Crosstalk between Nutrient Signalling Pathways and Immune Responses in Rice

Beatriz Val-Torregrosa 1, Mireia Bundó 1,† and Blanca San Segundo 1,2,‡

1 Centre for Research in Agricultural Genomics (CRAG) CSIC-IRTA-UAB, Campus Universitat Autònoma de Barcelona (UAB), 08193 Bellaterra (Cerdanyola del Vallès), Spain; beatriz.val@cragenomica.es (B.V.-T);
2 Consejo Superior de Investigaciones Científicas (CSIC), 08016 Barcelona, Spain
* Correspondence: blanca.sansegundo@cragenomica.es or blanca.sansegundo@csic.es; Tel.: +34-935636600 † Current affiliation: Laboratori Agroalimentari, Departament d’Agricultura, Ramadera, Pesca i Alimentació (DARP), C/Cami de Mataró Núm. 1, 08348 Cabrils, Spain.

Abstract: Rice is a staple food for more than half of the global population. Rice production is, however, severely affected by biotic and abiotic stresses. Fertilisers and pesticides are widely used in rice farming to maintain optimal yield and to prevent losses caused by environmental stress. However, the indiscriminate use of agrochemicals has adverse effects on the environment and human health. Stress caused by nutrient excess or deficiency has an impact on plant disease resistance. The interference of plant responses induced by nutrient stress can result in a positive or negative impact on resistance to pathogen infection. In this review, we explore the effects of combined stresses in rice, focusing on nutrient stress, such as nitrogen and phosphorous supply, and infection by fungal pathogen Magnaporthe oryzae, the causal agent of rice blast disease. Crosstalk between nutrient stress-induced and pathogen-induced signalling pathways in rice is discussed, as well, with particular emphasis on regulatory microRNAs. Understanding the interconnected regulations between nutrient stress and disease resistance will lay a foundation for rationally optimising fertiliser and pesticide use in rice production.

Keywords: abiotic stress; biotic stress; Magnaporthe oryzae; microRNA (miRNA); Oryza sativa; pathogen; plant immunity; signalling crosstalk

1. Introduction

Rice (Oryza sativa L.) is one of the most important crops in the world and a staple food for more than half of the global population. China is the largest rice producer in the world, followed by India, Indonesia, and Bangladesh [1]. Rice is grown in a wide range of environments characterised by different climatic conditions, different cultivation systems such as paddy fields and rainfed uplands, and different irrigation or flooding conditions. To meet the forecasted demand of rice, posed by the increasing human population, rice production must significantly increase. This needs to be achieved within the declining arable land, due to urbanisation, which is often polluted by agrochemicals and industrialisation. Additionally, rice farmers are prone to face climate change and water scarcity factors.

Rice, a monocotyledonous plant species, originated around 9000 years ago in the Yangtze Valley (China) and it has a long history of natural selection and domestication [2]. It belongs to the genus Oryza which includes the two cultivated species, the Asian rice (Oryza sativa) and the African rice (Oryza glaberrima). O. sativa comprises japonica and indica subspecies [3–6]. O. rufipogon and O. nivara are considered to be the progenitors of the Asian cultivated rice, whereas O. glaberrima originated from its wild ancestor O. barthii [7,8]. O. sativa was the first crop species which was genome sequenced, and reference genomes are available for japonica and indica subspecies [9,10]. More recently, the 3 K Rice Genome Project sequenced more than 3000 rice accessions which provided a tremendous resource...
for rice research [11]. Rice has been adopted as the model plant for functional genomics in cereal species. Progress in agricultural research is needed for the development of necessary management strategies that can capitalise on enormous variety of rice production systems in very different climates. Sustaining rice production in adverse environmental conditions is a big challenge in the coming years. To overcome problems caused by biotic and abiotic stresses, new rice varieties with higher yield and improved tolerances need to be developed, taking advantage of all the resources and technologies currently available.

In this review, we present an overview of the major biotic stresses that affect rice production, with special emphasis on the rice blast disease. As pathogen infection causes major problems in rice cultivation, and nutrient stress (deficiency or excess) can be a factor in disease resistance, in this review we describe problems associated with inadequate supply of nutrients in resistance to pathogen, focusing on blast resistance. Our current knowledge on the molecular mechanisms of the plant immune system, including microRNA (miRNA)-mediated regulation of gene expression, is summarised. Lastly, we discuss the relevance of miRNAs in crosstalk between nutrient stress and immune signalling in plants, particularly in rice.

2. Biotic and Abiotic Stresses in Rice

Rice is prone to various types of stresses, both biotic and abiotic. Biotic stresses, caused by fungi, bacteria, viruses, insect pests, or nematodes, represent a serious threat to sustainable rice production. To reduce losses caused by pathogen infections and to obtain maximum yields, high rates of pesticides and fertilisers continue to be applied in rice farming. This has been harming to the environment and human health. Furthermore, climate change is promoting the northerly movement of pests, which means that temperate rice agriculture should be concerned about tropical pests and diseases.

Abiotic stresses, such as drought, salinity, cold, high temperature, submergence, nutrient deficiencies, and heavy metal toxicity are also factors negatively affecting rice growth and productivity worldwide. As most of the global rice supply originates from flooded ecosystems, rice is one of the most affected crops by drought stress. High salinity is also a major stress factor that can seriously affect rice growth, development, and yield. Indeed, rice is rated as the most salt-sensitive cereal crop. Nutrient stress can be also a major cause of decline in rice production. Nutrient stress refers to deficiencies of essential plant nutrients as well as excess of nutrients caused by over-fertilisation. Under nutrient stress conditions, the plant must respond dynamically to maintain nutrient levels in the optimal range for normal growth and development. Physiological disorders, also called physiological diseases, are quite common in rice grown under poor soil conditions [12].

It is also true that, in nature, plants are simultaneously exposed to varying combinations of biotic and abiotic stresses. However, most studies to determine the effects of environmental stress have been performed on plants exposed to an individual stress. So far, there is limited research focusing on the impact of combined abiotic and biotic stresses in rice. Crosstalk between plant responses induced by one stress may result in positive or negative impact over the other stress [13].

2.1. Biotic Stress

One of the most important pathogens affecting rice is the fungus *Magnaporthe oryzae*. This fungus can infect the leaves, leaf collars, necks, and panicles of rice plants and it causes rice blast. *M. oryzae* is an ascomycete fungus with a hemibiotrophic lifestyle that involves initial proliferation inside living host cells before switching to a destructive necrotrophic mode [14,15]. The blast disease cycle starts when a conidium reaches the host tissue and develops a germ tube, which grows on the leaf surface before differentiating into the dome-shaped appressorium (Figure 1). A penetration peg develops from the appressorium to enter into the epidermal cells, where it expands to become an infective hypha. During the biotrophic stage, the hyphae move from cell to cell through plasmodesmata [16]. At this stage of the infection process, the fungus redirects nutrients from living plant cells
to facilitate nutrient acquisition from the host tissue (Figure 1) [15,17]. Several days after infection, the fungus switches to a necrotrophic lifestyle and blast lesions appear on the leaf surface, typically diamond-shaped with a grey or white centre and brown or reddish-brown border (Figure 1). The fungus sporulates profusely in these lesions, thus, facilitating the disease to spread rapidly to adjacent rice plants. Neck rot and panicle blast are particularly devastating diseases that cause major yield losses in severe epidemics. Furthermore, M. oryzae is capable of infecting different crops apart from rice, such as barley or wheat, which highlights its incredible ability to infect monocotyledonous plant species. The M. oryzae genome has been sequenced, and functional genomic approaches have led to the identification of hundreds of genes involved in its pathogenesis. M. oryzae has become a model fungus for studying host-pathogen interactions, at both cell biology and molecular levels [18].

Rice blast is a classical gene-for-gene interaction system, and numerous blast resistance (R) genes are identified in rice [19,20]. Furthermore, numerous avirulence (AVR) genes have been genetically and functionally described in M. oryzae [21]. However, traditional resistance conferred by R genes often breaks down in a few years due to the high variability and fast evolving fungal populations. Therefore, breeders usually prefer rice varieties that contain multiple R genes and have broad-spectrum blast resistance [22]. Chemical fungicides acting on the M. oryzae life cycle (e.g., inhibitors of melanin biosynthesis) are currently used for the control of the blast disease [23]. Chemical treatments aiming at the activation of plant defences, such as probenazole or spermidine, are also used for blast control [24,25]. The wide use of these chemicals might, however, provoke the development of resistance in the pathogen and/or cause side effects on non-target organisms and the environment. As there is a deep public concern about the side effects of chemical fungicides, it is critical to have effective methods to control the rice blast disease in an environmentally friendly way.

Figure 1. Colonisation of rice leaves by the hemibiotroph fungus Magnaporthe oryzae. The fungus first establishes a biotrophic interaction with the rice plant and later switches to a necrotrophic lifestyle. During the biotrophic phase, the fungus grows inside living host cells. The hyphae are encased by the extra-invasive hyphal membrane (EIHM) which is continuous with the rice plasma membrane. At this stage of the infection process, the fungus uses plasmodesmata for cell-to-cell movement [16,26], and absorbs nutrients from the plant living cells. The host vacuolar integrity in the first invaded rice cells is important for M. oryzae early stages of infection [27]. In the necrotrophic phase, the fungus colonises the host tissue and obtains nutrients from the host dead cells. Ap, appressorium; EIHM, extra invasive hyphal membrane; IH, invasive hyphae; Pd, plasmodesmata; PP, penetration peg; Sp, spores; Vac, vacuole. Black triangles represent plant nutrients.
Bakanae ("foolish seedling" in Japanese) disease, caused by one or more seed-borne *Fusarium* species, mainly *F. fujikuroi*, is a disease of increasing economic importance in major rice-producing areas. The fungus infects rice plants from the seedling stage to the mature stage, with severe infection of rice seeds [28]. Symptoms of the disease are poor seedling emergence, abnormal elongation of internodes in the seedlings due to the ability of *F. fujikuroi* to produce gibberellic acids (GAs), and grain contamination. The most common management practices to limit bakanae are based on thermal seed treatment (hot water immersion) and application of fungicides. On the other hand, very few QTLs (Quantitative Trait Loci) for resistance to bakanae have been mapped in the rice genome, and only a small number of rice varieties showing resistance to the disease are identified [29]. *Fusarium* species also produce mycotoxins in rice grains that represent serious food safety concerns in animal and human health [30].

Another economically significant rice disease affecting rice worldwide is rice sheath blight, which is caused by the fungus *Rhizoctonia solani*. This fungus is a necrotrophic soil fungus that infects rice plants by vegetative mycelia and sclerotia that is able to survive for long periods in the soil. Its dispersal is facilitated by water management practices and infected plant debris. Protection of rice from sheath blight relies on the use of chemical fungicides and bacterial biocontrol agents (e.g., commercial formulations of *Bacillus subtilis*) [31].

Bacterial blight and bacterial leaf streak, caused by *Xanthomonas oryzae pv. oryzae* and *X. oryzae pv. oryzicola*, respectively, are considered as major bacterial diseases in rice [32]. *X. oryzae* enters into the host through wounds or natural openings on the surface of the plant. Once inside, this pathogen moves across the plant through the xylem and colonises different host tissues. Other bacterial pathogens of rice are *Dickeya zeae* that causes bacterial foot rot [33] and *Burkholderia glumae* that causes bacterial panicle blight of rice [34]. Fungicides are being used for the control of bacterial diseases in rice.

Rice is also vulnerable to a number of diseases caused by viruses that occur especially in the tropics [35]. The distribution of viruses affecting rice is generally restricted to one of the continents in which rice is grown, e.g., Rice hoja blanca virus (RHBH) in South America and Rice Yellow Mottle Virus (RYMV) in Africa. Rice stripe virus (RSV) and Rice tungro viruses (RTVB and RTSV) are found in Asia. Most of the rice viruses are transmitted by insect vectors in a persistent or non-persistent manner.

Insect pests are also important biological constraints limiting rice yield. Stem borers are generally considered the most damaging insect pest of rice. In particular, the striped stem borer (SSB) (*Chilo suppressalis*) is a major constrain on rice production. Stem borers are difficult to control with insecticides, as once they are inside the stem, they are protected from chemical sprays. In order to be successful with the chemical control, repeated applications of appropriate insecticide are needed. The insecticidal Cry proteins produced by *Bacillus thuringiensis* have provided a tool for the control of a wide diversity of insect pests, including the rice SSB, and rice varieties protected from SSB damage through the expression of *Bt cry-endotoxin* genes (*Bt* rice) have been developed [36]. The public controversy over genetically modified (GM) crops is, however, limiting the cultivation of *Bt* rice varieties. Conventional breeding for *C. suppressalis* resistance has proven to be difficult due to the poor knowledge of the genetic basis of this trait. It is also worthy to mention the problem with the herbivore *Nilaparvata lugens*, also known as the brown planthopper [37].

The root knot nematode *Meloidogine graminicola* is the most serious nematode pest of rice. This nematode is well-adapted to several scenarios where rice is grown, thus causing severe losses of this crop. Among the symptoms, it is possible to find stunted growth, yellowing, and gall formation on the roots of rice plants, symptoms that are greatly dependent on the environmental conditions [38].

Clearly, integrative strategies are required for successful control of rice diseases, including the use of advanced breeding lines and cultivars, forecasting distribution of the disease and pathogen races, examining wild *Oryza* species for novel sources of resistance, as well as using appropriated disease and water management systems. Modern biotechnology
offers new opportunities for the development of rice varieties with enhanced disease resistance, which will be beneficial if rice yields are increased using sustainable strategies.

2.2. Plant Responses to Nutritional Stresses

Nutrients, macronutrients (nitrogen, phosphorus, potassium) and micronutrients (iron, copper, zinc, manganese, nickel, etc.) are taken from the soil and play a very important role in plant growth and development. Plants naturally evolved physiological mechanisms and adaptation strategies to cope with nutrient deprivation. Under limiting nutrient conditions, plants change the root system architecture by stimulating root branching and can also secrete small molecules into the soil to enhance the nutrient’s bioavailability. On a molecular level, nutrient limitation associates to the induction of genes encoding nutrient transporters to increase uptake, as well as genes involved in recycling or reallocation of nutrients to maintain nutrient levels in the optimal range for plant growth. Under nutrient excess, the plant responds through modification of nutrient acquisition programs, detoxification pathways, and sequestration into cellular compartments (e.g., vacuoles).

Nitrogen (N) is a component of important molecules, such as amino acids, proteins, or nucleic acids. The most available forms of N in the soil are nitrate and ammonium. Plants have developed different strategies to overcome low N in the soil, aiming to enhance N use efficiency through the regulation of N utilisation and acquisition. Alterations in root architecture and increased activity of influx transporters are the major determinants to increase N acquisition efficiency [39].

Phosphorus (P), like N, is a main macronutrient in plants. It is a component of nucleic acids and phospholipids, and also plays crucial roles in enzymatic reactions and signal transduction cascades (e.g., protein phosphorylation). P is an integral component of adenosine triphosphate (ATP), the universal cellular energy cofactor in biological processes. In plants, P is absorbed through the roots in the form of inorganic phosphate (Pi). Even though the overall content of Pi in the soil is generally high, its low bioavailability represents a limiting factor for plant growth in many agricultural ecosystems [40]. P deficiency is widespread in major rice ecosystems and is the major growth-limiting factor in acid upland soils where soil P-fixation capacity is often important. To explore the soil, the Pi-starved plants change their root architecture by increasing lateral roots and root hairs [41]. The most evident symptoms of Pi deficiency in plants are the reduction in plant growth and development of thinner leaves with darker appearance [42]. Pi-deprived plants experience important transcriptional reprogramming aiming to increase Pi acquisition in the root, translocation from the root to the shoot, and mobilisation of Pi stored in the vacuole, the so-called Phosphate Starvation Response (PSR) [43–45]. Another strategy developed by Pi-starved plants is an increase in RNAses and phosphatase activities, these enzymes being involved in metabolic Pi recycling [46,47]. Pi excess in the soil also has an effect on root development, as the plants show a reduction in primary root growth and less meristematic activity [48]. The excess of Pi also provokes alterations in the content of other nutrients such as iron or zinc [48]. Excessive Pi accumulation leads to toxicity at the tip of rice leaves (leaf tip necrosis), being more evident in older leaves.

Potassium (K) is another important inorganic nutrient for plant growth and production. It plays important roles in plant cell processes related with stomata closure, photosynthesis, turgor pressure, ion transport, or enzyme reactions as a cofactor. Although most soils are rich in K minerals, relatively little K⁺ is present in forms that are available to plants. When the plant senses the lack of this nutrient in the soil, it triggers the activation of channels and K transporters and, in some cases, is accompanied by the induction of other cellular processes which are also regulated by Ca²⁺, Reactive Oxygen Species (ROS), and hormones such as abscisic acid (ABA) [49].

Although required in smaller amounts than macronutrients (N, P, K), micronutrients are also essential for plant growth and play an important role in balanced crop nutrition. Iron (Fe), copper (Cu), zinc (Zn), manganese (Mn), molybdenum (Mo), and nickel (Ni) are metal micronutrients that function in a range of cell’s chemical reactions. Silicon (Si) and
boron (B) mainly contribute to the plant’s structure, while the major role of the non-metal chlorine (Cl) serves in osmotic balance.

Fe performs a number of important functions in plant cellular processes, and like other elements, it functions as a cofactor of many enzymes required for a wide range of biological functions. Oxidation-reduction reactions are most commonly associated with Fe-containing enzymes, such as catalases and peroxidases. Fe is a component of ferredoxin, which is indispensable for photosynthesis, and cytochromes, which are necessary for the electron transport system in mitochondria and chloroplasts. Symptoms of Fe deficiency include pronounced interveinal chlorosis in which photosynthetic tissues turn yellow due to the lack of chlorophyll. Although Fe is generally present at high quantities in soils, its bioavailability is limited in aerobic conditions where iron is predominantly found in the Fe$^{3+}$ form, mainly as a constituent of polymers with extremely low solubility [50]. Rice is, however, typically cultivated under anaerobic conditions (paddy fields) where abundant Fe$^{2+}$ is readily available to the plant, especially in soils with a low pH in which Fe$^{3+}$ is reduced to the more soluble ferrous ion Fe$^{2+}$. Under such conditions, unlike other graminaceous plants, the absorption of Fe$^{2+}$ by the rice roots might cause severe Fe toxicity. Excessive Fe accumulation in rice tissues causes overproduction of ROS, which, in turn, causes oxidative damage to macromolecules (nucleic acids, lipids, proteins, etc.) and damage to cellular structures [51]. Rice responds to Fe toxicity by means of different adaptive mechanisms. Firstly, rice can retain Fe in the root tissues and decrease Fe translocation from roots to shoots to avoid excessive Fe accumulation in leaves. Secondly, the Fe$^{2+}$ ions, once in the cells, might associate with proteins (e.g., ferritins) or can be stored in cellular compartments (vacuoles). Under Fe excess, the rice plant activates the expression of genes associated with protection against oxidative stress [51]. High iron content in the soil also impairs the absorption of Cu in rice.

Cu serves primarily as a cofactor of metabolic enzymes and protein complexes in the electron transport chain. Cu is also an integral member of plastocyanin, which actively participates in the electron transport of chloroplast grana during photosynthesis. On the other hand, zinc is the cofactor for enzymes involved in N metabolism, and Zn deficiency declines anaerobic root metabolism and lessens the capacity of rice seedlings to cope with anaerobic soil conditions. Rice plants are more susceptible to Zn deficiency in early growth phases. On the other hand, B helps in the biosynthesis of cell walls, and thereby participates in cell division and elongation processes. B is critical for pollen tube elongation and B deficiency results in minimal pollen viability and lower grain yield [52].

Finally, rice is particularly susceptible to heavy metals contamination, in particular, cadmium (Cd) and arsenic (As) coming from contaminated soil, river, or groundwater. Upon exposure to heavy metals, plants experience oxidative stress that leads to cellular damage and disturbance of cellular ionic homeostasis. These toxic metals accumulate in rice grains and are significant risks for human and animal health. Water management practices significantly influence the Cd and As concentration in rice grains [53].

In the following sections, we present an introduction to the principles of plant immune responses. As nutrient stress can have an impact on resistance to pathogen infection in plants, we discuss interactions between nutrient responses and immunity in rice. Understanding how rice plants respond to pathogen infection in heterogeneous and fluctuating environments is crucial for the development of efficient means of disease control.

3. Plant Innate Immunity

Plants have evolved multiple mechanisms to sense and respond to pathogen infection. Resistance to pathogen infection relies on numerous changes at the biochemical, physiological, and molecular level, largely coordinated by a massive and fast reprogramming of the host transcriptome. Upon contact with the plant, conserved molecular signatures derived from microbes, known as pathogen-associated molecular patterns (PAMPs; or microbe-associated molecular patterns, MAMPs) are recognised by Pattern Recognition Receptors (PRRs) localised in the plasma membrane [13,54–57]. Examples of MAMPs include
bacterial flagellin, lipopolysaccharide, EF-Tu, lipoproteins, peptidoglycans, and fungal chitin [58]. Plant PRRs fall into the receptor-like kinase (RLK) or receptor-like protein (RLP) families [59]. PAMP recognition triggers the activation of a general defence response referred to as PAMP-triggered immunity (PTI), in which multiple signal transduction pathways operate. Plants can also detect their own molecules such as oligogalacturonides released from plant cell walls as a consequence of physical damage caused by the pathogen infection, referred to as Damage-Associated Molecular Patterns (DAMPs), that are also able to initiate innate immune responses [58,60]. Extracellular ATP and endogenous elicitor peptides can also function as danger signals for the activation of the innate plant immune response [61]. Some pathogens are able to produce effector proteins that are delivered into the host cell to suppress PTI responses. Therefore, the pathogens carrying these effectors can cause diseases in their host plants [56]. During plant–pathogen coevolution, however, plants developed resistance (R) proteins, encoded by resistance (R) genes that recognise microbial effectors (or host proteins modified by effectors). This recognition triggers a rapid and robust host defence response, known as Effector-Triggered Immunity (ETI) [54,56]. Most R proteins belong to the nucleotide-binding site (NB)/leucine-rich-repeats (LRR) receptor family (NLR) [19]. ETI is often accompanied by a form of programmed cell death at the site of pathogen infection, the hypersensitive response (HR), which restricts the spread of infection. Accumulating evidence also supports that PTI and ETI share a number of signalling components [62]. Timing and intensity of the host responses, however, differ between PTI and ETI [63].

One of the early events in the plant response to pathogen infection is the increase in cytoplasmic Ca\(^{2+}\) for the activation of kinase cascades in which calcium-dependent protein kinases (CDPKs) and mitogen-activated protein kinases (MAPKs) participate [64]. Pathogen-induced defence responses include the production of ROS, such as hydrogen peroxide (H\(_2\)O\(_2\)), superoxide radicals (O\(^{2-}\)), and hydroxyl radicals (·OH) [65]. ROS are reactive molecules with multiple functions in plant defence reactions to pathogen infection. In particular, H\(_2\)O\(_2\) possess antimicrobial activity and contributes to cell wall reinforcement (e.g., lignification and oxidative cross-linking of cell-wall polymers) [66]. ROS also function as signalling molecules for the induction of defence responses. Another free radical that is produced in the plant during infection is nitric oxide (NO). NO seems to work together with ROS radicals in triggering cell death upon pathogen challenge and is also strongly related with Ca\(^{2+}\) and ABA signalling for regulation of the stomata closure [67].

Plants also produce proteins and secondary metabolites that can be toxic to phytopathogens. Perhaps the best-known examples of defensive metabolites are phytoalexins, which are synthesised de novo in response to infection. Major phytoalexins accumulating in rice are the diterpene phytoalexins momilactones, phytocassenes, and oryzalexins, and the flavanone phytoalexin sakuranetin [68,69]. However, Brassicaceae plant species accumulate camalexin, a tryptophan-derived phytoalexin with antifungal and antibacterial activity against phytopathogens [70]. Indole glucosinolates are also effective to limit colonisation by necrotrophic fungi in plants [71]. The accumulation of Pathogenesis-Related (PR) proteins is also a ubiquitous response of plants to pathogen infection, some of these proteins exhibiting antimicrobial activity [72].

Hormones play a crucial role in disease resistance by regulating plant immune responses [73]. Immune responses are coordinated mainly by the phytohormones salicylic acid (SA), ethylene (ET), jasmonic acid (JA and JA-derivatives), as well as by auxins and brassinosteroids. Plant hormones do not function independently, as synergistic and antagonistic interactions between hormone signalling pathways ultimately drive the fine-tuning of plant defence responses. Generally, the JA signalling pathway synergistically cross-talks with the ET signalling pathway, whereas the SA and JA pathways show negative crosstalk [73]. Hormone signalling networks in plant defence appear to be dependent on both the lifestyle of the pathogen in the host plant and the host plant itself. Whereas SA has been generally associated with plant defence against biotrophic pathogens, JA and ET are associated with protection against necrotrophic pathogens.
Historically, plant immune responses to fungal and bacterial pathogens have been considered to rely on the function of protein-coding genes, whereas RNA-based mechanisms largely regulate plant–virus interactions. Along with this, the transcriptional control of protein-coding defence genes has been studied extensively, but less attention has been paid to the post-transcriptional regulation of defence-related processes. Nowadays, the contribution of host endogenous small RNAs in the regulation of plant defence responses to infection by fungal and bacterial pathogens is well demonstrated [74–77]. Accordingly, the involvement of microRNAs (miRNAs), a class of small RNAs from plants, in post-transcriptional regulation of gene expression in the rice immune system is discussed below, with special emphasis on mechanisms involved in the rice response to infection by the blast fungus *M. oryzae*.

3.1. Regulatory Role of microRNAs in Rice Immunity

MiRNAs are short non-coding small RNAs that direct post-transcriptional gene silencing through cleavage or translational repression mechanisms [78,79]. In addition to miRNAs, plants possess another class of small RNAs, the small interfering RNAs (siRNAs) [80,81]. The two classes of small RNAs, miRNAs and siRNAs, are distinguished by their mode of biogenesis, function, and mechanisms of action. MiRNAs derive from long single-stranded RNA precursors with imperfect stem-loop structure (primary miRNAs, or pri-miRNAs) which are processed in a two-step process by RNAse III DICER-like proteins (DCL, typically DCL1) to produce a precursor miRNA (pre-miRNA) and then a double-stranded duplex, the miRNA-5p/miRNA-3p. The functional strand of the duplex is incorporated into an ARGONAUTE (AGO) protein in the RNA-induced silencing complex (RISC) where the miRNA recognises target transcripts by sequence complementarity [80]. Contrary to miRNAs, siRNAs derive from perfectly paired, long double-stranded RNA precursors that are produced by the activity of an RNA-dependent RNA polymerase (RDR).

According to their nucleotide sequence, miRNAs are categorised into different families, which comprise one or more members. The *miRBase* is a repository of *miRNA* sequences and annotations ([www.mirbase.org](http://www.mirbase.org)) (accessed on 3 June 2021) [84]. Currently, the *miRBase* contains 738 mature *rice* (*O. sativa*) miRNAs, following *Medicago truncatula* and *Glycine max* with 756 miRNAs in both species (*miRBase* release 22.1).

Plant miRNAs play a critical role in controlling diverse developmental processes and adaptation to environmental stress, both biotic and abiotic stresses [74,75,85–88]. Certain miRNAs have been shown to control traits of agronomic importance in rice, such as tillering, flowering, panicle and grain production [89,90]. Examples of miRNAs involved in tolerance to abiotic stress in rice are miR398 (drought, salt, and oxidative stresses), miR319 (cold stress), and miR393 (salinity), or miR399 (nutrient homeostasis) [85,91].

The role of miRNAs in disease resistance was described for the first time in the model plant *Arabidopsis thaliana* [92]. Here, perception of the *Pseudomonas syringae* flagellin peptide flg2 induces miR393 accumulation, and down regulation of auxin receptors, which in turn results in bacterial resistance. Since then, other miRNAs have been reported to play a key role in *Arabidopsis* immunity, such as miR160, miR167, miR393, miR396, miR398b, miR400, miR472, miR773, miR844, miR858, and miR1916, among others [75,87,93–96]. Increasing evidence supports that plant miRNAs play a role in PTI and ETI. Depending on the target gene that they regulate, miRNAs might function as positive or negative regulators of immune responses [97]. Most of our current understanding of miRNAs in plant immunity comes from studies on the interaction of *Arabidopsis* plants with the bacterial pathogen *P. syringae*. Limited information is available on miRNAs involved in disease resistance in other plant species and/or during interaction with other types of pathogens.

Even though a number of miRNAs have been shown to respond to pathogen infection in rice, the biological role of these pathogen-regulated miRNAs in rice immunity is not completely understood. Indeed, the biological role of only few miRNAs has been characterised
in the rice/\textit{M. oryzae} interaction (Figure 2). Some of these miRNAs function as positive regulators of immune responses (miR7695, miR160a, miR162a, miR398b, miR166k-166h and miR812w), whereas other miRNAs function as negative regulators of immune responses (miR156fh1, miR164a, miR167d, miR169a, miR319b, miR396, miR399, miR444b.2, miR439, and miR1873) \cite{77, 98-109}. To note, miR398 has been reported to function as a positive regulator of immune responses in rice against \textit{M. oryzae} while negatively regulating immune responses against \textit{P. syringae} in Arabidopsis \cite{110}.

![Figure 2. miRNAs with a known function in the response of rice plants to infection by the blast fungus \textit{M. oryzae}. The miRNA/target gene modules and the biological processes that they regulate are indicated. ACO, 1-aminocyclopropane-1-carboxylase oxidase3; ARF12, Auxin response factor; Chl, Chloroplast; CIPK, Calcineurin B-Like (CBL)-interacting serine–threonine protein kinase; CSD, Copper superoxide dismutase; DCL1, Dicer-like protein; EIN, Ethylene-insensitive protein 2; GRF, Growth regulating factors; LRR, Leucine-rich repeat; NAC, No apical meristem; NF-YA, Nuclear transcription factor Y subunit alpha; NRAMP, Natural resistance-associated macrophage protein; Nuc, Nucleus; PHO, Phosphate; PTs, Phosphate transporters; ROS, Reactive oxygen species; SPL, SQUAMOSA promoter-binding protein-like transcription factor; Sp, Spores; TCP, Teosinte branched/cycloidea/pcf; Vac, Vacuole.

In other studies, the activation of \textit{MIR166k-166h} encoding a polycistronic miRNA from rice was found to enhance resistance to infection by \textit{M. oryzae} and \textit{F. fujikuroi} in rice \cite{103}. The regulatory role of \textit{miR166k-166h} relies on the activity of \textit{miR166k-5p} (encoded by the \textit{miR166k-166h} precursor) on \textit{EIN2} (\textit{ETHYLENE INSENSITIVE 2}) transcripts, the \textit{EIN2} gene being a component of the ethylene signalling pathway in plants. Ethylene signalling is required for resistance against the blast fungus \textit{M. oryzae} in rice \cite{111}.

Regarding miR812, a new member of the miR812 family in rice, miR812w, involved in blast resistance was recently described \cite{99}. Whereas overexpression of miR812w increased blast resistance, CRISPR/Cas9-mediated \textit{MIR812w} editing enhances disease susceptibility. Of interest, miR812 originated from a miniature-inverted repeat element (MITE) of the Stowaway family of DNA transposons from rice, and the incorporation of cognate MITEs
into protein-coding genes would have created target sites for miR812w. Moreover, miR812w directs methylation in trans at its target genes, as well as in cis at the MIR812w locus [99].

Not only miRNAs themselves, but also components of the miRNA biogenesis machinery appear to play a role in blast resistance. Whereas silencing of OsDCL1 enhances resistance to the rice blast fungus, its activation compromises resistance against this fungus [112,113]. OsDCL1 activation represses pathogen-inducible rice defence responses, impairs diterpenoid phytoalexin production, and provokes important alterations in the population of rice miRNAs [113].

The functional characterisation of miRNAs, and components of the miRNA biogenesis and function machinery involved in disease resistance, also opens the possibility of using miRNAs as biomarkers in the management of disease resistance in rice breeding programs.

3.2. Impact of Nutrient Stress on Rice Immunity

Nutrient stress caused by excess or deficiency of nutrients might affect the plant’s response to pathogen infection, thus, determining the outcome of the interaction, resistance or susceptibility. The impact of nutrient stress on disease resistance is, however, difficult to predict, as different results are observed in the outcome of the interaction (resistance, susceptibility) depending on the identity of the interacting partners (host and pathogen) [114]. Although adaptation to nutrient stress and immunity are not independent processes, the molecular mechanisms involved in nutrient responses and innate immunity have been so far investigated separately from each other. To better understand how rice plants defend themselves during pathogen attack, it is essential to consider that they can be exposed to adverse environmental conditions, including improper nutrient supply and nutritional imbalances. Being a foliar pathogen, the fungus M. oryzae must acquire nutrients from the host, hence, the availability of host nutrients can be an important factor contributing to the establishment of the rice blast disease [17].

Plants can develop different nutritional-based defence mechanisms based on sequestration of essential nutrients away from the invading pathogen (withholding strategies), or localised accumulation of nutrients at the infection site that become toxic to the pathogen (i.e., hyper-accumulation of Fe for activation of an oxidative burst). The term “nutritional immunity” is used in human health as a non-specific host immune response and refers to mechanisms for withholding iron availability by the host that becomes inaccessible to the pathogen [115]. The term nutritional immunity can be also applied to restriction mechanisms of essential nutrients in plants to defend against pathogens. In plant/pathogen interactions, the host and the pathogen compete with each other for essential nutritional resources. This dynamic interaction is particularly relevant when plants are grown under nutrient limiting conditions. On the other hand, an increase in nutrient supply might create a more favourable environment for pathogen growth that would allow the pathogen to improve nutrient acquisition from the host plant, thus, promoting pathogenicity. However, the massive use of fertilisers might also have a negative impact on nutrient use efficiency in the plant and might be responsible for metabolic changes that jeopardise the plant defence responses. On the contrary, the accumulation of certain nutrients that can be toxic for the pathogen (e.g., iron) might protect the plant from infection. Figure 3 illustrates how interactions between nutrient and immune signalling pathways, particularly N, Pi, and Fe signalling, might lead to increased resistance or susceptibility to M. oryzae infection in rice.

It is generally assumed that N nutrition has an effect on disease incidence in different plant species [116]. During infection, pathogens require N sources, including NH$_4^+$ and NO$_3^-$, as well as amino acids. It has long been recognised that high N fertilisation enhances susceptibility to the blast fungus in rice plants [117]. It is also known that N supply has an impact on the two interacting partners: host and pathogen (Figure 3). From the perspective of the host plant, N availability has an effect on plant primary and secondary metabolism which, in turn, might affect host defence responses. While promoting plant growth, high N input might result in less formation of lignin and reduction in the thickness of the secondary cell wall, the plant’s physical barrier to pathogen infection [118]. From the perspective of
the pathogen, fungal pathogenicity can be affected by N availability. The nitrogen-induced susceptibility (NIS) to rice blast has been shown to be associated with the induction of rice genes implicated in N recycling and an increase in *M. oryzae* pathogenicity [119]. Contrary to what is observed in rice, N fertilisation reduces disease severity caused by *Verticillium* spp. in *Solanum* species, indicating that no generic model can describe the role of N in a given interaction [114].

![Figure 3. Crosstalk between nutrient signalling pathways and immune response. Nutrients are essential for plant growth and development. They are obtained from the soil through the plant root system. However, insufficient or excessive amounts of certain nutrients can affect disease resistance in plants. In rice, high nitrogen (N) supply not only promotes *M. oryzae* growth but also induces the expression of fungal effector genes, thus making the plant more susceptible to infection [117]. Growing rice plants under high phosphate (Pi) conditions leads to an increase in leaf Pi content, which is associated with a weaker activation of defence responses and susceptibility against the rice blast fungus [98]. Ferroptosis, an iron (Fe)- and ROS-dependent cell death process, contributes to arrest *M. oryzae* infection in incompatible rice/*M. oryzae* interactions [120]. Thus, iron levels must be strictly controlled in the host plant to maintain the subtle balance between growth and defence.](image)

Regarding Pi nutrition, emerging evidence supports the existence of crosstalk between Pi starvation signalling machinery and immune responses in plants [121,122]. However, our knowledge of interactions between plant adaptive mechanisms to Pi excess and immunity is still limited. Only recently, it was described that high Pi fertilisation compromises the expression of immune responses and enhances susceptibility to infection by *M. oryzae* in rice plants [98] (Figure 3). In other studies, overexpression of the *OsPT8* phosphate transporter, and subsequent increase in Pi content, was reported to enhance susceptibility to *M. oryzae* infection [123]. Therefore, over-use of fertilisers in rice fields might have unintended consequences in rice production by facilitating blast infection, while contributing to water and soil pollution.

Rice plants are also able to establish symbiotic associations with arbuscular mycorrhizal (AM) fungi [124]. Root colonisation by AM fungi improves the uptake of mineral nutrients in the host plant, mainly P and N [125–128]. Even though root colonisation by AM fungi is reduced in rice plants grown under permanent flooded conditions, evidence supports that, once the AM symbiosis is established, its functional capacities are not af-
fected by flooding [129]. During colonisation by AM fungi, the rice root system architecture undergoes drastic morphological and transcriptional alterations to accommodate fungal structures [130,131]. AM colonisation is, however, unevenly distributed within the rice root system: the crown roots are weakly colonised, the large lateral roots are strongly colonised, and fine lateral roots are not colonised [132,133].

Moreover, there is an important genetic variation within rice varieties for root colonisation by AM fungi [133]. Beneficial effects of root colonisation by AM fungi include growth stimulation, increased yield, and blast resistance, but differences are observed among cultivars in terms of blast resistance [134–137]. Recent studies in leaves of mycorrhizal rice plants revealed that the AM symbiosis is accompanied by a coordinated systemic regulation in the expression of genes involved in hormone signalling (JA, ET) and lipid signalling [136]. Then, systemic transcriptional responses occurring in leaves of mycorrhizal rice plants might be potentially important for defence against foliar pathogens. The AM symbiosis might as well contribute to the sustainability of rice production, an aspect that remains poorly explored.

As previously mentioned, Fe is an essential element for plants and a co-factor for a variety of proteins involved in redox reactions. Plants have developed strategies to mobilise Fe from the rhizosphere, referred to as Strategy I and Strategy II [138]. Typically, dicotyledonous species (such as Arabidopsis) and non-gramineous monocotyledonous species use the strategy I, in which protons are released into the rhizosphere to acidify the soil and to increase ferric iron (Fe$^{3+}$) solubility. Iron is then reduced to ferrous form (Fe$^{2+}$) by a ferric reductase oxidase (FRO) and enters into the root cells by the activity of the Fe$^{2+}$ transporters (IRTs). A different strategy, Strategy II, is used by graminaceous monocotyledonous plants, in which phytosiderophores are released into the rhizosphere and then reabsorption of the metal-phytosiderophore complex occurs by root membrane transporter proteins. Rice employs both strategies for iron uptake [138,139]. When in excess, however, iron becomes toxic to the plant cells due to the formation of ROS, which might cause oxidative damage to macromolecules and cellular structures, eventually leading to death. Therefore, iron homeostasis needs to be tightly controlled in the plant to avoid toxicity.

In plant–pathogen interactions, iron homeostasis must be carefully regulated to allow normal plant growth while providing a way to arrest pathogen growth. The molecular mechanisms underlying Fe homeostasis in the host plant during pathogen infection are, however, poorly understood. Evidence supports a relationship between iron content and expression of immune responses in plants, including rice [108,140–142]. In M. oryzae-infected rice leaves, Fe has been found to accumulate at the sites of attempted penetration by the fungus (appressoria), and in cells in the vicinity of the infection site, further supporting that Fe accumulation and redistribution in the host tissue might be important for blast resistance [108]. Iron- and ROS-dependent signalling cascades are known to be involved in the ferroptotic cell death pathway, a form of non-apoptotic, iron-dependent cell death that rice plants use to prevent M. oryzae infection [120] (Figure 3). Studies in wheat plants revealed that infection with Blumeria graminis f. sp. tritici elicits Fe accumulation in the apoplast, which in turn mediates apoplastic oxidative burst and induction of defence gene expression [143]. It has been proposed that disturbance of Fe homeostasis is sufficient to prime the plant immune system for enhanced defence [144].

Cu is an important element in a number of pesticides in agriculture because it interferes with numerous enzymatic reactions and blocks respiratory activity in bacterial, fungal, and oomycete pathogens. Accordingly, Cu is widely used in crop protection because of its antimicrobial activity against phytopathogens. In rice, Cu application is very effective to suppress bacterial blight, caused by Xanthomonas oryzae pv oryzae (Xoo). However, several bacterial pathogens have developed resistance to the metal, and Cu-resistant pathogenic bacteria have been reported because of the wide application of Cu-containing pesticides in agriculture worldwide [145].
In summary, evidence gathered over years indicates that nutrient stress has an impact on disease resistance in rice. Our understanding of the molecular mechanisms underlying the innate ability of rice plants to cope with pathogen infection in heterogeneous environments is, however, far from complete. Further investigation on these processes in different rice/pathogen/environment interactions would provide a solid foundation for the development of novel strategies for rationally optimising fertiliser and pesticide use in rice production and protection against diseases.

3.3. miRNAs in the Crosstalk between Nutrient Stress and Immune Signalling

Understanding how plants cope with pathogens under nutrient stress conditions and how nutrient signalling pathways interact with immune signalling is an issue of great importance in crop protection. Clearly, a dynamic regulation of gene expression is required for optimal plant responses to nutrient conditions that are diverse in time and space, as well as during pathogen infection. Being versatile regulators of gene expression, miRNAs are well suited for proper integration of these inputs.

Several miRNAs have been shown to play a key role in regulating nutrient homeostasis in plants [146,147]. Some of them are specifically responsive to a specific nutrient condition, but others can be regulated by different nutrient conditions. For instance, miR167, miR169, miR393, and miR827 have been described as nitrogen-responsive miRNAs, whereas miR395 (targeting ATP sulfurylase genes) functions in sulphur assimilation [146,147]. Nutrient-responsive miRNAs might then play a role in controlling nutrient homeostasis as well as crosstalk among nutrient stress responses. So far, most research on miRNAs involved in nutrient homeostasis focused on plants under nutrient-limiting conditions, and less is known about their implication in plants that encounter excess of nutrients in the environment. It is also true that, although a large number of miRNAs have been shown to be responsive to nutrient stress, how such alterations affect nutrient homeostasis remains to be determined.

The implication of miR399 in the phosphate starvation response of plants is well documented [43,91,148–150]. Most of our knowledge on this issue comes from studies in the model plant *Arabidopsis thaliana*, where Pi starvation is accompanied by an increase in miR399 accumulation in roots which, in turn, down-regulates the expression of *PHO2* (PHOSPHATE 2). *PHO2* encodes an ubiquitin E2 conjugating enzyme that mediates degradation of the Pi transporters. In this way, miR399 accumulation relieves negative post-transcriptional control of Pi transporters for an increase in Pi uptake. Moreover, miR399 moves from roots to shoots and serves as a signalling molecule in regulating Pi homeostasis during Pi deprivation at the whole-plant level. This allows the plant to increase uptake and root-to-shoot allocation of Pi in Arabidopsis plants under Pi-limiting conditions [91,148,151]. The miR399/PHO2 module has a conserved function in controlling phosphate homeostasis in Arabidopsis and rice plants.

Another miRNA controlling Pi homeostasis is miR827, which appears to function via different pathways in Arabidopsis and rice plants. In Arabidopsis, miR827 targets NITROGEN LIMITATION ADAPTATION (NLA) encoding an ubiquitin E3 ligase that mediates degradation of the plasma membrane-localised Pi transporter PHT1 [152,153], whereas in rice, miR827 targets two different OsSPX-MFS genes (*OsSPX-MFS1* and *OsSPX-MFS2*) that localise at the tonoplast [154,155]. It has been proposed that in rice, miR827 functions in Pi compartmentalisation and Pi storage [154–157]. For additional information on Pi starvation responses in rice, we refer to [43,44,158].

Recent studies have demonstrated that the miR399 might function as a potential integrator of Pi starvation responses and immune-defence responses in rice plants. Thus, transgenic rice plants overexpressing miR399 showed increased phosphate content, weaker induction of defence-related genes, and enhanced susceptibility to *M. oryzae*, which also correlates with the susceptibility phenotype observed in rice plants grown in Pi excess conditions [98]. Other components of the Pi starvation signalling also regulate the expression of genes involved in immune responses [121,122].
On the other hand, miR398 functions in plant stress responses by regulating the expression of genes encoding copper superoxide dismutase (CSD1, CSD2) and Cu chaperones for SOD1 involved in protection against oxidative stress in plants [159,160]. Overexpression in rice of miR398 results in up-regulation of defence-related genes, which in turn enhances resistance to infection by *M. oryzae* [110].

In other studies, a rice miRNA, miR7695, has been reported to regulate iron homeostasis and rice immunity. MiR7695 negatively regulates the expression of *OsNramp6* (Natural resistance-associated macrophage protein 6), encoding an iron and manganese transporter from rice [102,140]. miR7695 regulates not only Fe content, but also defence responses during infection of rice plants with *M. oryzae*. Both MIR7695 activation and growing rice plants under high Fe supply is accompanied by a superinduction of *PR* and diterpenoid phytoalexin biosynthesis genes which, in turn, results in enhanced resistance to infection by *M. oryzae* [108].

Thus, from what we have learned so far, distinct miRNAs can play a role in the crosstalk between nutrient stress signalling and immune signalling. MiRNAs might potentially work as molecular switches between these two intimately connected processes. As miRNAs are responsible for fine regulation of gene expression, rather than on-off regulations, these riboregulators are well suited to function in the crosstalk between stress-induced signalling pathways. At present, however, our knowledge on interconnected regulations between nutrient signalling and immune signalling, in which miRNAs participate, is still in its infancy.

4. Conclusions and Perspectives

With the predicted global demand for rice, in an ever-growing population, there is an increasing pressure to develop rice cultivars with improved stress responses and increased yield. The potential impact of climate change on rice production and the emergence of new diseases in a changing climate are also major issues that need to be addressed to ensure global food security. Towards this end, understanding the molecular mechanisms underlying disease resistance under nutrient stress conditions is essential for the development of effective strategies aiming at combating pathogen infection, an aspect that remains poorly explored. Fertilisers containing plant nutrients are commonly used to maintain optimal yield in rice cultivation. Ironically, excess nitrogen and phosphate fertilisation has been shown to increase susceptibility to infection with the blast fungus *M. oryzae* in rice [98,117]. Thus, the indiscriminate use of fertilisers might have adverse effects on the rice plant by increasing the likelihood of blast disease. There is the need to develop eco-friendly technologies to maintain rice productivity while reducing inputs of agrochemicals, fertilisers and pesticides, also in response to legislative and public pressure.

Regarding miRNA function, this is a revolutionary field of research, expanding in recent years. Distinct miRNAs are known to function in crosstalk between nutrient stress and immune responses. However, to fully exploit the potential of miRNAs as regulators of rice immune responses, a better knowledge of interconnected regulations between nutrient homeostasis and immune responses is needed. Here, it is worth mentioning that because of the small size of *MIRNA* genes, the functional analysis of *MIR* genes has historically been a challenge (e.g., lack of T-DNA insertional knock-out mutants). Alternative approaches have been used to repress endogenous miRNA activity which includes miRNA target mimicry, transcriptional gene silencing of miRNA promoters, and artificial miRNA-directed silencing of miRNA precursors. New technologies for precise, efficient gene targeting or genome editing such as the CRISPR/Cas9 system has been developed for genome editing and targeted mutagenesis of plant miRNAs, which have been successfully used for the functional analysis of rice miRNAs [99,161,162]. Using this methodology will enable us to expand our knowledge on the regulatory role of miRNAs in adaptive processes to environmental stress in rice plants, including pathogen resistance under nutrient stress conditions. Unravelling miRNA-mediated mechanisms governing pathogen resistance and nutrient homeostasis will be useful in designing novel strategies.
to improve not only disease resistance but also the nutritional status of rice plants with exclusion of use and release of agrochemicals. Since diseases caused by pathogens are major threats to rice production, high rates of pesticides and fertilisers are currently applied in rice production. As a consequence, environmental problems have arisen in rice growing areas due to excessive use of pesticides and fertilisers. The combination of novel molecular and biotechnological tools, screening technologies, advanced breeding, and cultural practices, also considering the plant microbiome, should become the main goal in agriculture in the 21st century.

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