九州大学学術情報リポジトリ Kyushu University Institutional Repository

Pathogenic and Genetic Diversity in Asian Strains of Xanthomonas oryzae pv. Oryzae

Jyufuku, Shinyou

Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University

Furuya, Naruto

Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University

Goto, Takahiro

Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University

Tsuchiya, Kenichi Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University

他

https://doi.org/10.5109/14031

出版情報:九州大学大学院農学研究院紀要. 54 (1), pp.19-23, 2009-02-27. Faculty of Agriculture, Kyushu University

バージョン:

権利関係:



Pathogenic and Genetic Diversity in Asian Strains of Xanthomonas oryzae pv. oryzae

Shinyou JYUFUKU¹, Naruto FURUYA*, Takahiro GOTO¹, Kenichi TSUCHIYA and Atsushi YOSHIMURA²

Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science,
Department of Applied Genetics and Pest Management, Faculty of Agriculture,
Kyushu University, Fukuoka 812–8581, Japan
(Received November 11, 2008 and accepted December 5, 2008)

The 57 strains of $Xanthomonas\ oryzae\ pv.\ oryzae\ (Xoo)\ collected$ from rice growing countries of India, Indonesia, Malaysia, Thai, Taiwan and Philippines were characterized by using polymerase chain reaction fingerprinting and virulence analysis. The strains of Xoo were grouped into 13 races on the basis of their pathogenicity to international differential lines. Distribution of races was quite specific to the country. Three strains from India and two strains from Indonesia were virulent to cultivars containing the bacterial blight resistance gene xa5, while most strains from other countries were avirulent to xa5. The strains from India showed high virulence and broad range of pathogenicity, in contrast to those from Malaysia. Two varieties of containing Xa21 and xa5 gene, respectively, were resistant to almost Asian Xoo strains. Our Xoo collections from Asian countries were also divided into 4 genetic groups by clustering statistics on the basis of the results from PCR-based RFLP with IS1112 primers. A partial relationship was found among the genetic groups, countries and races, suggesting that strategies that target regional resistance breeding and gene deployment are feasible. The results of this study will facilitate the further understanding of the population structure of Xoo in Asia.

INTRODUCTION

Xanthomonas oryza pv. oryzae (Xoo) causes bacterial blight, the most important bacterial disease of rice in Asia (Mew, 1987). The disease can cause 30 to 50% yield loss (Adhikari et al., 1994; Exconde et al., 1971). Although some chemicals have been developed to control the disease, none of them has been fully effective under very severe conditions. Host resistance is an important component of an integrated disease management program for bacterial blight (Mew et al., 1992 and 1993). Compared to the long history of rice cultivation, the deployment of genes for resistance to Xoo in commercial rice cultivars is relatively recent. The introduction of these genes for resistance into rice is correlated with a change in the pathogenic diversity of Xoo populations, that is, new races of the pathogen emerge and overcome deployed resistance. These observation have stimulated much curiously concerning the contribution of host genotype and other factors to the genetic diversity of the pathogen. Although, so far, more than 30 resistance genes have been identified and utilized in rice breeding programs (Khush, et al., 1990; Ogawa et al., 1991; Yoshimura et al., 1992 and 1995; Lin et al., 1996; Nagato and Yoshimura, 1998; Zhang et al., 1998; Khush and Angeles, 1999; Chen et al., 2002; Lee et al., 2003;

Thus it is important to understand the structure of pathogen population to determine the best strategy for deployment of resistance. Information on pathogen population structure would include knowledge of pathogen diversity, phylogeny, and the partitioning of variation in time and space. Knowledge of the spatial distribution of pathogen population can aid in the selection of disease resistance sources for a regional crop breeding program. Unfortunately, detail information on pathogen populations is rarely available. Although Xoo populations have been or are being studied in individual countries (Adhikari et al., 1999; Ochiai, et al., 2000), only one comparative studies of molecular variation at an international level had been carried out (Adhikari, et al., 1995). In this article, the population structure of Xoo collected from 1960s to 1970s in several major rice–growing countries countries of South and East Asian is assessed by using PCR-RFLP and virulence analyses.

MATERIAL AND METHODS

Isolation of causal bacterium

Rice leaves affected by bacterial blight were collected from 1960s to 1970s in India, Thailand, Indonesia, Taiwan, Malaysia and Philippine. Diseased leaf samples were cut into small pieces, 1 cm in length including the margin portion of fresh lesions. They were placed in 70% ethyl alcohol for a few seconds, dipped in 1% sodium

Tan et al., 2004; Xiang et al., 2006; Singh et al., 2007), the effectiveness of resistance genes varies over locations due to geographical structuring of the pathogen population. Information on the existing population structure of the pathogen in a region can be useful in the identification and characterization of useful in resistant germ plasm (Leung et al., 1993).

¹ Laboratory of Plant Pathology, Division of Plant Pathology and Pesticide Science, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University, Fukuoka 812–8581, Japan

² Laboratory of Plant Breeding, Division of Genetics and Plant Breeding, Department of Applied Genetics and Pest Management, Faculty of Agriculture, Kyushu University, Fukuoka 812–8581, Japan

 $[\]hbox{$*$ Corresponding author (E-mail: nafuruya@agr.kyushu-u.ac.jp)}\\$

20 S. JYUFUKU et al.

Table 1. Strains of *Xanthomonas oryzae* pv. *oryzae* used in this experiment

| Strain | Place, Country | Year isolated | Strain | Place, Country | Year isolated | |
|--------|---------------------|---------------|--------|-----------------------------|---------------|--|
| 1–1 | Chiangmai, Thailand | 1979 | 11–1 | Subang, Indonesia | 1979 | |
| 1–3 | Chiangmai, Thailand | 1979 | 12-4 | Cirebon, Indonesia | 1979 | |
| 1-6 | Ang-thon, Thailand | 1979 | 14–4 | Tulis, Batang, Central Jawa | 1979 | |
| 6-1 | Chiangmai, Thailand | 1979 | 15-1 | Central Jawa, Indonesia | 1979 | |
| 7–16 | Chai Nat, Thailand | 1979 | 16-1 | Central Jawa, Indonesia | 1979 | |
| TB7309 | Thailand | 1973 | 28-1 | Balaraja, Indonesia | 1979 | |
| TB7421 | Thailand | 1974 | 29-1 | Kragilang, Indonesia | 1979 | |
| TB7538 | Thailand | 1975 | 32-1 | Palangbesi, Indonesia | 1979 | |
| TB7545 | Thailand | 1975 | 34-1 | Padangbesi, Indonesia | 1979 | |
| TB7642 | Thailand | 1976 | 35-2 | Sumatra, Indonesia | 1979 | |
| TB7702 | Thailand | 1977 | Xo7604 | Indonesia | 1976 | |
| TB7809 | Thailand | 1978 | Xo7710 | Indonesia | 1977 | |
| TB7810 | Thailand | 1978 | | | | |
| | | | N6821 | Taiwan | 1968 | |
| 1-1 | Iloilo, Philippine | 1978 | N6822 | Taiwan | 1968 | |
| 1–4 | Iloilo, Philippine | 1978 | | | | |
| 2-4 | Iloilo, Philippine | 1978 | 2-4 | Hyderabad, India | 1979 | |
| 2-6 | Iloilo, Philippine | 1978 | 2-3 | Hyderabad, India | 1979 | |
| 3-1 | Iloilo , Philippine | 1978 | 5–2 | CRRI, India | 1979 | |
| 3–3 | Iloilo, Philippine | 1978 | 6–2 | CRRI, India | 1979 | |
| Pxo8 | Philippine | 1970s | 7-2 | CRRI, India | 1979 | |
| Pxo30 | Philippine | 1970s | 9–3 | Cuttack, India | 1979 | |
| Pxo50 | IRRI, Philippine | 1970s | N6914 | Rapsagar, India | 1969 | |
| Pxo81L | Philippine | 1970s | N6915 | Dumraan, India | 1969 | |
| Pxo87 | Isabela, Philippine | 1974 | N6917 | India | 1969 | |
| Isal 2 | Philippine | 1970s | | | | |
| Kr4 | Philippine | opine 1970s | N6807 | Malaysia | 1968 | |
| | | | N6808 | Malaysia | 1968 | |
| 5—4 | Pagelaran, Lampung | 1979 | N6809 | Malaysia | 1968 | |
| 7–4 | Simpang Kanan | 1979 | N6810 | Malaysia | 1968 | |
| 10-2 | Karawang, Indonesia | 1979 | N6811 | Malaysia | 1968 | |

hypochlotite solution for 1 minute and rinsed in sterilized distilled water. Each sample was then homogenized with 10 ml sterilized distilled water. The resulting suspension was diluted with sterilized distilled water and $100\,\mu$ l of appropriate dilution were spread on PSA medium (Wakimoto, 1955), and the plates were incubated at 30 °C for 4 days. Through this experiment, single-colony isolation was made by using Suwa's medium (Suwa, 1962). The viscous and yellow bacterial colonies that subsequently developed was subcultured on PSA medium and grown at 30 °C for 2 days. For long-term preservation, the bacterial cells suspended in 10% (w/v) skim-milk containing 0.05% L-glutamic acid were lyophilized. For inoculation to rice plant, the bacterium was grown on PSA medium at 30 °C for 2 days, and the culture was suspended in sterilized distilled water to reach a concentration of ca 10⁸ cfu/ml. The suspension was used as inoculum. All strains of Xoo used in this study was presented in Table 1.

Pathogenicity test

The near–isogenic lines have been employed to determine the race composition of *Xoo* in different rice growing countries of Asia (Ogawa *et al.*, 1991). This international differential lines (IR–BB series) and one cultivar, Taichung Native 1 (TN1) (Taura *et al.*, 1987) were used for the experiments. IR24 was used as a susceptible check. The rice seedlings were grown in seedling box, and were transplanted individually to plastic pots (1/50000a) in air conditioned greenhouse. Inoculation

was preformed by the leaf clipping method (Kauffman et al., 1973) at booting stage. Tips of rice leaves were clipped off with scissors dipped in the bacterial suspension. Two weeks after inoculation, lesion length on 10 inoculated leaves were measured. For typing of disease response, each plant was classified as resistant (R) if the mean lesion length was between 0 and 5 cm. Plants with lesion length from 5 to 15 were classified as moderately resistant (M). Plants with lesion length more than 15 cm were classified qualitatively as susceptible (S).

Isolation of genomic DNA

Genomic DNA of Xoo strains were prepared from 5—ml PS broth cultures grown overnight. The DNA from each bacterial strain was extracted by DNeasy Tissue Kit (Qiagen) according to the manufacture's instructions.

PCR fingerprinting

Genotypic diversity was evaluated by PCR-based assay using IS1112 primers set as described by George et al. (1995) and Shanti et al. (2001). The IS1112 primer sequences corresponding to JEL 1 (5'-CTCAGGTCAGGTCGCC-3') and JEL 2 (5'-GCTCTACAATCGTCCGC-3') were used to determine if they could reveal polymorphism in Xoo isolates in Asia.

All amplifications was carried out in a final volume of $25\,\mu l$ and were performed in a programmable thermal cycler (MyCycler, BIO–RAD). The reaction mixtures for PCR contained (final concentration) 50 pmol of primer, 50 ng of template DNA, $312.5\,\mu M$ each deoxynucleoside

triphosphate (dNTP) (Sigma Chemical Co.,), two units of Taq polymerase (Promega Corp.), and 10% (vol/vol) dimethyl sulfoxide (DMSO). The 5 × reaction buffer stock solution contained 10 mM Tris-HCl (pH8.3), 25 mM KCl, 3.5 mM MgCl₂, and 160 ng of bovine serum albumin per ml. PCR conditions were as followed, initially denatured for 1 min at 94 °C, and then subjected to 30 cycles of PCR (10 s of denaturation at 94 °C, 1 min of annealing at 62 °C, and 8 min of extension at 65 °C). After completion of PCR, samples were stored at 4 °C until gel electrophoresis. A $10-\mu l$ portion of each amplified PCR product was resolved on a gel containing a mixture of 0.75% agarose in $0.5 \times \text{Tris-borate-EDTA}$ buffer (89 mM Tris, 89 mM boric acid, and 0.5 M EDTA, pH 8.0), stained with ethidum bromide, and photographed on an UV transilluminator. Experiments were repeated three times to confirm DNA band identities and differences.

The banding pattern of each isolate was recorded in binary form, 1 representing the presence and 0 the absence of each band. A cluster analysis of 44 strains of *Xoo* was performed by an unweighted pair group method with arithmetic averages (UPGMA) using the statistics software package STAT Partner NEC (2.0).

RESULTS

Race determination

Variation in the pathogenicity of bacterial strains from several South and East Asian countries was examined. As indicated in Table 2, the strains of *Xoo* were polymorphic variable for virulence on the ten near–isogenic lines and TN1 which carry with resistance genes *Xa1*, *Xa2*, *Xa3*, *Xa4*, *xa5*, *Xa7*, *Xa10*, *Xa11*, *Xa14*, *Xa21* and *Xa14*. Among 57 strains, three strains from Thailand (TB7421, 6–1, 1–3), five strains from Indonesia (5–4, 7–4, 14–4, 29–1, 34–1), three strains from Philippine (2–4, Pxo87, Isal 2) and two strains from India (2–4, 7–2) were not determined race since their lesion lengths on IR24 as a check variety were less than 15 cm. The 44 strains of *Xoo* were classified into 13 races. This diversity is influenced by the country of collection. Of these, races A (two strains from India), B (two strains from Indonesia

and 3 strains from India) and D (strain from India) were virulent on all host differentials, while the other races were incompatible with at least one of the hosts.

DNA fingerprinting and population substructure

PCR fingerprinting of Asian strains of *Xoo* were generated with the IS*1112*–based PCR primers (Fig. 1). There were 10 major band positions scored in the PCR–based fingerprints.

The genetic relationships among strains and the banding patterns were analyzed by cluster and phylogenic analyses. At 12.6 of genetic distance in cluster analysis of Fig. 2, Asians 44 strains of Xoo were divided into four clusters named from L1 to L4, and percentage of strains belonging to these clusters were 21.0% (n=9), 11.4% (n=5), 25.0% (n=11), and 43.2% (n=19), respectively. Cluster L1 contained 5 strains from the Philippines. Cluster L2 contained 5 strains from Thailand and Philippines which were race H. Almost strains from India formed cluster L3 and varied in broad–spectrum

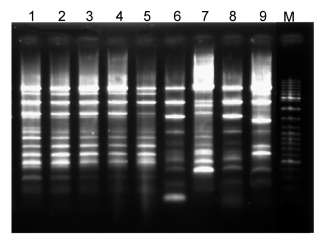


Fig. 1. Polymerase chain reaction fingerprint patterns of Xanthomonas oryzae pv. oryzae strains generated with IS1112-based primers (JEL1/JEL2). Lane 1, Philippines PXO50; Lane 2, Philippines PXO81L; Lane 3, Philippines PXO8; Lane4, Philippines PXO30; Lane 5, Philippines Kr4; Lane 6, Philippines 1-4; Lane 7, Indonesia 10-2; Lane 8, Indonesia 11-1; Lane 9, Indonesia12-4; M, DNA Ladder

Table 2. Pathogenicity analysis of the 44 strains of Xanthomonas oryzae pv. oryzae on the 12 rice cultivars

| Differential variety | Resistance gene | Race | | | | | | | | | | | | |
|---------------------------|--------------------|------|---|---|---|---|---|---|---|---|---|---|---|---|
| | | A | В | С | D | Е | F | G | Н | I | J | K | L | M |
| IR24 | Xa16, Xa18 | S | S | S | S | S | S | S | S | S | S | S | S | S |
| IR-BB 1 | Xa1 | S | S | S | S | S | S | S | S | S | S | S | S | R |
| IR-BB 2 | Xa1, Xa2 | S | S | S | S | S | S | S | S | S | S | S | S | R |
| IR-BB 3 | Xa3 | S | S | S | S | S | S | M | M | S | S | M | M | M |
| IR-BB 4 | Xa4 | S | S | S | M | S | M | S | M | M | S | S | M | M |
| IR-BB 5 | xa5 | M | S | M | M | R | R | R | R | R | R | R | R | R |
| IR-BB 7 | Xa7 | S | S | S | S | R | R | R | R | R | R | R | R | R |
| IR-BB 10 | Xa10 | S | S | S | S | S | S | S | S | S | R | R | S | S |
| IR-BB11 | Xa11 | S | S | R | S | S | S | S | S | S | S | S | S | R |
| IR-BB 14 | Xa14 | S | S | S | M | S | S | S | S | R | S | S | R | R |
| IR-BB 21 | Xa21 | S | M | M | M | M | M | M | M | M | M | M | R | R |
| TN1 | Xa14 | S | S | S | S | S | S | S | S | S | S | S | R | R |
| Number of present strains | | 2 | 5 | 2 | 1 | 9 | 5 | 3 | 7 | 2 | 3 | 3 | 1 | 1 |

22 S. JYUFUKU et al.

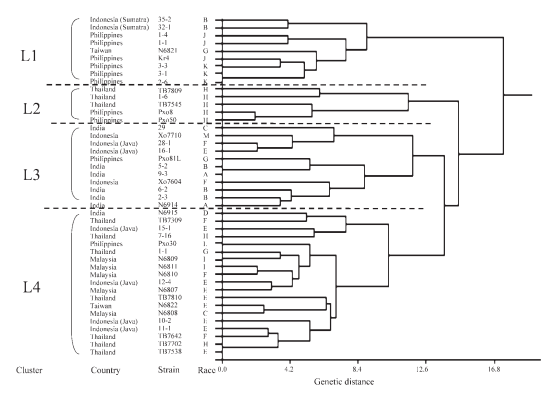


Fig. 2. Dendrogram construction with the statistics software package STAT Partner NEC (2.0) using polymerase chain reaction—based fingerprinting of 44 strains of *Xanthomonas oryzae* pv. *oryzae* from Asian countries. Race numbers correspond to those in Table 2.

pathogenicity. Custer L4 was the most heterogeneous group and contained almost strains of race E. The collection of strains from two different islands (Sumatra and Jawa) in Indonesia was clearly distinct in the genetic and pathogenic characters. A partial relationship was found the genetic groups determined using PCR-based finger-printing by IS1112 primer, races based on virulence against 11 resistance genes and regional origins of each strain isolated.

DISCUSSION

The pathotype (race) of Xoo strains obtained from 1960s to 1970s in South and East Asian countries was examined using the international differential cultivars carrying the Xa1, Xa2, Xa3, Xa4, xa5, Xa7, Xa10, Xa11, Xa14, Xa16, Xa18 and Xa21 genes for resistance (Ogawa, et al., 1991). The strains were divided into 13 races on the basis of their pathogenicity to 10 international differential lines and TN1. Virulence to IR-BB5 (xa5) and IR-BB21 (Xa21) was regionally differentiated. Many south Asian rice cultivars possess xa5, and most of the strains from India were virulent to xa5. Strains from Malaysia, Philippines, Thai and Taiwan were not virulent to xa5 and Xa21. Race distribution was specific to each country. Cultivated cultivars in certain region could be considered as a determining factor of race distribution. For example, Japanese race II which is virulent to rice varieties with the resistant gene Xa1 was generally found in northern Japan where Kinmaze group cultivars, possessing no resistant gene, are cultivated extensively (Horino, 1978). Thus, other factors may possibly be associated with race distribution.

Multilocus molecular makers (insertion sequence elements) has been used in conjunction with virulence typing to evaluate the diversity and structure of *Xoo* populations within and among countries in Asia. Based on the cluster analysis of the RFLP data, a partial relationships among clusters, virulence, and national or regional origin were found. The clonal populations within a country or region are likely a consequence of at least two factors, physical geographic barriers and the similarity of rice varieties grown on a national basis.

In general, regionally defined pathogen populations in Asia were found to be distinct. This finding could be either to show pathogen migration or dispersal or to spatial partitioning of host genotypes (different cultivar preference between regions). Although populations within a region generally were similar, in some cases genetically similar strains were detected in different regions, suggesting the migration of strains between counties, possibly as a consequence of germ plasm exchange. Adhikari et al. (1995) grouped strains from different Asian countries using probes IS1112 and avrXa10 and suggested that the pathogen migrated within Asia. George et al. also showed that the same strains from Indonesia and the Philippines were very similar using PCR- and RFLP-based fingerprinting of the insertion sequence. The same haplotypes might have widespread in these two countries. Our study indicated the strains from both Indonesia and Philippines were relatively diverse in phylogenic relationship. Systematic sampling of Xoo populations within the various countries is needed to determine whether the differences in pathogenic diversity among strains suggested by these results reflect true differences in diversity between regions.

Our results show that although population of the bacterial blight pathogen of rice are very diverse, they exhibit regional differentiation. This information may provide a preliminary basis to design strategies for usable different sources of resistance to the pathogen. For example, although recent studies showed that strains which are virulent to Xa21 widely distributed in Korea (Lee $et\ al.$, 1999) and Nepal (Adhikari $et\ al.$, 1999), the xa5 and Xa21 genes for resistance might be useful in many other countries. The results of this and further DNA fingerprinting analyses would be useful in the selection of strains for additional resistance screening and investigation of pathogenic evolution.

REFERENCES

- Adhikari, T. B., T. W. Mew, and P. S. Teng 1994 Progress of bacterial blight on rice cultivars carrying different Xa—genes for resistance in the field. *Plant Dis.*, **78**: 73–77
- Adhikari, T. B., C. M. Vera Cruz, Q. Zhang, R. J. Nelson, D. Z. Skinner, T. W. Mew and J. E. Leach 1995 Genetic diversity of *Xanthomonas oryzae* pv. oryzae in Asia. Appl. Environ. Microbiol., **61**: 966–971
- Adhikari, T. B., T. W. Mew and J. E. Leach 1999 Genotypic and pathotypic diversity in Xanthomonas oryzae pv. oryzae in Nepal. Phytopathology, 89: 687–694
- Chen, H., S. Wang and Q. Zhang 2002 A new gene for bacterial blight resistance in rice located on chromosome 12 identified from Mingui 63, and elite restore line. *Phytopathology*, **92**: 750–754
- Exconde, O. R., O. S. Opina, and A. Phanomsawara 1971 Yield losses due to bacterial leaf blight. *Philipp. Agric.*, **57**: 120–140
- George, M. L. C., J. E. Leach and R. J. Nelson 1995 DNA fingerprinting of *Xanthomonas oryzae* pv. *oryzae* using IS*1112* based polymerase chain reaction. Int. *Rice Res. Notes*, **20**(3): 30–31
- George, M. L. C., M. Bustamam, W. T. Cruz, J. E. Leach and R. J. Nelson 1986 Movement of *Xanthomonas oryzae* pv. oryzae in southeast Asia detected using PCR-based DNA fingerprinting. *Phytopathology*, **87**: 302–309
- Horino, O. 1978 Distribution of pathogenic strains of *Xanthomonas oryzae* (Uyeda et Ishiyama) Dowson in Japan in 1873 and 1975. *Ann. Phytopatol. Soc. Jpn.*, **44**: 297–304 (Jpn. Engl. Sum)
- Kauffman, H. E., A. P. K. Reddy, S. P. Y. Hsieh and S. D. Merca 1973 An improved technique for evaluating resistance of rice varieties to *Xanthomonas oryzae*. *Plant Dis. Rep.*, **57**: 537–541
- Khush, G. S., E. R. Angeles 1999 A new gene for resistance to race 6 of bacterial blight in rice, Oryzae sativa L. Rice Genet. Newsl., 16: 92–93
- Khush, G. S., E. Bacalangco and T. Ogawa 1990 A new genes for resistance to bacterial blight from O. longisaminata. Rice Genet. Newsl., 67: 121–122
- Lee, K. S., S. Rasabandith, E. R. Angeles, G. S. Khush 2003 Inheritance of resistance to bacterial blight in 21 cultivars of rice. *Phytopathology*, **93**: 147–152
- Lee, S. W., S. H. Choi, S. S. Han, D. G. Lee and B. Y. Lee 1999 Distribution of *Xanthomonas oryzae* pv. *oryzae* strains virulent to *Xa21* in Korea. *Phytopathology*, **89**: 928–933
- Leung, H., R. J. Nelson and J. E. Leach 1993 Population structure of plant pathogenic fungi and bacteria. $Adv.\ Plant$ Pathol., ${\bf 10}:$ 157–205

- Lin, X. H., D. P. Zhang, Y. F. Xie, H. P. Gao, Q. Zhang 1996 Identifying and mapping a new gene for bacterial blight resistance in rice based on RFLP markers. *Phytopathology*, 86: 1156-11593
- Mew, T. W. 1987 Current status and future prospects of research on bacterial blight of rice. Annu. Rev. Phtytopathol., 25: 359–382
- Mew, T. W., C. M. Vera Curz and E. S. Medalla 1992 Changes in race frequency of *Xanthomonas oryzae* pv. oryzae in response to rice cultivars planted in the Philippines. Plant Dis., 76: 1029–1032
- Mew, T. W., A. M. Alvarez, J. E. Leach and J. Swing 1993 Focus on bacterial blight of rice. *Plant Dis.*, **77**: 5–12
- Nagato, Y. and A. Yoshimura 1998 Report of the committee on gene symbolization, nomenclature and linkage groups. *Rice Genet. Newsl.*, 15: 13–74
- Ochiai, H., O. Horino, K. Miyajima and H. Kaku 2000 Genetic diversity of *Xanthomonas oryzae* pv. *oryzae* strains from Sri Lanka. *Phytopathology*, **90**: 415–421
- Ogawa, T., T. Yamamoto, G. S. Khush, and T. W. Mew 1991 Breeding of near-isogenic lines of rice with single genes for resistance to bacterial blight pathogen (*Xanthomonas* campestris pv. oryzae). Jpn. J. Breed., **41**: 523–529
- Saitou, N. and M. Nei 1987 The neighbor–joining method: a new method for reconstructing phylogenetic trees. *Mol. Biol. Evol.*, **4**: 406–425
- Shanti, M. L., M. L. C. George, C. M. Vera Cruz, M. A. Bernardo, R. J. Nelson, H. Leung, J. N. Reddy and R. Sridhar 2001 Identification of resistance genes effective against rice bacterial blight pathogen in eastern India. *Plant Dis.*, 85: 506–512
- Singh, K., Y. Vikal, R. Mahajan, K. K. Cheema, D. Bhatia, R. Sharma, J. S. Lore, T. S. Bharaj 2007 Three novel bacterial blight resistance genes identified, mapped and transfer to cultivated rice O. sativa L. In: "The 2nd International Conference on Bacterial Blight of Rice", Nanjing, China, October 1–3, pp. 76–77
- Suwa, T. 1962 Studies on the culture media of Xanthomonas oryzae (Uyeda et Ishiyama) Dowson. Ann. Phytopathol. Soc. Jpn., 27: 165–171 (in Japanese with English summary)
- Taura, S., T. Ogawa, R. E. Tabien, G. S. Khush, A. Yoshimura and
 T. Omura 1987 The specific reaction of Taichung Native 1
 to Philippine races of bacterial blight and inheritance to race
 5 (PX0112). Rice Genet. Newsl., 4: 101–102
- Tan, G. X., X. Ren, Q. M. Weng, Z. Y. Shi, L. L. Zhu, G. C. He 2004 Mapping of a new resistance gene to bacterial blight in rice line introgressed from *Oryzae officinalis*. Acta Genet. Sin., 31: 724–729
- Wakimoto, S. 1955 Studies on the multiplication of OP, phage (Xanthomonas oryzae bacteriophage) 1. One–step growth experiment under various conditions. Sci. Bull. Fac. Agric. Kyushu Univ., 15: 151–160
- Xiang, Y., Y. L. Cao, C. Q. Xu, X. H. Li, S. P. Wang 2006 Xa3, conferring resistance for rice bacterial blight and encoding a receptor kinase-like protein, is the same as Xa26. Theor Appl Genet., 113: 1347-1355
- Yoshimura, S. A. Yoshimura, R. J. Nelson, T. W. Mew and N. Iwata 1992 RFLP analysis of introgressed chromosomal segments in three near–isogenic lines of rice for bacterial blight resistance genes, *Xa1*, *Xa3* and *Xa4*. *Jpn. J. Genet.*, **67**: 29–37
- Yoshimura, S., A. Yoshimura, A., N. Iwata, S. R. McCouch, M. L. Rbenes, M. R. Baraoidan, T. W. Mew and R. J. Nelson 1995 Tagging and combining bacterial blight resistance genes in rice using RAPD and RFLP markers. *Mol. Breed.*, 1: 375–387
- Zhang, Q., S. C. Lin, B. Y. Zhao, C. L. Wang, W. C. Yang, Y. L. Zhou, D. Y. Li, C. B. Chen, L. H. Zhu 1998 Identification and tagging a new gene for resistance to bacterial blight (Xanthomonas oryzae pv. oryzae) from O. rufipogon. Rice Genet. Newsl., 15: 138–142