated that the *SspI*-C fragment should be located upstream of *PmeI*-A, -C fragments, and therefore the fragment must overlap the *PmeI*-B fragment. A 90-kb subfragment was derived from the *SspI* digest of *PmeI*-C fragment (Fig. 4, lane 4). Since *PmeI* did not cleave the 90-kb *SspI*-D fragment, the data suggested that the *SspI*-D fragment was located within the *PmeI*-C fragment (Fig. 4, lane 11). From the previous data, we were able to determine the order of the *PmeI* restriction fragments to be -B-D-A-C-E, which overlaps the ordering of the *SspI* restriction fragments -C-A-B-D-E (Fig. 6).

3.4. Ordering the *PcoI* fragments

*PcoI* digested *X. campestris* pv. *glycines* YR32 genomic DNA genomic DNA into three restriction fragments (Fig. 1). The order of the *PcoI*-A, -B, and -C fragments relative to those *PmeI* fragments was determined using multiple digestion of *PcoI-*

*PmeI* digests of genomic DNA, 2-D PFGE analysis and Southern hybridization analysis, as previously described for the *PmeI* and *SspI* fragments. We also determined the order of the *PcoI* fragments relative to those of *SspI*. Any ambiguities that appeared in 2-D PFGE analysis were resolved by Southern hybridization analysis using selectable genetic markers (Table 1) and by multiple digestion of *X. campestris* pv. *glycines* YR32 genomic DNA.

Physical mapping of the *X. campestris* pv. *glycines* YR32 chromosome reveals a genomic structure that consists of a single circular chromosome. There was no evidence of any extrachromosomal DNA larger than 10.5 kb in this strain.
3.5. Constructing a partial genetic map

Southern hybridization experiments were used to localize previously described genes of *X. campesiris* (Table 1). The *pat* (Fig. 4) and *opsX* probes hybridized strongly to the *PacI-A, Pmel-A, and SvaI-A* fragments. It also hybridized to the 1650-kb subfragment of *PacI-PmelI*, the 1400-kb subfragment of *Pmel-SvaI*, and the 1950-kb subfragment of *PacI-SvaI* digests of genomic DNA. The *xspD, xanAB*, and *rbcCDAB* probes (data not shown) hybridized strongly to the *PacI-B, Pmel-C*, and *SvaI-B* fragments. The probes also hybridized to the 1125-kb subfragment of *PacI-PmelI*, the 1035-kb subfragment of *Pmel-SvaI*, and the 1315-kb subfragment of *PacI-SvaI* digests of genomic DNA. Further Southern hybridization analysis with *I-CeuI* single and double digests of genomic DNA suggested that these genes were located within the larger *I-CeuI* fragment, the 1565-kb subfragment of *I-CeuI-PmelI*, the 800-kb subfragment of *I-CeuI-PmelI*, and the 710-kb subfragment of *I-CeuI-SvaI* digests of genomic DNA (Fig. 6).

All genetic markers used in this work are thought to be involved in plant pathogenesis (Table 1). The *pat* gene, a DNA fragment conserved in several plant and animal pathogens [9], was located on the same fragment as *opsX*, a gene which is involved in lipopolysaccharide biosynthesis and encodes a virulence factor [11]. *xspD*, whose gene product is required for the secretion of extracellular enzymes across the outer membrane of *X. campesiris* [8], was located on the same fragment as *rbcCDAB* and *xanAB* genes (see Fig. 6). *rbcCDAB* is a cluster of genes involved in the synthesis of the undecaprenyl phosphate unit necessary for lipopolysaccharide production, while the *xanAB* gene encodes proteins involved in synthesis of the UDP-glucose and GDP-mannose units required for xanthan gum production [12]. Localization of other pathogenesis genes will be required in order to explain any possible correlation between genome structure and the distribution of pathogenesis genes on the chromosome.

![Fig. 5. PFGE of *X. campesiris* pv. *glycines* YR32 digested with *I-CeuI* and subsequent digestion with *PacI, PmelI*, and *SvaI*. Lanes: 1, *R. solani* 2412 genome DNA digested with *PsmI* as molecular size markers; 2, *I-CeuI* digest; 3, *I-CeuI-PmelI* digest; 4, *I-CeuI-PmelI* digest; 5, *I-CeuI-SvaI* digest; 6, *I-CeuI-PmelI* digest; 7, *I-CeuI-PmelI* digest; 8, *I-CeuI-PmelI* digest; 9, yeast chromosomal standards.](image)

![Fig. 6. Physical map of *X. campesiris* pv. *glycines* YR32 chromosome. *PacI, PmelI, SvaI*, and *I-CeuI* restriction sites are indicated with designated fragments (see Table 2). Estimated sizes are indicated in the innermost circle. Description of genes is listed in Table 1.](image)

3.6. Two rRNA genes localization employing intron-encoded *I-CeuI* restriction endonuclease

To determine the number of rRNA genes and their locations on the chromosome, we utilized an intron-encoded restriction endonuclease *I-CeuI*, which recognizes a highly conserved 19-bp sequence in *rrl* genes for the large rRNA subunit (23S) [15]. Digestion with *I-CeuI* and other enzymes determined the precise location of the 23S rRNA genes on the chromosome (Fig. 5). *I-CeuI* cleaved *X. campesiris* pv. *glycines* YR32 genomic DNA into two fragments (Fig. 5, lane 2) with the smaller fragment estimated at approximately 430 kb. Based on previous results
[14,15], the data from this experiment indicated that X. campestris pv. glycines YR32 has two 23S rRNA genes, and they are 430 kb away from each other. From the estimated genome size, we therefore calculated the larger fragment to be 4590 kb in size.

The Pco1-B fragment was cleaved by I-CeuI into three subfragments, whose sizes were approximately 1565, 430, and 173 kb (Fig. 5, lane 3). The Pco1-A and -C fragments appeared undigested in I-CeuI-Pco1-digested genomic DNA, suggesting that these two fragments did not contain 23S rRNA genes. The data indicate that the two 23S rRNA genes are located within the Pco1-B fragment. I-CeuI did not cleave the Pco1-B, -D, and -E fragments (Fig. 5, lane 4), suggesting that these three Pco1 fragments also did not contain I-CeuI-cut sites. I-CeuI cleaved the Pco1-A fragment into 1825- and 105-kb subfragments, and the Pco1-C fragment into 800- and 325-kb subfragments. The summation of the 105-kb subfragment of Pco1-A and the 325-kb subfragment of Pco1-C, which totals approximately 430 kb, clearly indicated that one 23S rRNA gene was located within the Pco1-A fragment, and the other was within the Pco1-C fragment.

Digestion of Sva1 fragments with I-CeuI yielded seven fragments (Fig. 5, lane 5). I-CeuI did not cleave the Sva1-A, -B, -D, and -E fragments, indicating that these four fragments do not have any I-CeuI sites. The Sva1-C fragment was cleaved by I-CeuI into 715-, 430-, and 425-kb subfragments, with the latter two appearing as a single intensely stained band. These data showed that the two rRNA genes were located within the Sva1-C fragment. Southern hybridization analysis using a fragment derived from the P. aeruginosa rRNA operon as a probe has confirmed these results (data not shown). This probe hybridized to Pco1-A and -C fragments, to the 1035- and 530-kb subfragments of the Pco1-Sva1 digests, and to the 1315-kb subfragment of Pco1-Sva1 digests of genomic DNA.

Multiple digestions of I-CeuI genomic DNA fragments with two different enzymes, i.e. I-CeuI-Pco1-Pco1 (Fig. 5, lane 6), Pco1-Pco1-Sva1 (Fig. 5, lane 7), and I-CeuI-Pco1-Sva1 digests (Fig. 5, lane 8), also confirmed the positions of the 23S rRNA genes. This analysis clearly supports the positions of the 23S rRNA genes and verifies the total genome size of X. campestris pv. glycines YR32.

The finding that X. campestris pv. glycines (and perhaps all X. campestris pathovars) have only two 23S rRNA genes deserves special attention. Bacteria can contain up to 10 repeated rrr operons within their genomes [4,13]. Data correlating bacterial genome size suggest that there is no significant relationship between the number of rrr operons and genome size. The fast-growing E. coli, with a genome slightly smaller than X. campestris, has seven rrr operons [1], and Bacillus subtilis has a genome of approximately 4.1 Mb has 10 rrr operons [14], while P. aeruginosa with a genome relatively larger than X. campestris has four rrr operons [18].

This chromosome map will serve as a framework in our efforts to refine the physical map of X. campestris pv. glycines YR32, to localize transposon insertion sites and genes involved in pathogenesis, and to study the dynamics or the genome plasticity of this agriculturally important bacterium.

Acknowledgments

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References


