**EPPO Datasheet: *Xanthomonas oryzae pv. oryzicola***

Last updated: 2022-09-29

**IDENTITY**

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| **Preferred name:** *Xanthomonas oryzae pv. oryzicola* **Authority:** (Fang et al.) Swings et al. **Taxonomic position:** Bacteria: Proteobacteria: Gammaproteobacteria: Lysobacterales: Lysobacteraceae **Other scientific names:** *Xanthomonas campestris pv. oryzicola* (Fang et al.) Dye, *Xanthomonas oryzicola* Fang et al., *Xanthomonas translucens f. sp. oryzicola* (Fang et al.) Bradbury **Common names in English:** BLS, bacterial leaf streak of rice, leaf streak of rice [view more common names online...](https://gd.eppo.int/taxon/XANTTO/) **EPPO Categorization:** A1 list **EU Categorization:** A1 Quarantine pest (Annex II A) [view more categorizations online...](https://gd.eppo.int/taxon/XANTTO/categorization) **EPPO Code:** XANTTO | 1759.jpg [more photos...](https://gd.eppo.int/taxon/XANTTO/photos) |

**Notes on taxonomy and nomenclature**

Bacterial leaf streak of rice, caused by *Xanthomonas oryzae* pv. *oryzicola*, has quite similar symptoms to bacterial leaf blight of rice, caused by *Xanthomonas* *oryzae* pv. *oryzae*, [**see EPPO Datasheet on *X. oryzae* pv. *oryzae***](https://gd.eppo.int/taxon/XANTOR/datasheet). Bacterial leaf streak was first observed (but thought for a considerable time to be bacterial leaf blight) in the Philippines in 1918 (Reinking, 1918). It was ‘rediscovered’ in China in 1957, described as bacterial leaf streak of rice and the causal bacterium was named *Xanthomonas oryzicola* (Fang *et al*., 1957). *X. oryzicola* was reclassified in later years, first as *X. translucens* f. sp. *oryzicola,* and then as *X. campestris* pv. *oryzicola* (Bradbury, 1971; Aldrick *et al*., 1973; Dye, 1978). The combination *Xanthomonas translucens* (Jones *et al*., 1917) f.sp. *oryzae*(Uyeda & Ishiyama, 1928) Pordesimo 1958 has been incorrectly used (see Bradbury, 1971, Aldrick *et al*, 1973).

On the basis of a polyphasic taxonomical study, Swings *et al*. (1990) placed both bacteria as pathogenic varieties within the species *Xanthomonas oryzae*as*X. oryzae*pv. *oryzicola*and*X. oryzae*pv*. oryzae.*

For a long time, and unlike *X. oryzae* pv. *oryzae*, it was not possible to discriminate pathogenic races for *X. oryzae* pv. *oryzicola*(Ou, 1985), but recently some race variation was reported from Southern China (Yang *et al*., 2020). Variability among *X. oryzae*pv. *oryzicola* strains based on genomic studies is very high (Adhikari & Mew, 1985; Gonzalez *et al*., 2007; Zhao *et al*., 2012; Wonni *et al*., 2011, 2014). Whole genome sequencing was performed with the pathotype strain of *X. oryzae* pv. *oryzicola* (WHRI 5234 = NCPPB 1585 = ICMP 5743, isolated in Malaysia in 1964, Michalopoulou *et al*., 2018).

A strain slightly deviating from *X. oryzae* pv. *oryzicola* and *X. oryzae* pv. *oryzae* isolated from the (invasive) perennial grass weed species *Leersia hexandra* (southern cutgrass or rice swamp grass) was described in 1957 from China by Fang *et al*. (1957) as *X. leersiae*. Based on comparative genomics of strains from China, Burkina Faso, India, Mali and Uganda it was later described as *X. oryzae* pv. *leersiae*. *X. oryzae* pv. *leersiae* is most closely related to *X. oryzae* pv. *oryzicola*, but it is also a close relative of *X. oryzae* pv. *oryzae* (Lang *et al*., 2019).

*X. oryzae* strains occurring in the United States, and first reported in 1989 (Jones *et al*., 1989), appear to be (slightly) different from *X. oryzae*pv*. oryzicola*, *X. oryzae*pv*. oryzae,* and *X. oryzae* pv. *leersiae*. These strains have low virulence on rice, they have not yet been distinguished at pathovar level and are called (also in this document) *X. oryzae* ‘USA’ (Xu & Gonzales, 1991; Gonzalez *et al*., 2007; Triplett *et al*., 2011; Hajri *et al*., 2012; Lang *et al*., 2019).

*X. oryzae* as a species, is genomically closely related to *X. vasicola* pv. *vasculorum*, causing leaf scald of maize, sugarcane and some other Poaceae and *X. vasicola* pv. *musacearum*, causing banana xanthomonas wilt. It is only distantly related to other *Xanthomonas* species and pathovars pathogenic to Poaceae, such as the host specialized pathovars of *X. translucens*and *X. albilineans*(Rodriguez *et al*., 2012; Hersemann *et al*., 2017; Sapkota *et al*., 2020).

For additional taxonomic and nomenclatorial information see CABI (2022a and b) and Niño-Liu *et al*. (2006).

**HOSTS**

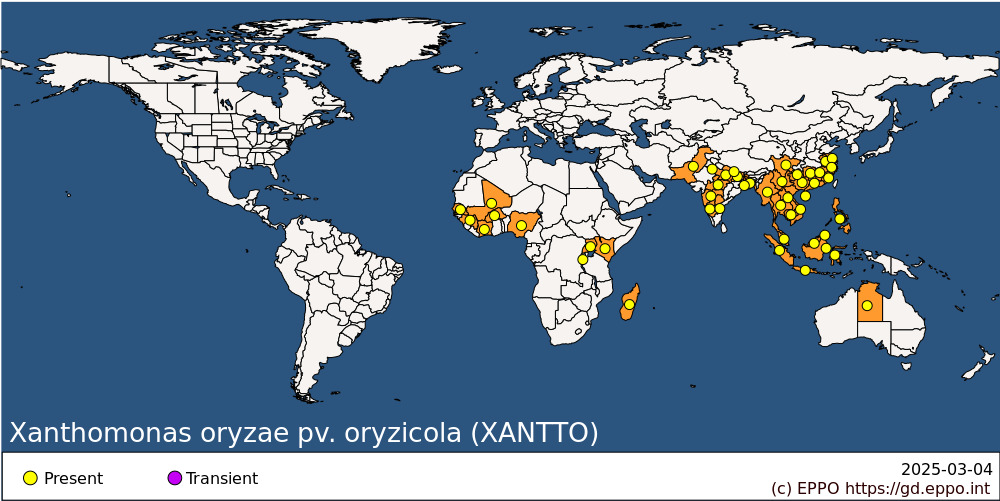
The principal host of *X. oryzae* pv. *oryzicola* is rice, *Oryza sativa*. The sticky, short-grained *O. sativa* subsp. *Japonica* (syn. *O. oryza* subsp. *Sinica*) is less susceptible to *X. oryzae* pv. *oryzicola* than the non-sticky, long-grained *O. sativa* subsp. *Indica*. In Europe, *O. sativa* subsp. *Japonica* is mainly grown (Agri-food Data Portal, 2022; Cai *et al*., 2013; Kraehmer *et al*., 2017).

Other important hosts belong to the Poaceae family, both wild and cultivated, annual and perennial species (Reddy & Nayak, 1975; Leyns *et al*., 1984; CABI, 2021; EFSA, 2018)*.*

**Host list:** *Brachiaria lata*, *Digitaria horizontalis*, *Echinochloa colonum*, *Eleusine indica*, *Leersia hexandra*, *Leptochloa mucronata*, *Oryza barthii*, *Oryza glaberrima*, *Oryza latifolia*, *Oryza longistaminata*, *Oryza minuta*, *Oryza officinalis*, *Oryza sativa*, *Paspalum scrobiculatum*, *Paspalum vaginatum*, *Rottboellia cochinchinensis*, *Zizania aquatica*, *Zizania palustris*, *Zoysia japonica*

**GEOGRAPHICAL DISTRIBUTION**

Bacterial leaf streak was first reported in the Philippines in 1918 and is widely present in tropical and subtropical Asia, including China, Malaysia, India, Indonesia, and also in Northern Australia (under the old and incorrect name *Xanthomonas translucens* f.sp. *oryzae*,Aldrick *et al*., 1973) and West and East Africa, including Madagascar (CABI/EPPO, 2015). It has not been reported from temperate regions, and unlike *X. oryzae* pv. *oryzae* ([**see EPPO Datasheet on *X. oryzae* pv.** ***oryzae***](https://gd.eppo.int/taxon/XANTOR/datasheet)) no geographically distinct groups have been determined (Ou, 1985; Mew, 1991).

 **Africa:** Burkina Faso, Burundi, Cote d'Ivoire, Guinea, Kenya, Madagascar, Mali, Nigeria, Senegal, Uganda **Asia:** Bangladesh, Cambodia, China (Anhui, Fujian, Guangdong, Guangxi, Guizhou, Hainan, Hunan, Jiangsu, Jiangxi, Sichuan, Yunnan, Zhejiang), India (Andhra Pradesh, Bihar, Haryana, Karnataka, Madhya Pradesh, Maharashtra, Uttar Pradesh, West Bengal), Indonesia (Java, Kalimantan, Sulawesi, Sumatra), Laos, Malaysia (Sabah, Sarawak, West), Myanmar, Nepal, Pakistan, Philippines, Thailand, Vietnam **Oceania:** Australia (Northern Territory)

**BIOLOGY**

*X. oryzae* pv. *oryzicola* usually enters the host plant through stomata or leaf lesions caused by insects, heavy rain and/or wind. It multiplies in the apoplast of mesophilic parenchyma cells and spreads actively in the intercellular spaces. It causes linear water-soaked to necrotic leaf streaks, without entering the vascular tissues (Mew, 1993). *X. oryzae* pv. *oryzicola* has a strong cell-wall degrading (cellulose) activity. This differs from *X. oryzae* pv. *oryzae* which mainly infects the plant via hydathodes (water pores, connected to vascular tissue) and multiplies and spreads mainly in the vascular tissue (Tsuno & Wakimoto, 1983; Zou *et al*., 2012, Cao *et al*., 2020).

In severe infections, *X. oryzae* pv. *oryzicola* may produce typical yellow orange (amber-coloured) exudate in the form of tiny droplets from stomata on the leaf surface. The droplets dry in the form of sticky tiny beads with or without small stalks, or also in strands. These strands may be spread by dry wind (Ou, 1985; Mew, 1991).

Both *X. oryzae* pv *oryzicola* and *X. oryzae* pv. *oryzae* can be isolated from the rice seed coat (Sakthivel *et al*., 2001; Niño-Liu *et al*., 2006), but only *X. oryzae* pv. *oryzicola* has been reported to be seed transmitted (Fang *et al*, 1957; Shekhawat, 1969; Mew, 1993; Xie & Mew, 1998; EFSA 2018). The bacterium can survive up to 5 months in seeds stored at 15-30 °C and seed transmission is efficient when sown under conditions of high humidity (Devadath, 1984).

The bacterium can persist from one season to the next on infected leaves and leaf debris, but was found not to survive in non-sterile soil (Devadath & Dath, 1970). The bacteria may survive on and in alternate hosts, such as *Leersia hexandra*and *Zizania aquatica* (Reddy & Nayak, 1975; Leyns *et al*., 1984), but this has been infrequently and/or inadequately reported (Ou, 1985; Niño-Liu *et al*., 2006).

Spread within a crop occurs by mechanical contact and via rain and irrigation water. Under favourable conditions (warm and wet with heavy winds) rapid and severe disease development can occur. The bacterium survives for up to 90 days in water at 15-20°C and up to 60 days at 25-45°C (Devadath, 1984). Contaminated irrigation water may spread the bacterium to adjacent fields (Devadath, 1984).

*X. oryzae* pv. *oryzicola* occurs mostly in tropical and subtropical climates and causes damage only under very wet conditions. Without continuous rain, secondary infections no longer occur (Mew, 1993; EFSA 2018).

After infection, temperature is the main determinant of disease development. Higher temperatures (26 - 32°C) favour disease development, lower temperature (below 22°C) restrain it (Devadath, 1984). Heavy nitrogenous fertilization favours disease development as is the case for *X. oryzae* pv. *oryzae*(Devadath, 1984). Insects (such as leafhoppers and grasshoppers), humans, and agricultural equipment can mechanically transmit the bacterium (Devadath, 1984). There is an apparent connection with pest damage since the bacterium readily enters insect-damaged tissue, but the exact role of these insects and that of man and machines is poorly understood.

The pathogenicity of *X. oryzae*pv*. oryzicola,*as for*X. oryzae*pv*. oryzae,* is based on a type-3 secretion system, that injects a range of type-3 effectors into rice cells (Niño-Liu *et al*., 2006; Jiang *et al*., 2020). This includes members of the Transcription Activator-Like Effector family (TALEs), major virulence factors, activating susceptibility genes of the host (Hutin *et al*., 2015). Contrary to *X. oryzae* pv. *oryzae,*which has widely present gene-for-gene resistance based on an avirulence gene (bacterium) and a resistance gene (plant), so called avr-R gene interactions, this has not been identified in the *X. oryzae* pv. *oryzicola*-rice pathosystem. Resistant rice varieties therefore only show (partial), so-called quantitative resistance (Niño-Liu *et al*., 2006; Zhao *et al*., 2004; Hajri *et al* 2012; Cai *et al*, 2017). However, to date, the avrRxo1 effector gene was found to be present in all Asian *X. oryzae* pv. *oryzicola*strains, and as it is likely to be involved in fitness/pathogenicity it is therefore important for resistance breeding (Zhao *et al*. 2004).

A high degree of genetic diversity was observed among Asian (Philippines) and African strains of *X. oryzae* pv. *oryzicola*. Strains from Mali were found to be closely related to those from Malaysia, implicating a possible transfer of the bacterium with planting material from Asia to Africa (Raymundo *et al*. 1999; Gonzalez *et al*., 2007; Wonni *et al*., 2014). In an extensive study, using 75 *X. oryzae* pv. *oryzicola* strains from South-West China and 6 differential rice varieties, Wang *et al.* (2010) discriminated 13 race groups, that showed some geographical differentiation. Yang *et al*. (2020) could discriminate 6 pathotypes of *X. oryzae* pv. *oryzicola* in Southern China, using differential varieties, and these local rice varieties showed various levels of resistance against *X. oryzae* pv. *oryzicola*.

**DETECTION AND IDENTIFICATION**

**Symptoms**

Early symptoms are narrow, dark-green, water-soaked, interveinal streaks of various lengths, initially restricted to the leaf blades. The lesions enlarge, often showing a yellow halo and later turn yellowish-orange to brown (depending on the rice cultivar) and may coalesce. Bacterial ooze is often present on the streaks, visible as tiny amber-coloured drops. In advanced stages, the disease is difficult to distinguish from that caused by *X. oryzae*pv*. oryzae* but lesion margins remain linear (rather than wavy for those caused by *X. oryzae* pv. *oryzae*). It can be noted also that both *X. oryzae*pv*. oryzicola*and*X. oryzae*pv*. oryzae* may occur simultaneously in the same field, and sometimes even in the same plant (Goto, 1992; Mew, 1993). In a final stage, streaks become brown to greyish and may completely wither. Infected florets turn brown or black and the ovary and stamens die. Symptomatic infected seeds show browning of glumes and necrotic endosperm. Symptoms are often associated with those caused by larvae of lepidopterous leaf rollers/folders (e.g., *Cnaphalocrocis medinalis*), and of the rice hispa beetle (*Discladispa armigera*), because bacteria readily enter the damaged tissue resulting from these insect infestations (Ou, 1985; Niño-Liu *et al*., 2006; EFSA, 2018).

**Morphology**

*X. oryzae*pv.*oryzicola*is an aerobic, motile, Gram-negative, non-spore-forming, capsulated rod, occurring singly or in pairs, 1.0-2.5 x 0.4-0.6 µm in size, with one polar flagellum (Bradbury 1970, 1986).

Most of the procedures described for the isolation of *X. oryzae*pv. *oryzae*from rice plants, can also be applied for the isolation of *X. oryzae*pv. *oryzicola*. (EPPO, 2007). Faster growing contaminants often occurring on and in diseased tissues, such as species of *Pantoea* or xanthomonad-like saprophytes may overgrow the slow growing *X. oryzae* pv. *oryzicola*colonies and hinder its isolation from diseased material.

Isolation of *X. oryzae*pv. *oryzicola*from symptomatic material is possible on Peptone Sucrose Agar (PSA), Nutrient Broth Yeast Extract agar medium (NBY), Growth Factor (GF) agar or otherwise using semi- selective media (Agarwal *et al*., 1989; Sakthivel *et al*.,2001; EPPO, 2007). A semi-selective medium, called XOS, is available for detection of *X. oryzae* pv. *oryzicola* from rice seed (Di *et al*., 1991; EPPO, 2007). On nutrient agar (NA), after 3 days of growth, colonies of *X. oryzae* pv. *oryzicola*are circular, entire, smooth, convex, opaque, and pale to straw yellow, 1-2 mm in size. Optimum growth temperature is between 25 and 30°C. For growth on other media, see EPPO, 2007.

**Detection and identification methods**

Detection of *X. oryzae* pv. *oryzicola* in seed, using a detached leaf inoculation method was described by Xie & Mew (1998). The method is based on inoculating leaf segments on agar with seed washings in a moist chamber. For selective recovery from seed, this method and the XOS semi-selective medium of Di *et al*. (1991) can be used.

Furthermore poly- and monoclonal antibodies (genus and pathovar specific) can be used in Immuno-fluorescence and ELISA tests on seed extracts and/or colonies isolated from seeds or leaf/stem material and isolated bacterial cells (Benedict *et al*., 1989). An ELISA kit is commercially available for the detection of *X. oryzae* pv. *oryzicola* (EPPO, 2007). A padlock probe (PLP)-based PCR with dot blot hybridisation was developed for simultaneous detection of *X. oryzae*pv*. oryzicola* and *X. oryzae*pv*. oryzae* by Tian *et al*., 2014. A specific TaqMan probe for its detection in seed was developed by Zhao *et al*. (2007).

Leach *et al*. (1990) used a repetitive DNA sequence (pJEL 101) to distinguish *X. oryzae*pv. *oryzae*from other pathovars and species of *Xanthomonas*. Kang *et al*. (2008) developed a specific PCR detection system (targeting a membrane fusion protein gene) for *X. oryzae*pv. *oryzicola*. Other specific TaqMan-based multiplex PCRs for detection and discrimination of *X. oryzae*pv*. oryzicola*and *X. oryzae*pv*. oryzae* were developed and validated by Lang *et al*. (2010), Noh *et al*. (2012), Kang *et al* (2012) and Lee & Vera Cruz (2014).

Lang *et al*. (2014) developed a sensitive and rapid loop-mediated isothermal amplification (LAMP) test, using primer sets to distinguish not only *X. oryzae*pv*. oryzicola* and *X. oryzae pv. oryzae,* but also the Asian and African lines of *X. oryzae*pv*. oryzae*.

A SYBR green-based multiplex PCR for the detection and identification of *X. oryzae*pv*. oryzicola, X. oryzae*pv*. oryzae* and *Burkholderia glumae* (causing bacterial grain rot of rice) was developed by Lu *et al*. (2014). Kang *et al*. (2016) also developed a multiplex PCR for the detection of the same three bacteria. Cui *et al*. (2016) developed a multiplex conventional and real-time PCR for the simultaneous detection of six bacterial pathogens of rice, including *X. oryzae* pv. *oryzicola, X. oryzae* pv. *oryzae*, *Pseudomonas fuscovaginae* (rice sheath brown rot), *Burkholderia glumae*, *B. gladioli* (bacterial panicle blight of rice) and *Acidovorax avenae* subsp. *avenae* (bacterial brown stripe of rice). A validated multiplex PCR to detect *P. fuscovaginae*, *X. oryzae*pv. *oryzicola*and*X. oryzae*pv. *oryzae*, *Burkholderia* (both *B. glumae*and *B. gladioli*) as well as *Sphingomonas* and *Pantoea* spp. was published by Bangratz *et al*. (2020).

The two pathovars of *X. oryzae*differ in the symptoms induced (Ou, 1985), phenotypic characters (Reddy & Ou, 1974; Vera Cruz *et al*., 1984; Vauterin *et al*., 1995), polyacrylamide gel electrophoresis protein fingerprints (Mew & Vera Cruz, 1979; Kersters *et al*., 1989), serological behavior (Benedict *et al*., 1989) and phage typing (EPPO 2007). Also, on the basis of rep-PCR using BOX-primers discrimination of *X. oryzae* pv. *oryzicola* and *X. oryzae* pv. *oryzae* is possible (Raymundo *et al*., 2008).

As for the whole genus *Xanthomonas*, *X. oryzae* is catalase-positive, unable to reduce nitrate and a weak producer of acids from carbohydrates. Pathovars *oryzicola* and *oryzae* can be differentiated by (a) acetoin production (*X. oryzae*pv*. oryzicola*+, *X. oryzae*pv*. oryzae*–), (b) growth on l-alanine as sole carbon source (*X. oryzae*pv*. oryzicola*+, *X. oryzae*pv*. oryzae*–), (c) growth on 0.2% vitamin-free casamino acids (*X. oryzae*pv*. oryzicola*+, *X. oryzae*pv*. oryzae*–) and (d) resistance to 0.001% Cu (NO3)2 (*X. oryzae*pv*. oryzicola*–, *X. oryzae*pv*. oryzae*+) (Dye & Lelliott, 1974; Reddy & Ou, 1974; Gossele *et al*., 1985; Niño-Liu *et al*., 2006; EPPO 2007). Extensive characterization of *X. oryzae* pv. *oryzicola*, using biochemical, physiological tests and PAGE was performed by Vera Cruz *et al.* (1984). Wonni *et al*. (2014) determined extensive variability between African strains of *X. oryzae* pv. *oryzicola*. Restriction fragment length polymorphism (RFLP) analysis using the effector *avrXa*7 as probe resulted in the identification of 18 haplotypes. PCR using two conserved type III effector (T3E) genes (*xopAJ*and *xopW*) differentiated the strains into an African group where the *xopAJ*was generally not detected, and a group of possible Asian origin.

Six housekeeping genes— *atpD*(ATP synthase β chain), *dnaK*(chaperone protein), *efP*(elongation factor P), *gyrB*(DNA gyrase subunit B), *lepA*(GTP binding protein), and especially *rpoD*(RNA polymerase σ-70 factor) are useful for identification and phylogenetic studies of *X. oryzae* pv. *oryzicola* strains (Afolabi *et al*., 2014; Wonni *et al*., 2014).

Isolates can be tested for pathogenicity on susceptible rice cultivars. For *X. oryzae*pv. *oryzicola*30–45-day old plants of cultivars IR24 or IR50 (International Rice Institute) or local, susceptible varieties can be used. Leaf clipping and spray inoculation methods are available for inoculations (Kauffman *et al*., 1973; Cottyn *et al*., 1994; EPPO, 2007; Afolabi *et al*., 2014). Niño-Liu *et al*. (2005) inoculated plants by dipping them in bacterial mixture and incubating in a growth chamber. Symptoms developed within 6 days.

**PATHWAYS FOR MOVEMENT**

*X. oryzae* pv. *oryzicola* can only move short distances within infected crops. The bacterium is found in association with weeds, even if their role in the disease cycle is less clear than for *X. oryzae* pv. *oryzae* (Leyns *et al*., 1984; Reddy & Nayak, 1975).

There are little substantiated data on spread or transmission in the field by animals other than insects (Ou, 1985; Niño-Liu *et al*., 2006; EFSA, 2018).

Long distance spread can take place via infected rice seeds, and seed transmission is regarded as the main means of dispersal. The planting of disease-free seed is considered of utmost importance in control (Rao, 1987; Xie *et al*., 1990, 1991; Ming *et al*., 1991; Mew, 1993; Xie & Mew, 1998).

**PEST SIGNIFICANCE**

**Economic impact**

Bacterial leaf streak is only of importance in some areas during very wet seasons and where high levels of nitrogen fertilization are used. It does not usually reduce yields if low levels of nitrogen fertilization are applied. In general, bacterial leaf streak is a much less important disease than bacterial leaf blight. In Central India, losses ranged from 5 to 30% depending upon environmental factors and cultivars (Naik *et al*., 1973). In Northern India, disease intensity affecting 80% of leaf area resulted in 61 percent yield loss (Singh *et al*., 1980). In the Philippines, no significant losses were reported in either the wet or dry seasons (Opina & Exconde, 1971).

In West Africa outbreaks of *X. oryzae* pv. *oryzicola* usually showed lower incidence and severity than those of *X. oryzae* pv. *oryzae*, as determined in a 10-year survey (Awoderv *et al*., 1991).

In China, however, *X. oryzae*pv*. oryzicola* has sometimes been more damaging than *X. oryzae*pv*. oryzae*. In Southern China, epidemics of *X. oryzae* pv. *oryzicola* have repeatedly been reported, reducing yield by 10-20% and in some cases reaching up to 40% losses (Xie & Mew 1998; Niño-Liu *et al*., 2006; Cai *et al*., 2017). In Uganda, under favourable conditions (wet/windy/warm temperatures/susceptible varieties) bacterial leaf streak has caused major crop losses (up to 60%) (Andaku *et al*., 2016; EFSA, 2018).

**Control**

The bacterial leaf streak pathogen hardly requires any particular control measures except the use of healthy seed and prevention measures (see below). Neither treatments nor resistance are mentioned to any significant extent in the literature.

***Chemical control***

Chemical seed treatment and field sprays have been reported from India (Shekhawat & Srivastava 1971), using a combination of antibiotics (streptomycin sulphate and tetracycline) and copper-oxychloride. It was also reported that when yield is affected, a copper-based fungicide applied at heading stage can be effective in controlling the disease (ICAR/TNAU, 2022; CABI Plantwise, 2022). The use of antibiotics against plant pathogens is not permitted in many EPPO countries, although in Asia their use is still ongoing and resistance against streptomycin has been reported in China (Xu *et al*., 2010). Recently Chen *et al*. (2019) reported a strong bactericidal effect (in vitro and in vivo) of the bactericide melatonin (N-acetyl-5-methoxytryptamine) on *X. oryzae* pv. *oryzicola* and a reduction of disease incidence by 17%.

***Heat treatment***

Hot water treatment of rice seeds at 52-54°C for 30 min, preceded by 8-10 hour of presoaking at room temperature in water, has been advised and used to cure seeds of *X. oryzae* pv. *oryzae*(Jain, 1970; Reddy, 1983) and is also expected to be effective for *X. oryzae* pv. *oryzicola*.

***Biological control***

Hata *et al*. (2015) found an antagonistic effect on *X. oryzae* pv. *oryzicola*of *Streptomyces*spp. in vitro. In a follow-up (greenhouse) study two strains showed a suppressive effect on bacterial leaf streak due to induction of systemic resistance and growth promoting activity (Hata *et al*., 2021)

Zhang *et al*. (2012) reported promising biocontrol effect of strain Lx-11 of *Bacillus amyloliquefaciens*. This strain appears also to trigger a systemic immunization activity and significantly reduced disease incidence in field experiments (from 60% to 71%) which was better than the effect of a chemical spray with thiadiazole-copper (a bactericide often used in China).

***Plant resistance***

In contrast to bacterial leaf blight, native major resistance genes controlling resistance to bacterial leaf streak have not yet been identified in rice. There are, however loci determining quantitative resistance, such as qBLSR-11-1 (Chen *et al*., 2006) and qBlsr5a, which had a relatively large impact in breeding lines, where the broadly effective rice recessive gene xa5 is involved (Xie *et al*., 2020). In a genome-wide resistance-gene analysis in rice Sattayachiti *et al*. (2020) and Thianthavon *et al*. (2021) stated that this recessive xa5 gene is a very promising candidate to be used in breeding for broad-spectrum resistance. A non-host resistance gene, Rxo1, isolated from maize, and present in transgenic rice was shown to confer high level resistance to bacterial leaf streak (Zhao *et al*., 2005; Jiang *et al*., 2020). Using CRISPR/Cas9 gene editing of two rice varieties Ni *et al*. (2021) obtained rice lines that proved to be resistant to *X. oryzae* pv. *oryzicola* and *X. oryzae* pv. *oryzae*. The original agronomic traits of these lines were not diminished. The dominant locus Xo1 apparently confers complete resistance to African strains of *X. oryzae* pv. *oryzicola* (Triplett *et al*., 2016, Cai *et al*., 2017).

***Prevention and cultural control***

Prophylactic measures (such as use of healthy seeds, adequate fertilization and irrigation, destruction or ploughing under of crop residues, disinfection of machinery and equipment, production of seedlings in boxes and removal of diseased plants and weed hosts from fields and along irrigation canals) have all been found useful in the control of bacterial leaf streak (Devadath, 1984; Goto, 1992; Ou 1985; Shekhawat *et al*., 1972).

**Phytosanitary risk**

Rice cultivation in Europe occurs in Bulgaria, France, Greece, Hungary, Italy Portugal, Romania, the Russian Federation, Spain, Turkey and Ukraine. About 80% of the European Union rice production takes place in Italy (>220 000 ha and Spain (>115 000 ha), another 12% in Greece and Portugal (some 20-25 000 ha each). The remainder is cultivated in Bulgaria, France, Hungary and Romania, (10-20 000 ha each). In non-EU European countries, rice is grown in the Russian Federation (120 000 ha in the Krasnodar region) as well as in Ukraine (25 000 ha). In those countries all rice fields are under irrigation, planted in spring and harvested in autumn. (Agri-food Data Portal, 2022; Ferrero & Nguyen, 2004; Kraehmer *et al*., 2017).

Resistance of European varieties against *X. oryzae*pv*. oryzicola* is unknown. Non-European varieties are only introduced, in small quantities, for breeding (Cai *et al*., 2013; Kraehmer *et al*., 2017). No interceptions of *X. oryzae* pv *oryzicola* have been reported in the EU from 1995 to April 2022 (European Commission, 2022). However, no systematic surveying and monitoring for *X. oryzae*pv*. oryzicola* takes place in Europe

The main risk of introduction is via imported rice seed (germplasm) used for breeding purposes and therefore direct sowing. Milled rice poses a negligible risk, because hulls are removed, and endosperm infection is very rare. Moreover, milled rice has its main destination outside growing areas.

Once introduced by infected seed, further spread could take place via newly infected seed and contaminated water and the bacterium could survive in stubble, straw, weed hosts and volunteer plants.

**PHYTOSANITARY MEASURES**

Phytosanitary (quarantine) measures can be implemented to reduce the risk of long-distance dissemination of the pathogen. It can be recommended that consignments of rice seeds should have been produced from pest-free areas, or from pest-free places of production.

General inspection and sampling procedures for imported rice, which include *X. oryzae*pv*. oryzicola* are described in EPPO Standard PM 3/78(2) *Consignment inspection of seed and grain of cereals*. Seed inspections of rice intended for breeding purposes in international trade may assist in preventing spread of the pathogen to areas with no history of the disease. However, visual inspection of imported seeds is not very reliable due to the occurrence of latent infections and therefore, when material is imported from areas where the disease is known to occur, certification for disease freedom via field inspections and laboratory testing are necessary.

A contingency plan to prepare for possible introductions of *X. oryzae*pv. *oryzicola*in the USA, was developed by the USDA (USDA, 2013).

**REFERENCES**

Adhikari, TB & Mew TW (1985) Host-parasite relationship in bacterial leaf streak of rice caused by *Xanthomonas oryzae* pv. *oryzicola*. *Nepal Journal of Agriculture* **16**, 134-141.

Agarwal PC, Mortensen CN & Mathur SB (1989) Seed-borne diseases and seed health testing of rice. *Phytopathological Papers* No. 30. CAB International, Wallingford, UK.

Agri-food Data Portal (2022) Agri-Food Markets. Rice. European Commission. <https://agridata.ec.europa.eu/extensions/DataPortal/rice.html>

Afolabi O, Milan B, Amoussa R, Koebnik R, Szurek B, Habarugira G, Bigirimana J & Silue D (2014) First report of *Xanthomonas oryzae* pv. *oryzicola*causing bacterial leaf streak of rice in Burundi. *Plant Disease* **98,**1426. <https://doi.org/10.1094/PDIS-05-14-0504-PDN>

Aldrick SJ, Buddenhagen W & Reddy APK (1973) The occurrence of bacterial leaf blight in wild and cultivated rice in Northern Australia. *Australian Agricultural Research* **24**,219-227. <https://doi.org/10.1071/AR9730219>

Andaku JL, Tusiime G, Tukamuhabwa P & Onaga G (2016) Bacterial leaf streak disease of rice: A silent constraint to rice production in Uganda. *RUFORUM Working Document Series* **14**, 523-527.

Awoderv VA, Bangura N & John VT (1991) Incidence, distribution and severity of bacterial diseases on rice in West Africa. *Tropical Pest Management* **37**, 113-117.

Bangratz M, Wonni I, Kini K, Sondo M, Brugidou C, Béna G, Gnacko F, Barro M, Koebnik R, Silué D & Tollenaere C (2020) Design of a new multiplex PCR assay for rice pathogenic bacteria detection and its application to infer disease incidence and detect co-infection in rice fields in Burkina Faso. *PloS One* **15**, e0232115. <https://doi.org/10.1371/journal.pone.0232115>

Benedict AA, Alvarez AM, Berestecky J, Imanaka W, Mizumoto CY, Pollard LW, Mew TW & Gonzalez CF (1989) Pathovar-specific monoclonal antibodies for *Xanthomonas campestris* pv. *oryzae* and for *Xanthomonas campestris* pv. *oryzicola*. *Phytopathology* **79**, 322-328.

Bradbury JF (1970) *Xanthomonas oryzicola*. *CMI Descriptions of Pathogenic Fungi and Bacteria* No. 240. CAB International, Wallingford, UK.

Bradbury JF (1971) Nomenclature of the bacterial leaf streak pathogen of rice. *International Journal of Systematic Bacteriology* **21**, 72.

Bradbury JF (1986) Guide to plant pathogenic bacteria. Farnham Royal, Slough, UK: CAB International, 332pp.

CABI/EPPO (1998) Distribution maps of quarantine pests for Europe (edited by Smith IM, Charles LMF). Wallingford, UK: CAB International, xviii + 768 pp.

CABI/EPPO (2015) *Xanthomonas oryzae* pv. *oryzicola*. [Distribution map]. Distribution Maps of Plant Diseases, No 463 October. Wallingford, UK: CABI, Map 463 (Edition 4). CABI Plantwise (2022) Pest management decision guide: green and yellow list. Bacterial leaf streak on rice. <https://www.plantwise.org/KnowledgeBank/pmdg/20147801491> (last accessed 2022-09).

Cai X, Fan J, Jiang Z, Basso B, Sala F, Spada A, Grassi F & Lu B-R (2013) The puzzle of Italian rice origin and evolution: determining genetic divergence and affinity of rice germplasm from Italy and Asia. *PloS ONE* **8**, e80351. <https://doi.org/10.1371/journal.pone.0080351>

Cai L, Cao Y, Xu Z, Ma W, Zakria M, Zou L, Cheng Z & Chen G (2017) A transcription activator-like effector Tal7 of *Xanthomonas oryzae* pv. *oryzicola* activates rice gene *Os09g29100* to suppress rice immunity. *Science Reports* **7,**5089. <https://doi.org/10.1038/s41598-017-04800-8>

Cao J, Chu C, Zhang M, He L, Qin L, Li X & Yuan M (2020) Different cell wall-degradation ability leads to tissue-specificity between *Xanthomonas oryzae* pv*. oryzae* and *Xanthomonas oryzae* pv*. oryzicola*. *Pathogens,* **9**,187. <https://doi.org/10.3390/pathogens9030187>

Chen C-h, Wei Zheng W, Huang X-m, Zhang D-p & Lin X-h (2006) Major QTL conferring resistance to rice bacterial leaf streak. *Agricultural Sciences in China* **5**, 216-220. <https://doi.org/10.1016/S1671-2927(06)60041-2>

Chen X, Sun C, Laborda P, He Y, Zhao Y, Li C. & Liu F (2019) Melatonin treatments reduce the pathogenicity and inhibit the growth of *Xanthomonas oryzae* pv. *oryzicola*. *Plant Pathology* **68**, 288-296. <https://doi.org/10.1111/ppa.12954>

Cottyn B, Cerez MT & Mew TW (1994) Chapter 7: Bacteria. In: *A Manual of Rice Seed Health Testing* (eds Mew TW & Misra JK), pp.29-46. IRRI, Manila, Philippines.

Cui Z, Ojaghian MR, Tao Z, Kakar KU, Zeng J, Zhao W, Duan Y, Vera Cruz CM, Li B, Zhu B & Xie G (2016) Multiplex PCR assay for simultaneous detection of six major bacterial pathogens of rice. *Journal of Applied Microbiology* **120**, 1357–1367. <https://doi.org/10.1111/jam.13094>.

Devadath S (1984) Bacterial leaf streak of rice. In: Raychaudhuri SP, Verma JP, eds. *Review of Tropical Plant Pathology.*Volume **1**. New Delhi, India: Today and Tomorrow's Printers and Publishers, pp. 155-179.

Devadath S & Dath AP (1970) Epidemiology of *Xanthomonas translucens* f.sp. *oryzae*. *Oryza* **7**, 13-16.

Di M, Ye HZ, Schaad NW & Roth DA (1991) Selective recovery of *Xanthomonas*spp. from rice seeds. *Phytopathology***81**, 1358–1363.

Dye DW (1978) Genus *Xanthomonas*Dowson 1939. In: Young JM, Dye DW, Bradbury JF, Panagopoulos CG & Robbs CF (1978) A proposed nomenclature and classification for plant pathogenic bacteria. *New Zealand Journal of Agricultural Research* **21**, 153-177.

Dye DW & Lelliott RA (1974) Genus II. *Xanthomonas*Dowson 1939. In: *Bergey’s Manual of Determinative Bacteriology*. 8th edition: 243- 249. Eds. R. E. Buchanan and N. E. Gibbons. Williams & Wilkins Co., Baltimore, U.S.A.

EFSA (2018) EFSA Panel on Plant Health: Jeger M, Candresse T, Chatzivassiliou E, Dehnen-Schmutz K, Gilioli G, Gregoire J-C, Jaques Miret JA, MacLeod A, Navajas Navarro M, Niere B, Parnell S, Potting R, Rafoss T, Rossi V, Urek G, Van Bruggen A, Van der Werf W, West J, Winter S, Bragard C, Szurek B, Hollo G and Caffier D. Scientific Opinion on the pest categorisation *of Xanthomonas oryzae* pathovars *oryzae* and *oryzicola*. *EFSA Journal* **16**, 5109, 25 pp. <https://doi.org/10.2903/j.efsa.2018.5109>

EPPO (2007) EPPO Standards. PM7/80 (1) Diagnostic standard *Xanthomonas oryzae*. *EPPO Bulletin* **37**, 543–553.

European Commission (2022) Interceptions of harmful organisms in imported plants and other objects. <https://ec.europa.eu/food/plants/plant-health-and-biosecurity/european-union-notification-system-plant-health-interceptions-europhyt/interceptions_en#plant-interceptions--monthly-reports-2022>

Fang CT, Ken HC, Chen TY, Chu YK, Faan HC & Wu SC (1957) A comparison of the rice bacterial leaf blight organism with the bacterial leaf streak organisms of rice and *Leersia hexandra* Swartz. *Acta Phytopathologica Sinica* **3**. 99-124.

Ferrero A & Nguyen NV (2004) The sustainable development of rice-based production systems in Europe. *Proceedings of the FAO Rice Conference ‘Rice is Life’* **53**, 115-124.

Gonzalez C, Szurek B, Manceau C, Mathieu T, Séré Y & Verdier V (2007) Molecular and pathotypic characterization of new *Xanthomonas oryzae* strains from West Africa. *Molecular Plant-Microbe Interactions* **20**, 534–546.

Goto M (1992) Fundamentals of Bacterial Plant Pathology. Academic Press, San Diego, 342 pp.

Hajri A, Brin C, Zhao S, David P, Feng J, Koebnik R, Szurek B, Verdier V, Boureau T & Poussier S (2012) Multilocus sequence analysis and type III effector repertoire mining provide new insights into the evolutionary history and virulence of *Xanthomonas oryzae*. *Molecular Plant Pathology* **13**, 288–302. <https://doi.org/10.1111/j.1364-3703.2011.00745.x>

Hata EM, Kamaruzaman S, Zainal AMA Mohd TY, Noor AA (2015) In vitro antimicrobial assay of Actinomycetes in rice against *Xanthomonas oryzae* pv*. oryzicola* and as potential plant growth promoter. *Brazilian Archives* *of Biology and Technology* **58**, 821-832. <https://doi.org/10.1590/S1516-89132015060263>

Hata EM, Yusof MT & Zulperi D (2021) Induction of Systemic resistance against bacterial leaf streak disease and growth promotion in rice plant by *Streptomyces shenzhenesis* TKSC3 and *Streptomyces* sp. SS8. *Plant Pathology Journal* **37**, 173-181. <https://doi.org/10.5423/PPJ.OA.05.2020.0083>

Hersemann L, Wibberg D, Blom J, Goesmann A, Widmer F, Vorhölter F-J & Kölliker R (2017) Comparative genomics of host adaptive traits in *Xanthomonas translucens* pv. *graminis*. *BMC Genomics* **18,**35, 18pp. <https://doi.org/10.1186/s12864-016-3422-7>

Hutin M, Pérez-Quintero M, Lopez Camilo AL & Boris S (2015) MorTAL Kombat: the story of defense against TAL effectors through loss-of-susceptibility. *Frontiers in Plant Science* **6**, 535. <https://doi.org/10.3389/fpls.2015.00535>

ICAR/TNAU (2022) Expert System for Paddy. Bacterial leaf streak. *Xanthomonas oryzae*pv*. oryzicola*. [www.agritech.tnau.ac.in/expert\_system/paddy/cpdisbls.html](http://www.agritech.tnau.ac.in/expert_system/paddy/cpdisbls.html) (last accessed 2022-09).

Jain SS (1970) Relation efficacy of some chemicals against bacterial leaf blight of rice caused by *Xanthomonas oryzae*(Uyeda and Ishiyama). *Oryza* **7,** 19-32.

Jiang N, Yan J, Liang Y, Shi Y, He Z, Wu Y, Zeng Q, Liu X & Peng J (2020) Resistance genes and their interactions with bacterial blight/leaf streak pathogens (*Xanthomonas oryzae*) in rice (*Oryza sativa* L.)—an updated review. *Rice* **13,** 12 pp <https://doi.org/10.1186/s12284-019-0358-y>

Jones LR, Johnson AG & Reddy CS (1917) Bacterial blight of barley. *Journal of Agricultural Research* **11**, 625–643.

Jones RK, Barnes LW, Gonzalez CF, Leach JE, Alvarez AM & Benedict AA (1989) Identification of low-virulence strains of *Xanthomonas campestris* pv. *oryzae* from rice in the United States. *Phytopathology* **79**, 984–990

Kang MJ, JK Shim, MS Cho, YJ Seol, JH Hahn, DJ Hwang & DS Park (2008) Specific detection of *Xanthomonas oryzae* pv. *oryzicola*in infected rice plant by use of PCR assay targeting a membrane fusion protein gene. *Journal of Microbiology and Biotechnology* **18**, 1492-1495

Kang MJ, Kima MH, Hwang DJ, Cho MS, Seol Y, Hahn JH, Ryu S & Park DS (2012) Quantitative *in planta* PCR assay for specific detection of *Xanthomonas oryzae* pv. *oryzicola* using putative membrane protein-based primer set. *Crop Protection* **40**, 22-27.Kang IJ, Kang MH, Noh TH, Shim HK, Shin DB & Heu S (2016) Simultaneous detection of three bacterial seed-borne diseases in rice using multiplex polymerase chain reaction. *Plant Pathology Journal* **32**, 575-579. <https://doi.org/10.5423/PPJ.NT.05.2016.0118>

Kauffman HE, Reddy APK, Hsieh SPY & Merca SD (1973) An improved technique for evaluating resistance of rice varieties to *Xanthomonas oryzae*. *Plant Disease Reporter***57**, 537-541.

Kersters K, Pot B, Hoste B, Gillis M, de Ley J (1989) Protein electrophoresis and DNA:DNA hybridizations of xanthomonads from grasses and cereals. *EPPO Bulletin* **19**(1), 51-55.

Kraehmer H, Thomas C & Vidotto F (2017) Rice Production in Europe. In: Chauhan B, Jabran K, Mahajan G (eds) Rice Production Worldwide. Springer, Cham. <https://doi.org/10.1007/978-3-319-47516-5_4>

Lang ML, Hamilton JP, Diaz MGQ, Van Sluys MA, Burgos MRG & Vera Cruz CM (2010) Genomic-based diagnostic marker development for *Xanthomonas oryzae*pv.*oryzae* and *X. oryzae*pv.*oryzicola*. *Plant Disease* **94**, 311–319.

Lang JM, Langlois P, Nguyen MHR, Triplett LR, Purdie L, Holton TA, Djikeng A, Vera Cruz CM, Verdier V & Leach JE (2014) Sensitive detection of *Xanthomonas oryzae* pathovars *oryzae*and *oryzicola* by loop-mediated isothermal amplification. *Applied and Environment Microbiology* **80**, 4519–4530. <https://doi.org/10.1128/aem.00274-14>

Lang JM, Pérez-Quintero A L, Koebnik R, DuCharme E, Sarra S, Doucoure H, Keita I, Ziegle J, Jacobs JM, Oliva R, Koita O, Szurek B, Verdier V & Leach JE (2019) A pathovar of *Xanthomonas oryzae* Infecting wild grasses provides Insight into the evolution of pathogenicity in rice agroecosystems. *Frontiers in Plant Science* **10**, 507,15 pp. <https://doi.org/10.3389/fpls.2019.00507>

Lee D-Y & Vera Cruz CM (2014) Specificity of multiplex PCR in the detection of *Xanthomonas oryzae* pv. *oryzae* and *Xanthomonas oryzae* pv. *oryzicola* in rice (*Oryza sativa* L.) seeds. *Journal of the Korean Society of International Agriculture* **26**, 425-429. <https://doi.org/10.12719/KSIA.2014.26.4.425>

Leyns F, De Cleene M, Swings J & De Ley J (1984) The host range of Genus *Xanthomonas. Botanical Review* **50**, 308-356.

Li ZZ, Zhao H & Ying XD (1985) [The weed carriers of bacterial leaf blight of rice]. *Acta Phytopathologica Sinica* **15**, 246-248.

Lu W, Pan L, Zhao H, Jia Y, Wang Y, Yu X & Wang X (2014) Molecular detection of *Xanthomonas oryzae* pv. *oryzae*, *Xanthomonas oryzae* pv. *oryzicola*, and *Burkholderia glumae* in infected rice seeds and leaves. *The Crop Journal* **2**, 398-406. <https://doi.org/10.1016/j.cj.2014.06.005>.

Mew TW (1991) Bacterial Leaf Streak. In: Rice, Volume I. Production. Chapter 5, 192-194. BS Luh ed. Springer Science+Business Media New York

Mew TW (1993)*Xanthomonas oryzae* pathovars on rice: cause of bacterial blight and bacterial leaf streak. In: *Xanthomonas*. JG Swings & EL Civerolo, eds. Chapman and Hall, New York. pp. 30–40.

Mew TW & Vera Cruz CM (1979) Variability of *Xanthomonas oryzae*: specificity in infection of rice differentials. *Phytopathology***69**, 152–155.

Michalopoulou VA, Vicente JG, Studholme DJ & Sarris PF (2018) Draft genome sequences of pathotype strains for three pathovars belonging to three *Xanthomonas* species. *Microbiology Resource Announcements* **7**, e00923-18. <https://doi.org/10.1128/MRA.00923-18>.

Ming D, Ye HZ, Schaad NW & Roth DA (1991) Selective recovery of *Xanthomonas* spp. from rice seed. *Phytopathology***81**, 1358-1363.

Naik SL, Nema KG, Kulkarni SN & Shrivastava PS (1973) Susceptibility of rice varieties to the attack of bacterial streak caused by *Xanthomonas translucens* Dowson f. sp. *oryzicola*(Fang *et al*.) Bradbury. *Indian Journal of Agricultural Sciences* **43**,590-594.

Ni Z, Cao Y, Jin X, Fu Z, Li J, Mo X, He Y, Tang J & Huang S (2021) Engineering resistance to bacterial blight and bacterial leaf streak in rice. *Rice (N Y)***14**, 38, 5pp. <https://doi.org/10.1186/s12284-021-00482-z>

Niño-Liu D, Ronald PC & Bogdanove AJ (2005) A simple method of mass inoculation of rice effective for both pathovars of *Xanthomonas oryzae*, and the construction of comparable sets of host cDNA libraries spanning early stages of bacterial leaf blight and bacterial leaf streak. *Journal of Phytopathology* **153,**500-504.

Niño-Liu DO, Ronald PC & Bogdanove AJ (2006) *Xanthomonas oryzae* pathovars: model pathogens of a model crop. *Molecular Plant Pathology* **7**, 303-324. <https://doi.org/10.1111/j.1364-3703.2006.00344.x>

Noh T-H, Shim H-K, Kang M-H, Park Y-J, Lee D-K, Lee BM, Tyagi K, Paik C-H & Lee G-H (2012) Rapid identification and validation of *Xanthomonas oryzae* pv. *oryzae* (*Xoo*) by using PCR-amplified phage integrase and transposase A gene*. Journal of the Korean Society of International Agriculture* **24**, 219-225.

Opina OS & Exconde OR (1971) Assessment of yield loss due to bacterial leaf streak of rice. *Philippine Phytopathology* **7**, 35-39.

Ou SH (1985) Rice Diseases. 2nd ed. Commonwealth Mycological Institute, Kew, Surrey, United Kingdom, pp 61-96.

Rao PS (1987) Across-season survival of *Xanthomonas campestris* pv. *oryzicola*, causal agent of bacterial leaf streak. *International Rice Research Newsletter* **12**, 2, 27.

Raymundo AK, Briones AM Jr, Ardales EY, Perez MT, Fernandez LC, Leach JF, Mew TW, Ynalvez MA, McLaren CG & Nelson RJ (1999) Analysis of DNA polymorphism and virulence in Philippine strains of *Xanthomonas oryzae* pv. *oryzicola*. *Plant Disease* **83**, 434-440.

Raymundo AK, Perez MT & Leach JE (2008) Detection and analysis of genetic polymorphism in *Xanthomonas oryzae* pv. *oryzicola* using repetitive sequence-based polymerase chain reaction. *Rice Genetics***III,** pp. 934-938 <https://doi.org/10.1142/9789812814289_0131>

Reddy PR (1983) Evidence of seed transmission of *Xanthomonas campestris*pv. *oryzae*. *Current Sciences***52**, 265–266.

Reddy RP & Nayak P (1975) Natural occurrence of bacterial leaf streak disease of rice on *Oryza perennis*. *Current Science* **44**, 131.

Reddy PR & Ou SH (1974) Differentiation of *Xanthomonas translucens* f.sp. *oryzicola* (Fang *et al*.) Bradbury, the leaf-streak pathogen, from *Xanthomonas oryzae* (Uyeda and Ishiyama) Dowson, the blight pathogen of rice, by enzymatic tests. *International Journal of Systematic Bacteriology* **24**, 450-452.

Reinking OA (1918) Philippines economic plant diseases. *Oryza sativa* L. rice bacterial leaf stripe. *Philippines Journal of Science*, Section A **13**, 225-226.

Rodriguez-R LM, Grajales A, Arrieta-Ortiz ML, Salazar C, Restrepo S & Bernal A (2012) Genomes-based phylogeny of the genus *Xanthomonas*. *BMC Microbiology* **12,**43, 14pp. <https://doi.org/10.1186/1471-2180-12-43>

Sakthivel N, Mortensen C & Mathur S (2001) Detection *of Xanthomonas oryzae* pv. *oryzae* in artificially inoculated and naturally infected rice seeds and plants by molecular techniques. *Applied Microbiology and Biotechnology* **56**, 435–441. <https://doi.org/10.1007/s002530100641>

Sapkota S, Mergoum M & Liu Z (2020) The translucens group of *Xanthomonas translucens*: Complicated and important pathogens causing bacterial leaf streak on cereals. *Molecular Plant Pathology* **21**, 291-302. <https://doi.org/10.1111/mpp.12909>

Sattayachiti W, Wanchana S, Arikit S, Nubankoh P, Patarapuwadol S, Vanavichit A, Darwell CT & Toojinda T (2020) Genome-wide association analysis identifies resistance loci for bacterial leaf streak resistance in rice (*Oryza sativa* L.). *Plants* **9**, 1673. <https://doi.org/10.3390/plants9121673>

Shekhawat GS, Srivastava DN & Rao YP (1969) Seed infection and transmission of bacterial leaf streak of rice. *Plant Disease Reporter* **53**, 115-116.Shekhawat GS, Srivastava DN & Rao YP (1972) Host specialization in bacterial leaf streak pathogen of rice (*Oryza sativa*). *Indian Journal of Agricultural Sciences* **42**, 11-15.

Shekhawat GS & Srivastava DN (1971) Control of bacterial leaf streak of rice (*Oryza sativa* L.). *Indian Journal of Agricultural Sciences***41**, 1098-1101.

Singh DV, Banerjee AK, Kishun R & Abidi AB (1980) Effect of bacterial leaf streak on the quantitative and qualitative characters of rice. *Indian Journal of Mycology and Plant Pathology* **10**, 67-68.

Swings J, Van den Mooter M, Vauterin L, Hoste B, Gillis M, Mew TW & Kersters K (1990) Reclassification of the causal agents of bacterial blight (*Xanthomonas campestris* pv. *oryzae*) and bacterial leaf streak (*Xanthomonas campestris* pv. *oryzicola*) of rice as pathovars of *Xanthomonas oryzae* (ex Ishiyama 1922) sp. nov., nom. rev. *International Journal of Systematic Bacteriology* **40**, 309-311.

Thianthavon T, Aesomnuk W, Pitaloka MK, Sattayachiti W, Sonsom Y, Nubankoh P, Malichan S, Riangwong K, Ruanjaichon V, Toojinda T, Wanchana S & Arikit S (2021) Identification and validation of a QTL for bacterial leaf streak resistance in rice (*Oryza sativa* L.) against Thai *Xoc* strains. *Genes* **12**, 1587. <https://doi.org/10.3390/genes12101587>

Tian Y, Zhao Y, Xu R, Liu F, Hu B & Walcott RR (2014) Simultaneous detection of *Xanthomonas oryzae* pv. *oryzae* and *X. oryzae* pv. *oryzicola* in rice seed using a padlock probe-based assay. *Phytopathology* **104**, 1130-1137.

Triplett LR, Hamilton JP, Buell CR, Tisserat NA, Verdier V, Zink F & Leach JE (2011) Genomic analysis of *Xanthomonas oryzae* isolates from rice grown in the United States reveals substantial divergence from known *X. oryzae* pathovars. *Applied & Environmental Microbiology* **77**, 3930-3937. <https://doi.org/10.1128/AEM.00028-11>

Triplett LR, Cohen SP, Heffelfinger C, Schmidt CL, Huerta AI, Tekete C, Verdier V, Bogdanove AJ & Leach JE (2016) A resistance locus in the American heirloom rice variety Carolina Gold Select is triggered by TAL effectors with diverse predicted targets and is effective against African strains of *Xanthomonas oryzae* pv. *oryzicola*. *Plant Journal* **87**, 472-483. <https://doi.org/10.1111/tpj.13212>

Tsuno K & Wakimoto S (1983) Ultrastructural investigations on the host parasite interactions in the rice leaf parenchymatous tissue infiltrated with bacterial leaf blight and leaf streak pathogens. *Annals of the Phytopathological Society of Japan* **49**, 659-669.

USDA (2013) Recovery Plan for *Xanthomonas oryzae* causing bacterial blight and bacterial leaf streak of rice. 22 pp. <https://www.ars.usda.gov/ARSUserFiles/opmp/Rice%20Bacterial%20Blight%20and%20Streak%20Recovery%20Plan%20Final.pdf>

Uyeda E & Ishiyama S (1928) In: S. Ishiyama (ed.), Bacterial leaf-blight of the rice plant. *Proceedings Third Pan-Pacific Scientific Congress*, Tokyo, 0ct.-Nov. 1926, **2**,2112.

Vauterin L, Hoste B, Kersters K & Swings J (1995) Reclassification of *Xanthomonas*. *International Journal of Systematic Bacteriology* **45**, 472-489.

Vera Cruz CMV, Gossele F, Kersters K, Segers P, Van den Mooter M, Swings J & Ley J de (1984) Differentiation between *Xanthomonas campestris* pv. *oryzae*, *Xanthomonas campestris* pv. *oryzicola* and the bacterial 'brown blotch' pathogen on rice by numerical analysis of phenotypic features and protein gel electropherograms. *Journal of General Microbiology* **130**, 2983-2999. <https://doi.org/10.1099/00221287-130-11-2983>

Wang SX, Ma GZ, Wei LF & Ji GH (2010) [Virulence differentiation of strains *of Xanthomonas oryzae*pv. *oryzicola* in southwest of China]. *Journal of Hunan Agricultural University* **36**, 188-191 (in Chinese).

Wonni I Detemmerman L, Dao S, Ouedraogo L, Soungalo S, Koita O, Szurek B, Koebnik R, Triplett L, Cottyn B & Verdier V (2011) Genetic diversity of *Xanthomonas oryzae* pv. *oryzicol*a from West Africa. *Phytopathology* **101**, S193-S193.

Wonni I, Cottyn B, Detemmerman L, Dao S, Ouedraogo L, Sarra S, Tekete C, Poussier S, Corral R, Triplett L, Koita O, Koebnik R, Leach J, Szurek B, Maes M & Verdier V (2014) Analysis of *Xanthomonas oryzae* pv. *oryzicola*population in Mali and Burkina Faso reveals a high level of genetic and pathogenic diversity. *Phytopathology* **104**, 520-531.

Xie GL & Mew TW (1998) A leaf inoculation method for detection of *Xanthomonas oryzae* pv. *oryzicola* from rice seed. *Plant Disease* **82**, 1007-1011.

Xie GL, Sun SY, Wang GJ, Zhu XD, Chen JA, Ye YH, Feng ZM & Liang MX (1990) Studies on rice seed inspection of *Xanthomonas campestris* pv. *oryzicola*. 1. Immunoradiometric assay. *Chinese Journal of Rice Science* **4,** 127-132

Xie GL, Wang HR, Sun SY & Wang GJ (1991) Studies on rice seed inspection of bacterial leaf streak of rice: II. Rice leaf in vitro method. *Chinese Journal of Rice Science* **5**, 121-126

Xie X, Chen Z, Guan H, Zheng Y, Zhang J, Qin M & Wu W (2020) Transcriptome analysis of xa5-mediated resistance to bacterial leaf streak in rice (*Oryza sativa* L.).*Scientific Reports* **10,**19439. <https://doi.org/10.1038/s41598-020-74515-w>

Xu GW & Gonzalez CF (1991) Plasmid, genomic, and bacteriocin diversity in U.S. strains of *Xanthomonas campestris* pv. *oryzae*. *Phytopathology* **81**, 628-631.

Xu Y, Zhu X-F, Zhou M-G, Kuang J, Zhang Y, Shang Y & Wang J-X (2010) Status of streptomycin resistance development in *Xanthomonas oryzae* pv. *oryzae* and *Xanthomonas oryzae* pv. *oryzicola* in China and their resistance characters. *Journal of Phytopathology* **158**, 601-608. <https://doi.org/10.1111/j.1439-0434.2009.01657.x>

Yang B & Bogdanove A (2013) Inoculation and virulence assay for bacterial blight and bacterial leaf streak of rice. *Methods in Molecular Biology* **956,**249-255. Yang, B, ed. Springer. <https://doi.org/10.1007/978-1-62703-194-3_18>

Yang J, Wang X, Wang Y-F, Liu Q, Wang Y-T, Zhang J-H, Wei L-F & Ji G-H (2020) Pathotypes differentiation of *Xanthomonas oryzae* pv. *oryzicola* and identification of rice varieties resistant to bacterial leaf streak in Yunnan Province. *Acta Phytopathologica Sinica* **50**, 218-227.

Young JM, Park DC, Shearman HM & Fargier E (2008) A multilocus sequence analysis of the genus *Xanthomonas*. *Systematic & Applied Microbiology* **31**, 366–377.

Zhang RS, Liu YF, Luo CP, Wang XY, Liu YZ, Qiao JQ, Yu JJ & Chen ZY (2012) *Bacillus amyloliquefaciens* Lx-11, a potential biocontrol agent against rice bacterial leaf streak. *Journal of Plant Pathology* **94**, 609-619.

Zhao B, Ardales EY, Raymundo A, Bai J, Trick HN, Leach JE & Hulbert SH (2004) The *avr*Rxo1 gene from the rice pathogen *Xanthomonas oryzae*pv. *oryzicola*confers a nonhost defense reaction on maize with resistance gene *Rxo1*. *Molecular Plant-Microbe Interactions* **17**, 771-779.

Zhao B, Lin X, Poland J, Trick H, Leach J & Hulbert S (2005) A maize resistance gene functions against bacterial streak disease in rice. *Proceedings National Academy of Sciences, USA*, **102**, 15383–15388

Zhao WJ, Zhu SF, Liao XL, Chen HY & Tan TW (2007) Detection of *Xanthomonas oryzae* pv. *oryzae* in seeds using a specific TaqMan probe. *Molecular Biotechnology* **35**, 119-127.

Zhao S, Poulin L, Rodriguez-R LM, Serna NF, Liu S-H, Wonni I, Szurek B, Verdier V, Leach JE He Y-Q, Feng J-X & Koebnik R (2012). Development of a variable number of tandem repeats typing scheme for the bacterial rice pathogen *Xanthomonas oryzae* pv. *oryzicola*. *Phytopathology* **102**,948-956. <https://doi.org/10.1094/PHYTO-04-12-0078-R>

**CABI and EFSA resources used when preparing this datasheet**

CABI (2022)*Xanthomonas oryzae* pv. *oryzae* (bacterial leaf blight of rice). <https://www.cabi.org/isc/datasheet/56956> [Accessed: April 2022].

CABI (2022)*Xanthomonas oryzae* pv. *oryzicola* (bacterial leaf streak of rice). <https://www.cabi.org/isc/datasheet/56977> [Accessed: April 2022].

EFSA (2018) EFSA Panel on Plant Health: Jeger M, Candresse T, Chatzivassiliou E, Dehnen-Schmutz K, Gilioli G, Gregoire J-C, Jaques Miret JA, MacLeod A, Navajas Navarro M, Niere B, Parnell S, Potting R, Rafoss T, Rossi V, Urek G, Van Bruggen A, Van der Werf W, West J, Winter S, Bragard C, Szurek B, Hollo G and Caffier D. Scientific Opinion on the pest categorisation *of Xanthomonas oryzae* pathovars *oryzae* and *oryzicola*. *EFSA Journal* **16**, 5109, 25 pp. <https://doi.org/10.2903/j.efsa.2018.5109>

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**Datasheet history**

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CABI/EPPO (1992/1997) Quarantine Pests for Europe (1st and 2nd edition). CABI, Wallingford (GB).

EPPO (1980) Data sheets on quarantine organisms No. 2, *Xanthomonas oryzae. EPPO Bulletin* **10**(1), 4 pp. <https://doi.org/10.1111/j.1365-2338.1980.tb02685.x>

