

Academy of Scientific Research & Technology



Journal of Genetic Engineering and Biotechnology

Vol. 16 No 1, 2018

F

B

G

Available online at www.sciencedirect.com

ScienceDirect



10.51.01.5



Journal of Genetic Engineering and Biotechnology

Editor-in-Chief: Mahmoud Saker <u>View Editorial Board</u> <u>Submit Your Paper</u> <u>Open Access</u> <u>View Articles</u> <u>Guide for Authors</u> <u>Abstracting/ Indexing</u> <u>Track Your Paper</u> Journal Metrics

- <u>CiteScore</u>: 1.58 i
- Source Normalized Impact per Paper (SNIP): 0.993 i
- SCImago Journal Rank (SJR): 0.375 i
- <u>View More on Journal Insights</u>

Your Research Data

- <u>Share your research data</u>
- Visualize your data

Society Links



اكاديمية البحث العلمي والتكنولوجيا Academy of Scientific Research and Technology

Related Links

- <u>Author Stats</u> i
- Researcher Academy
- <u>Author Services</u>
- <u>Try out personalized alert features</u>

Production and Hosting by Elsevier B.V. on behalf of Academy of Scientific Research and Technology

Peer Review under the responsibility of the National Research Center

Journal of Genetic Engineering and Biotechnology is devoted to rapid publication of full-length research papers that lead to significant contribution in advancing knowledge in genetic engineering and biotechnology and provide novel perspectives in this research area. JGEB includes all major themes related to genetic engineering and recombinant DNA.

The areas of interest of JGEB include but not restricted to:

- Plant genetics,
- Animal genetics,
- Bacterial enzymes,
- Agricultural Biotechnology,
- Biochemistry,
- Biophysics,
- Bioinformatics,
- Environmental Biotechnology,
- Industrial Biotechnology,
- Microbial biotechnology,
- Medical Biotechnology,
- Bioenergy,
- Biosafety,
- Biosecurity,
- Bioethics,
- GMO<mark>S</mark>,
- Genomics,
- Proteomics

JGEB accepts research papers, short communications, reviews and mini-reviews.

Journal of Genetic Engineering and Biotechnology

Editorial Board

Editor-in-Chief Mahmoud Saker Academy of Scientifc Research and Technology, National Research Center, Egypt

Deputy Editor Sameh Soror Dept. of Genetic & Biotechnology, Helwan University, Cairo, Egypt

Associate Editors Yasser Abdel Fattah City of Scientific Research and Technological Applications (SRTA-City), Egypt Nabila Abdel Maksoud National Research Center, Egypt **Desouky** Abd-El-Haleem City of Scientific Research and Technological Applications (SRTA-City), Egypt Adel Abolesoud The Agricultural Research Center, Egypt Ahlam Abou Mossallam National Research Center, Egypt Mai Allam National Research Center, Egypt Shireen Assem Agricultural Genetic Engineering Research Institute, Egypt Abdel Fattah Badr Helwan University, Egypt Mahmoud Bahgat National Research Center, Egypt Danila Carbonara Università degli Studi di Pavia, Italy Sherif El-Khamisy Zewail City of Science and Technology, Egypt Ahmed Gaballa Cornell University, USA Yehia Zakaria Gad National Research Center, Egypt **Reda Gafaar** Tanta University, Egypt Hassan Ghareeb National Research Center, Egypt **Anil Grover** University of Delhi South Campus, India

Nada Hamza Commission of Genetic Engineering & Biotechnology, Sudan **Moemen Hanafy** National Research Center, Egypt **Harrison Hughes** Colorado State University, USA Yehia Mahumoud Tanta University, Egypt Hanan Malkawi Hamdan Bin Mohammed Smart University, United Arab Emirates **Osman El-Mahdy Osman** National Research Center, Egypt Mohamed Rady National Research Center, Egypt Hala Ragab National Research Center, Egypt **Ewald Schnug** Julius Kühn-Institut, Germany Mohamed Shaba Colorado State University, USA **Michel Smith** Kansas State University, USA

Volume 16, Issue 2 Pages 239-776 (December 2018)

Microbial/industrial Biotechnology

Review article Open access <u>Quorum sensing intervened bacterial signaling: Pursuit of its cognizance and repression</u> Kayeen Vadakkan, Abbas Alam Choudhury, Ramya Gunasekaran, Janarthanam Hemapriya, Selvaraj Vijayanand Pages 239-252

Short communicationOpen access <u>Phylogenetic diversity and biotechnological potentials of marine bacteria from continental slope</u> <u>of eastern Arabian Sea</u> Arakkaveettil Kabeer Farha, Thasneem TR, Aswathy Purushothaman, Jaseetha Abdul Salam, Abdulla Mohamed Hatha Pages 253-258

Research article Open access Valorisation of chicken feathers for xanthan gum production using *Xanthomonas campestris* <u>MO-03</u> Murat Ozdal, Esabi Basaran Kurbanoglu Pages 259-263

Research article Open access <u>Biolytic extraction of poly(3-hydroxybutyrate) from *Bacillus megaterium* Ti3 using the lytic enzyme of *Streptomyces albus* Tia1 Neetu Israni, Surabhi Thapa, Srividya Shivakumar Pages 265-271</u>

Research article Open access <u>Purification and characterization of alkaline soda-bleach stable protease from *Bacillus* sp. APP-<u>07 isolated from Laundromat soil</u> I.K. Shaikh, P.P. Dixit, T.M. Shaikh Pages 273-279</u>

Research article Open access Improvement of cellulose degradation by cloning of endo-β-1, 3-1, 4 glucanase (*bgls*) gene from *Bacillus subtilis* BTN7A strain

Wafaa K. Hegazy, Mohamed S. Abdel-Salam, Azhar A. Hussain, Hoda H. Abo-Ghalia, Safa S. Hafez

Pages 281-285

Research articleOpen access Antibacterial activity of soil bacteria isolated from Kochi, India and their molecular identification Davis Gislin, Dorairaj Sudarsanam, Gnanaprakasam Antony Raj, Kathirvelu Baskar

Pages 287-294

Research article Open access

Purification and characterization of alkaline protease with novel properties from *Bacillus cereus* strain S8

B.K.M Lakshmi, D. Muni Kumar, K.P.J Hemalatha Pages 295-304

Research article Open access

Enhancement of nematicidal potential through cloning and expression of *chitinase* gene from *Bacillus subtilis* subsp. *Subtilis* BTN7A strain

Mohamed S. Abdel-Salam, Hoda H. Ameen, Abdallah S.M. Kassab, Ahmed E.A. Mahgoob, Usama S. Elkelany

Pages 305-310

Research article Open access <u>Biodegradation of feather waste by keratinase produced from newly isolated *Bacillus* <u>licheniformis ALW1</u> Azza M. Abdel-Fattah, Mamdouh S. El-Gamal, Siham A. Ismail, Mohamed A. Emran, Amal M. Hashem</u>

Pages 311-318

Research article Open access <u>Study on the potential of cold-active lipases from psychrotrophic fungi for detergent formulation</u> Sanjay Sahay, Deepak Chouhan Pages 319-325

Research article Open access <u>Optimization of novel halophilic lipase production by *Fusarium solani* strain NFCCL 4084 using palm oil mill effluent Kiptoo Geoffry, Rajeshwara N. Achur Pages 327-334</u>

Research article Open access

<u>Cloning and expression of MPT83 gene from *Mycobacterium tuberculosis* in *E. coli* BL21 as vaccine candidate of tuberculosis: A preliminary study</u>

Ahyar Ahmad, Rosana Agus, Muh. Nasrum Massi, Rosdiana Natzir, ... Masugi Maruyama Pages 335-340

Research article Open access <u>Immobilization of thermostable exo-inulinase from mutant thermophilic Aspergillus tamarii-U4</u> <u>using kaolin clay and its application in inulin hydrolysis</u> Emmanuel O. Garuba, Abiodun, A. Onilude Pages 341-346

Research article Open access

High level expression and purification of recombinant flounder growth hormone in E. coli

Tae-Jin Choi, Temesgen Tola Geletu Pages 347-355

Research article Open access

Screening of potential probiotic lactic acid bacteria and production of amylase and its partial purification

Tallapragada Padmavathi, Rayavarapu Bhargavi, Purushothama Rao Priyanka, Naige Ranganath Niranjan, Pogakul Veerabhadrappa Pavitra

Pages 357-362

Research article Open access <u>Partial purification and characterization of exoinulinase produced from *Bacillus* sp. R. Ramapriya, A. Thirumurugan, T. Sathishkumar, D.R. Manimaran Pages 363-367</u>

Research article Open access

Effect of vitamins and cell constructions on the activity of microbial fuel cell battery Dena Z. Khater, K.M. El-Khatib, Rabeay Y.A. Hassan Pages 369-373

Research article Open access <u>Decolorization of Textile Reactive Dyes by Bacterial Monoculture and Consortium Screened</u> <u>from Textile Dyeing Effluent</u> Md. Ekramul Karim, Kartik Dhar, Md. Towhid Hossain Pages 375-380

Research article Open access

Optimization of quorum quenching mediated bacterial attenuation of *Solanum torvum* root extract by response surface modelling through Box-Behnken approach Kayeen Vadakkan, Selvaraj Vijayanand, Abbas Alam Choudhury, Ramya Gunasekaran, Janarthanam Hemapriya Pages 381-386

Research article Open access

Isolation and characterization of *Bacillus* sp. strain BC01 from soil displaying potent antagonistic activity against plant and fish pathogenic fungi and bacteria Md Javed Foysal, Asura Khanam Lisa Pages 387-392

Research article Open access

Expression of Leptospira membrane proteins Signal Peptidase (SP) and Leptospira Endostatin like A (Len A) in BL-21(DE3) is toxic to the host cells

Padikara K. Satheeshkumar, Prasannan V. Anu, Mohmed I. Junaida, Madathiparambil G. Madanan, ... Perumana R. Sudhakaran Pages 393-398

Research article Open access

<u>Scenedesmus obliquus:</u> Antioxidant and antiviral activity of proteins hydrolyzed by three enzymes

Abd El-Moneim M.R. Afify, Gamal S. El Baroty, Farouk K. El Baz, Hanaa H. Abd El Baky, Soha A. Murad

Pages 399-408

Research article Open access

Statistical optimization of crude oil bio-degradation by a local marine bacterium isolate

Pseudomonas sp. sp48

Soha Farag, Nadia A. Soliman, Yasser R. Abdel-Fattah Pages 409-420

Research article Open access

Influence of bioprocess variables on the production of extracellular chitinase under submerged fermentation by *Streptomyces pratensis* strain KLSL55

A. Shivalee, K. Lingappa, Divatar Mahesh Pages 421-426

Review article Open access

Recent advances in stem cells therapy: A focus on cancer, Parkinson's and Alzheimer's Dalia Fleifel, Mai Atef Rahmoon, Abdelrahman AlOkda, Mostafa Nasr, ... Sherif F. El-Khamisy Pages 427-432

Research article Open access

In vitro differentiation of human multilineage differentiating stress-enduring (Muse) cells into insulin producing cells

Ali M. Fouad, Mahmoud M. Gabr, Elsayed K. Abdelhady, Mahmoud M. Zakaria, ... Ayman F. Refaie

Pages 433-440

Research articleOpen access <u>Development and evaluation of latex agglutination test coating with recombinant antigen,</u> <u>LipL32 for serodiagnosis of human leptospirosis</u> Kotchakorn Thongsukkaeng, Rerngwit Boonyom

Pages 441-446

Research article Open access <u>PNME – A gene-gene parallel network module extraction method</u> Bikash Jaiswal, Kumar Utkarsh, D.K. Bhattacharyya Pages 447-457

Research article Open access

Expression, purification and biological characterisation of recombinant human irisin (12.5 kDa) Kalpana Panati, Venkata Ramireddy Narala, Vydyanath R. Narasimha, Madhavi Derangula, ... Suneetha Yeguvapalli

Pages 459-466

Research article Open access

Increased level of B cell differentiation factor in systemic lupus erythematosus patients Hala Zaki Raslan, Hiba Sibaii, Salwa Refat El- Zayat, Hagar Hassan, Mahitab El- Kassaby Pages 467-471

Research article Open access

Healthcare-associated (HA) and community-associated (CA) methicillin resistant <u>Staphylococcus aureus</u> (MRSA) in Bangladesh – Source, diagnosis and treatment Md. Anowar Khasru Parvez, Rabeya Nahar Ferdous, Md. Shahedur Rahman, Sohidul Islam Pages 473-478

Research article Open access

Assessment of Ki-67 as a potential biomarker in patients with breast cancer Halla Mohamed Ragab, Nervana Samy, Mie Afify, Nabila Abd El Maksoud, HebatAllah Mohamed Shaaban Pages 479-484

Research article Open access <u>Kolaviron and selenium reduce hydrogen peroxide-induced alterations of the inflammatory</u> <u>response</u> Tebekeme Okoko Pages 485-490

Animal Biotechnology

Research article Open access

Feline panleukopenia viral infection in cats: Application of some molecular methods used for its diagnosis

Romane A. Awad, Wagdy K.B. Khalil, Ashraf G. Attallah Pages 491-497

Research article Open access

Buffalo species identification and delineation using genetic barcoding markers Amal Ahmed Mohamed Hassan, Esraa Aly Balabel, Hanaa Abdel Sadek Oraby, Samy Anwar Darwish Pages 499-505

Research article Open access Detection of myostatin gene MSTN in some goat breeds (*Capra hircus*) Y.A. Dowidar, M.A. El-Sayed, Aly M. Elrefy, Hytham E. Shoura Pages 507-512

Research article Open access

Five BoLA-DRB3 genotypes detected in Egyptian buffalo infected with Foot and Mouth disease virus serotype O

Othman E. Othman, Muhammad G. Khodary, Ayman H. El-Deeb, Hussein A. Hussein Pages 513-518

Research article Open access <u>Cytogenetic effects of silver and gold nanoparticles on Allium cepa roots</u> Priyanka Debnath, Arghadip Mondal, Amita Hajra, Chittaranjan Das, Naba Kumar Mondal Pages 519-526

Research article Open access <u>Synthesis of silver nanoparticles by *Bacillus clausii* and computational profiling of nitrate reductase enzyme involved in production Koel Mukherjee, Rashmi Gupta, Gourav Kumar, Sarita Kumari, ... Padmini Padmanabhan Pages 527-536</u>

Plant Biotechnology

Review article Open access <u>Transgenic approaches for genetic improvement in groundnut (Arachis hypogaea L.) against</u> <u>major biotic and abiotic stress factors</u> Saikat Gantait, Suvendu Mondal Pages 537-544

Review article Open access *In vitro* biotechnological advancements in Malabar nut (*Adhatoda vasica* Nees): Achievements, <u>status and prospects</u> Saikat Gantait, Jitendriya Panigrahi Pages 545-552

Review article Open access <u>Elevated carotenoids in staple crops: The biosynthesis, challenges and measures for target</u> <u>delivery</u> Adebanjo Ayobamidele Badejo Pages 553-562

Review article Open access <u>In vitro culture, transformation and genetic fidelity of Milk Thistle</u> M.R. Rady, M.M. Saker, M.A. Matter Pages 563-572

Research article Open access <u>Cloning, transformation and expression of cell cycle-associated protein kinase OsWeel in indica</u> <u>rice (Oryza sativa L.)</u> Frengky H.H. Prasetyo, Bambang Sugiharto, Netty Ermawati Pages 573-579

Research article Open access

Optimization of indole acetic acid production by isolated bacteria from *Stevia rebaudiana* rhizosphere and its effects on plant growth

Sheela Chandra, Kazim Askari, Madhumita Kumari Pages 581-586

Research article Open access

Plant regeneration, developmental pattern and genetic fidelity of somatic embryogenesis derived *Musa* spp.

Natarajan Nandhakumar, Krish Kumar, Duraialagaraja Sudhakar, K. Soorianathasundaram Pages 587-598

Research article Open access <u>Population structure, morphological and genetic diversity within and among melon (*Cucumis* <u>melo L.) landraces in Iran</u> Masoud Maleki, Abdolali Shojaeiyan, Sajad Rashidi Monfared Pages 599-606</u>

Research article Open access <u>Influence of cold pretreatment on shoot regeneration from callus in date palm (*Phoenix* <u>dactylifera L.) cv. 'Barhee'</u> Ahmed Madi Waheed Al-Mayahi, Abdulminam Hussien Ali, Hussein J. Shareef</u>

Pages 607-612

Research article Open access

Screening of plant growth promoting traits in heavy metals resistant bacteria: Prospects in phytoremediation

N. Tirry, N. Tahri Joutey, H. Sayel, A. Kouchou, ... N. El Ghachtouli Pages 613-619

Research article Open access

<u>Phytochemical analysis, antioxidant and antimicrobial activity of wild and *in vitro* derived plants of *Ceropegia thwaitesii* Hook – An endemic species from Western Ghats, India
S. Muthukrishnan, T. Senthil Kumar, A. Gangaprasad, F. Maggi, M.V. Rao
Pages 621-630
</u>

Research article Open access

Molecular diversity of internal transcribed spacer among the monoconidial isolates of *Magnaporthe oryzae* isolated from rice in Southern Karnataka, India D. Jagadeesh, M.K. Prasanna Kumar, R. Chandrakanth, N.S. Devaki Pages 631-638

Research articleOpen access <u>Production of biomass and flavonoid of *Gynura procumbens* (Lour.) Merr shoots culture in temporary immersion system</u>

Ayu Dewi Pramita, Alfinda Novi Kristanti, Sugiharto, Edy Setiti Wida Utami, Yosephine Sri Wulan Manuhara Pages 639-643

Research article Open access

<u>Callus mediated shoot organogenesis and regeneration of cytologically stable plants of</u> <u>Ledebouria revoluta</u>: An ethnomedicinal plant with promising antimicrobial potency Sk Moquammel Haque, Avijit Chakraborty, Biswajit Ghosh Pages 645-651

Research article Open access

Evaluation of the alleviative role of Chlorella vulgaris and Spirulina platensis extract against ovarian dysfunctions induced by monosodium glutamate in mice Sekena H Abdel-Aziem, Heba A.M. Abd El-Kader, Faten M. Ibrahim, Hafiza A Sharaf, Aida I. El makawy Pages 653-660

Research article Open access

Assessment of genetic diversity in *Salvadora persica* L. based on inter simple sequence repeat (ISSR) genetic marker

Mohammad Asadi Monfared, Davood Samsampour, Gholam Reza Sharifi-Sirchi, Fatemeh Sadeghi

Pages 661-667

Research article Open access

Micropropagation protocol for Antigonon leptopus an important ornamental and medicinal plant Zenna Fawzia Ghareeb, Lobna S. Taha

Pages 669-<mark>675</mark>

Research article Open access

Total phenolic and flavonoid contents and antioxidant activity of ginger (*Zingiber officinale* Rosc.) rhizome, callus and callus treated with some elicitors

Ammar Mohammed Ahmed Ali, Mawahib ElAmin Mohamed El-Nour, Sakina Mohamed Yagi Pages 677-682

Research article Open access

Physiological and molecular studies on the effect of gamma radiation in fenugreek (*Trigonella foenum-graecum* L.) plants

Rania Samy Hanafy, Samia Ageeb Akladious Pages 683-692

Research article Open access <u>Rice straw fermentation by Schizophyllum commune ARC-11 to produce high level of xylanase</u> <u>for its application in pre-bleaching</u> Archana Gautam, Amit Kumar, Amit Kumar Bharti, Dharm Dutt Pages 693-701

Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579



Contents lists available at ScienceDirect

Journal of Genetic Engineering and Biotechnology

journal homepage: www.elsevier.com/locate/jgeb



Cloning, transformation and expression of cell cycle-associated protein kinase *OsWee1* in indica rice (*Oryza sativa L*.)



Frengky H.H. Prasetyo^a, Bambang Sugiharto^b, Netty Ermawati^{c,*}

^a Graduate School of Biotechnology Department, Jember University, JL. Kalimantan 37 Kampus Tegalboto, Jember 68121, Indonesia ^b Center for Development of Advanced Sciences and Technology, and Department of Biology, Faculty of Mathematic and Natural Sciences, Jember University, JL. Kalimantan 37 Kampus Tegalboto, Jember 68121, Indonesia

^c Department of Agricultural Production, and Central Laboratory for Biosciences, State Polytechnic of Jember, JL. Mastrip PO Box 164, Jember 68120, Indonesia

ARTICLE INFO

Article history: Received 1 June 2018 Received in revised form 23 September 2018 Accepted 1 October 2018 Available online 7 December 2018

Keywords: Agrobacterium Cell cycle Gene transformation OsWee1 Protein kinase Transgenic rice

ABSTRACT

The development process of seed in plants is a cycle of cells which occur gradually and regularly. One of the genes involved in controling this stage is the *Wee1* gene. *Wee1* encode protein kinase which plays an important role in phosphorylation, inactivation of cyclin-dependent kinase 1 (CDK1)-cyclin (CYC) and inhibiting cell division at mitotic phase. The Overexpression of *Wee1* leads to delaying entry into mitotic phase, resulting in enlargement of cell size due to suppression of cell division. Accordingly, the cloning and overexpressing of *Wee1* in rice plant is important aim of this research in achieving better quantity and quality of future rice. The main objective of this present study is to cloning and generate transgenic rice plants overexpressing of *Wee1* gene. *Wee1* was isolated from cDNA of indica rice (*Oryza sativa*), called *OsWee1*. The full length of *OsWee1* was 1239 bp in size and successfully inserted into plant expression vector pRI101ON. Seven-day-old rice seedlings were prepared for transformation of *OsWee1* gene using Agrobacterium-mediated transformation method. Four positive transgenic lines were identified through the presence of kanamycin resistance gene (*nptII*) using genomic PCR analysis. Southern blot analysis result provides evidence that four independent rice transformation is needed in order to obtain stable expression of *OsWee1*.

© 2018 Production and hosting by Elsevier B.V. on behalf of Academy of Scientific Research & Technology. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-

nd/4.0/).

1. Introduction

One of the parameters to increase the yield of rice is the seed sizes, which regulated by the seed development processes after pollination. The process is a part of cell cycle which occur gradually and regularly during its life cycle. The cycle is divided into four phases, the mitotic (M) phase which includes mitosis and cytokinesis, G1 (first gap) phase, S (synthesis) phase, and G2 (second gap) phase. The cell develops larger in the G1 phase, then in S phase, cell replicates the chromosome inside the cell. Furthermore, the cell will continue its growth in G2 phase, and divided. The daughter cells can repeat the cycle [1]. The cell cycle progression is controlled at distinct checkpoints which major checkpoints are synthesis phase (G1-S checkpoint), mitosis (G2-M checkpoint) and the spindle checkpoint [2].

Peer review under responsibility of National Research Center, Egypt. * Corresponding author.

E-mail address: netty@polije.ac.id (N. Ermawati).

In eukaryotes, the cell cycle is controlled by family of conserved cyclin-dependent protein kinases (CDKs). The activity of CDK fluctuates regularly during the cell cycle, triggering important processes [3]. Phosphorylation and dephosphorylation of the CDK catalytic subunit, threonine 14 and tyrosine 15, are able to regulate and determine the timing of G2 and mitosis [4]. Previous study reported that phosphorylation of CDKs at tyrosine 15 in *Schizosaccharomyces pombe* is mediated by *Wee1*, which causes a delay in mitosis by phosphorylating the M-phase promoting factor on tyrosine 15 [5].

Wee1 is a gene encoding protein kinase located in the nucleus. The expression of this gene in plants is strongly induced by DNA damage which can be caused by radiation, ionization, chemicals and other stresses [6,7]. When DNA is damaged, ataxia-telangiectasia mutated (ATM) or ATM- and Rad3-related (ATR) kinases will be expressed depending on the genotoxic type of stress. Futhermore, the ATM and ATR signals will phosphorylate and activated Chk1 and further phosphorylates the *Wee1*. Activation of Chk1 caused cell cycle delay in G2-M phase by increasing *Wee1* regulation and decreasing regulation of phosphatase (Cdc25) which

https://doi.org/10.1016/j.jgeb.2018.10.003

¹⁶⁸⁷⁻¹⁵⁷X/© 2018 Production and hosting by Elsevier B.V. on behalf of Academy of Scientific Research & Technology. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579

controls tyrosine-15 phosphorylation inhibitors on cyclindependent kinase (Cdc25) resulting in G2 phase arrest [8–10].

In fission yeast (S. pombe), loss of Wee1 activity causes unsufficient growth cells early enter to mitosis phase and cytokinesis, therefore it causes cells to produce two abnormally small daughter cells [5,11]. However, increasing expression of Wee1 causes delayed entry into mitosis and increase in cell size, this indicates that the activity levels of Wee1 plays a role in ensuring the entry time of the mitotic phase and having strong effect on cell size [12]. Sun et al. [13] reported that expression of *ZmWee1* was observed in endosperm tissue at 15 day after pollination, where this gene shows its role in endoreduplication in the endosperm and supposes to be a potential regulator of seed development. The similar result was reported in Wee1 tomato [14] and AtWee1 from Arabidopsis [15], that the expression levels of *Wee1* gene was found higher in the generative organs such as seed, fruit and flower compared to that the vegetative organs. In previous study, the expression of rice *OsWee1* was almost found in all the tissues; roots, stem, tiller, flowers, leaves and seeds. The highly expression of rice OsWee1 was found in 5 day after pollination of the seeds [16]. These results revealed that besides having an important role in seed developments, Wee1 has also influence in the growth and developments of plants.

Considering the important role of *Wee1* in the development of seed, cloning and transformation of *OsWee1* was conducted in order to have understanding of the superior potential of *OsWee1* overexpressing in rice. In this study, we present results of *OsWee1* cloning and overexpression of this gene in rice.

2. Materials and methods

2.1. Plant materials

The mature seeds of indica rice (cv. Mekongga) were used in this research. Dehulled seeds were sterilized with 70% ethanol for 2 min followed by 5.25% sodium hypochlorite for 10 min and then washing with sterile distilled water for 3–5 times. The sterilized seeds were placed on MS basal salt media (Table 1) pH 5.8, supplemented with 3% (w/v) sucrose, 100 mg/L L-glutamine, 0.25% phytagel, and cultured under continuous light at ±22 °C within a period of 7 days.

2.2. Plasmid construct

Cloning of *OsWee1* consists of 2 steps, first step was cloning *OsWee1* into pGEMT easy vector (Promega), and the second was

Table 1

MS	basal	salt	media	content	(In	mg/L	media).	

Components	mg/L
NH ₄ NO ₃	1650.0
KNO ₃	1900.0
MgSO ₄ ·7H ₂ O	370.0
MnSO ₄ ·4H ₂ O	22.3
ZnSO ₄	10.6
CuSO ₄ ·5H ₂ O	0.025
CaCl ₂ ·H ₂ O	440.0
KI	0.83
CoCl ₂ ·6H ₂ O	0.025
KH ₂ PO ₄	170.0
H_3BO_3	6.2
Na2MoO4·2H2O	0.25
FeSO ₄ ·7H ₂ O	27.85
Na2EDTA-2H2O	37.25
Nicotinic acid	0.5
Pyridoxine HCl	0.5
Thiamine HCl	1.0
Glycine	2.0

cloned into plant expression vector pRI101ON vector (TaKaRa) (Fig. 1A). The amplification fragment of OsWee1 was conducted from DNA recombinant pGEMT: OsWee1 which obtained from previous study [16] and deposited in GeneBank under Accession no. KX758541. PCR analysis were performed using the following a set of primer contain NdeI and BamHI sites (Table 2) overhang to ensure compatibility with pRI101ON vector. The fragment of OsWee1 was amplified as follows PCR Core Kit (Roche) manufacture's procedure, initial denaturation at 94 °C for 2 min, each with 25 cycles of denaturation at 94 °C for 15 sec, annealing at 57 °C for 20 sec, extension at 72 °C for 2 min, and a final extension at 72 °C for 7 min. Fragment obtained was then purified using GeneAll Expintm Combo GP and quantified using nanodrop (NanoVue Plus spectrophotometer, BioLab). The DNA fragment of OsWee1 was ligated into pRI1010N and the recombinant pRI1010N. OsWee1 was then transformed into E. coli XL10 gold competent cells through heat shock method [15].

2.3. Flanking analysis of OsWee1

A recombinant of *pRI1010N*. *OsWee1* was amplified and confirm the correct size by digestion using *HindIII*, *EcoRI*, *Nde1* and *BamHI* restriction enzymes (NEBr Inc.). The flanking frame of *OsWee1* in pRI1010N was checked and analyzed using Sanger dideoxy sequencing technology (The 1st BASE, Malaysia). The sequence was then analyzed using BLAST (www.ncbi.nlm.nih.gov/blast).

2.4. Transformation into Agrobacterium

DNA recombinant of *pRI1010N*. OsWee1 was transferred into Agrobacterium cells by heat shock method [17]. Aliquot of $100 \,\mu$ L freshly prepared competent cells and $1 \,\mu$ L of DNA recombinant were mixed, keep on ice for 5 min and chilled into liquid nitrogen. Heat shock was immediately conducted by heated at 42 °C in waterbath for 90 sec. Added 1 ml of YEP medium (10 g/L yeast extract, 10 g/L peptone, 5 g/L NaCl₂) and gently shaked at 28 °C for 1 h to allow bacteria and harboring DNA replication. Cultured bacteria was then collected by centrifuge at 5000 rpm for 3 min, colected pellet was disolved and spread on YEP agar medium supplement with 50 mg/L kanamycin, 100 mg/L rifampicin and 12.5 mg/L gentamicin. The bacteria cells were incubated at 28 °C for 2 d and the colony grown was identified using PCR with several specific primers (Table 2).

2.5. Agrobacterium-mediated transformation and putative rice transformants selection

Agrobacterium harboring the binary The construct pRI1010N: OsWee1 was inoculated in YEP medium supplemented with 100 mg/L rifampicin, 12.5 mg/L gentamicin and 50 mg/L kanamycin, then incubate at 28 °C by gently shake at 110 rpm for 48 h. The growth of Agrobacterium was checked for its optical density by spectrophotometer ($OD_{600nm} = 0.3$). Seven-day-old rice seedlings were soaked in the Agrobacterium suspension for 20 min. To reduce the growth of excessive bacteria, the infected seedlings were dried using sterilized filter papers for 5 min. The infected seedlings were grown into co-cultivation medium (MS basal salt (Table 1), 3% sucrose, 100 mg/L acetosyringone, 0.3% phytagel; pH 5.2) and incubated for 2 d in the dark condition. The co-cultivated rice seedlings were thoroughly washed with 500 mg/L cefotaxime followed by sterilized water for three times. Furthermore, the seedlings were cultured on selection medium (MS basal salt, 3% sucrose, 0.25% phytagel, 50 mg/L kanamycin and 250 mg/L cefotaxime; pH 5.8) under a 16/8-h (day/night) light cycle at 22 °C and periodically sub-cultured every 4 weeks into fresh media. The screening

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579

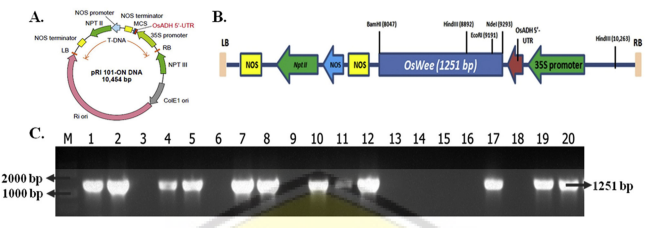


Fig. 1. Cloning OsWee1 into expression vector pRI101ON. A. A Map of pRI101ON vector; B. Schematic representation of pRI101ON. OsWee1 construct; C. Colony PCR analysis of OsWee1 amplified using 146-FNdel and 147-RBamHI primers.

Table 2

List of primers used in this study and corresponding sequences.

Primer name	Sequence (5'-3')	GC content (%)
OsWee-F	ATGGCACTTGGAATTAGTTGTGGTC	44.0
OsWee-R	TTATCGTGGCAAACCAACTGAGG	47.8
FndeI	GCCATATGGCACTTGGAATTAGTTGT	42.3
RBamHI	GCGGATCCTTATCGTGGCAAACCAA	52.0
CaMV-F	GAAGACGTTCCAACCACG	55.6
RV-F	CAGGAAACAGCTATGACC	50.0
nptII-F	GTCATCTCACCTTGCTCCTGCC	59.1
nptII-R	GTCGCTTGGTCGGTCATTTCG	57.1
OsActin-F	TCCATCTTGGCATCTCTCAG	50.0
OsActin-R	GTACCCGCATCAGGCATC	61.1

putative transformants were conducted by screening of rice explants on the selection media containing 50 mg/L kanamycin.

2.6. Molecular analysis of putative transformed plants by PCR and Southern blotting

Genomic DNA was isolated from 4 g of leaves of *wildtype* and putative transformants rice [18] with minor modification. To confirm the presence of transgene, the putative transformants and non-transformant rice samples were analyzed by PCR analysis using nptII-F and nptII-R primers (Tabel 2). About 50–100 ng of total genomic DNA from independent putative transformant lines and non-transformant were mixed with 50 μ l of reaction mix (KAPA Taq Extra HotStart ReadyMix) and subjected for PCR analysis under pre-denaturation condition at 95 °C for 3 min, followed by 30 cycles at 95 °C for 30 sec, 58 °C for 30 sec, 72 °C for 1 min and a final extension at 72 °C for 5 min. The results were visualized by loading on 1% (w/v) agarose gel electrophoresis.

Southern blot analysis was performed by method previously described [19]. Twenty μ g of the genomic DNA was digested with the restriction enzymes *BamHI* (Promega) at 37 °C overnight. The digested DNA was separated on a 1% agarose gel and shifted to an Amersham Hybond N+ membrane (GE Healthcare, UK) using capillary transfer. The membrane was hybridized with a DIG-labelled DNA probe (Roche, Germany) and incubated overnight at 42 °C with gentle agitation. The DNA probe was prepared by amplification of *pRI1010N*. *OsWee1* by PCR using FNdeI and RBamHI primers and the PCR product was then labelled with DIG. The processes of probe preparation and washing of the membrane to remove the unbound probe were performed according to the manufacturer's instructions (Roche). Hybridization was visualized by exposing the membrane to X-ray (Fuji Film).

2.7. RNA isolation and reverse transcriptase PCR (RT-PCR)

The total RNA was isolated from 100 mg of young leaves of putative transformant lines and non-transformant 30-day-old rice plants using RNAprep pure plant kit (Tiangen, Beijing). The first-strand cDNA was prepared from 1 μ g of total RNA using iScriptTM cDNA Synthesis Kit (BIO-RAD) according to the manufacturer's instructions. The synthesized cDNA was then used as template to check the expression of *OsWee1* using OsWee-F and OsWee-R primers (Table 2). The RT-PCR conditions were 95 °C for 3 min, followed by 25 cycles at 95 °C for 30 sec, 56 °C for 30 sec, 72 °C for 1 min and a final extension at 72 °C for 5 min. The primer pairs of OsActin (Table 2) was used to amplify the *OsActin* reference gene, at 95 °C for 3 min, followed by 25 cycles at 95 °C for 3 osec, 48 °C for 30 sec, 72 °C for 5 min. The results were visualized by loading on 1% (w/v) agarose gel electrophoresis.

3. Results and discussion

3.1. Cloning of OsWee1

The cloning of full length *OsWee1* gene (1239 bp) in the expression vector (pRI101-ON) was performed on the *NdeI* and *BamHI* restriction sites. Clone of *OsWee1* that obtained in previous study [14] was amplified using a pair of FNdeI and RBamHI primers (Table 2) in order to add those restriction sites on the open reading frame (ORF) of *OsWee1* before cloning. The construct of *OsWee1* gene in the expression vector was shown in Fig. 1A. The construct of *OsWee1* was then transformed into *E. coli* strain XL10 gold competent cells and selected using antibiotic kanamycin on the growth media. To verify the positive clones, 20 recombinant colonies were randomly selected and prepared for the colony PCR analysis using FNdeI and RBamHI primers (Table 2). As shown in Fig. 1B, the presence of 1251 bp of *OsWee1* gene was found in 12 clones which were confirmed by colony PCR and visualized on agarose gel. For further analysis, clone no. 7 was selected.

In order to evaluate the correct size and flanking of the *OsWee1* clones, PCR analysis, restriction enzymes digestion and sequencing were performed. The PCR reaction was conducted using 4 pairs of primers, CaMV-F/RV-R, FNdel/RBamHI, nptII-F/nptII-R, CaMV-F/OsWee-R (Table 2). The PCR result showed that expected bands appear at the appropriate sizes, 1868 bp, 1251 bp, 524 bp and 1504 bp, respectively (Fig. 2 A). Digestion was performed by single or combination of *HindIII*, *BamHI* and *EcoRI* restriction enzymes. Open reading frame of *OsWee1* contains *BamHI* and *EcoRI* enzymes,

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573–579

while *HindIII* was present two sites in the construct. Then, this clone was further conformed by sequencing (*data not shown*). The correct band sizes of *OsWee1* after digestion was shown in Fig. 2B. The nucleotide sequence analysis showed a full length region of *OsWee1* about 1239 bp and a 100% homolog with cDNA of *OsWee1* in GeneBank (KX758541). It suggested that the desired recombinant *pRI101ON*. *OsWee1* was successfully prepared to expressed in *Agrobacterium*.

3.2. Transformation of OsWee1 into Agrobacterium

The recombinant construct of *pRI1010N*..*OsWee1* was transformed into *Agrobacterium* strain GV3101 and selected on YEP solid medium supplemented with 50 mg/L kanamycin, 100 mg/L rifampicin and 12.5 mg/L gentamicin antibiotic (Fig. 3A). Twelve colonies were randomly selected for PCR analysis using FNdel and RBamHI primers and 4 positive clones harboring *OsWee1* were obtained by PCR (Fig. 3B). This results clearly confirm that recombinant *pRI1010N*..*OsWee1* was transformed into *Agrobacterium*, and it can be used for transformation in rice.

The successful Agrobacterium-mediated transformation in rice has been achieved using various methods. However, in monocot plants remains limited because it is not a natural host for *Agrobacterium* [20]. Many researchers have developed methods for Agrobacterium-mediated transformation in monocotyledons, especially in utilizing of different *Agrobacterium* strains [21,22], piercing method and vacum infiltration [23], choosing of different cultivars [24] and type of explants [25,26]. The basic protocol of Agrobacterium-mediated transformation [27] was develop for induction of callus derived from scutella seeds. However, since most of indica rice genotypes have less regeneration potential [28], we conducted Agrobacterium-mediated transformation in rice using 7-day-old rice sprouts as explants.

A hundred explants were infected by *Agrobacterium* suspension for 20 min and co-cultivation in dark condition for 2 d followed by cefotaxime-antibiotic treatment. Transformation was carried out twice with total 200 explants. However the percentage of rice transformation efficiency is very low (2%) (Table 3). Previous study reported [29] numerous factors that can improve transformation efficiency, was lighting condition, temperature, co-cultivation periods and *Agrobacterium*-density during co-cultivation step. The most critical factor reported was the period of co-cultivation. The successful integration of target gene into the plant genome occurs mainly during co-cultivation. Sahoo and Tuteja [29] found that suitable method for rice transformation was infection for 20 min followed by co-cultivation for 2 day. An extension of co-cultivation period, caused arising of excessive bacterial growth which results in inhibiting explants growth, reduce the number of shoots generation and finally causing death of explant. Rashid *et al.* [30] reported that to anticipate those problems, reducing the density of bacteria into 0.1 - 0.2 OD prevents overgrowth bacteria during co-cultivation.

The co-cultivated explants were transferred in the selection medium, to inhibit the formation of non-transformant explants and eliminate the residual of Agrobacterium. The first shoots produced from the explants were subjected to three successive propagation cycles with the same level concentration of antibiotic. This method is applied to select putative transformants and reduce false positive transformants or eliminate the chimeric of transgenic explants [31]. After three sub-cultures, the non-transformant plantlet were turned into white (chlorosis), while the putative transformants were able to survive and grow normally (Fig. 4A). The lacking of *nptII* gene in the non-transgenic plants caused inhibing of chlorophyll development and induce chlorosis at the shoot of rice plantlets. In contrast to the transgenic overexpressing OsWee1 showed normal green at the shoots. The absence of *nptII* gene will suppress the growth of non-transformant roots caused by unability of root to inactivated the kanamycin in the media (Fig. 4A). Similar results were also obtained in previous study [32] that increased concentration of kanamycin may lead to inhibition of root growth. Kanamycin inhibits the synthesis of protein in plastid and mitochondrial. Kanamycin acts as an inhibitor, active destroy the function of ribosome following by inhibition of translational initiation. Another way, the response was automatically inhibit protein synthesis and effects on reducing the growth and development of plants.

3.3. Expression of OsWee1 in rice

The putative transgenic plants which survived in selection *in vitro* medium were then transferred to soil and placed in the greenhouse under agronomic conditions (Fig. 4B). These plants were then analyzed for their transgenic status by PCR analysis. PCR analysis showed that the *nptll* gene 550 bp in size was found in 4 of putative transgenic lines (Fig. 4C). To determine the expression of *OsWee1* at the RNA levels, we performed a RT-PCR analysis. The quantity of RNA level is a reflection of the level of transcription. As shown in Fig. 5A, the transcript levels of rice overexpress

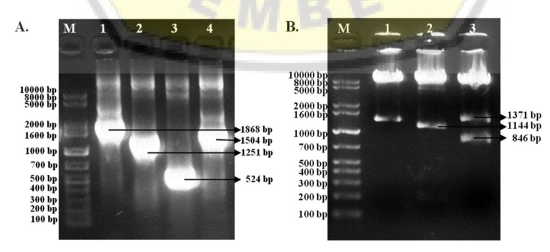


Fig. 2. Analyze of recombinant *pRI1010N*. *OsWee1*. A. PCR analysis of positive clones using several primers (M: 1 kb Tiangen Ladder, line 1–4: CaMV-F/RV-R primer, Fndel/RBamHI primer, nptII primer, and CaMV-F/OsWee-R, respectively); B. Restriction digestion with *HindIII* (line 1), *EcoRI/BamHI* (line 2) and *HindIII/BamHI* (line 3).

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579

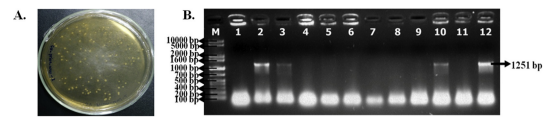


Fig. 3. Transformation pR11010N: OsWee1 into Agrobacterium. A. Agrobacterium colonies strain GV 3101 harboring recombinant DNA of OsWee1; B. PCR analysis of OsWee1 using a set of FNdel and RBamHI primers and the estimation size of OsWee1 fragment.

Table 3

The percentage of rice transformation efficiency.

Transformation number	Number of infected seedling	Number of selected plant	Number of transformant ^a	Transformation efficiency (%)
1	100	3	3	3%
2	100	2	1	1%
Total	200	5	4	2%

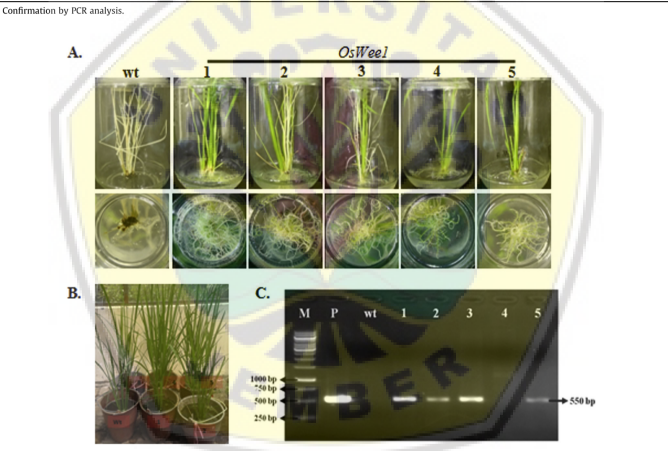


Fig. 4. Transgenic rice overexpressing of *OsWee1* gene. A. Phenotypic selection of transgenic rice overexpressing *OsWee1*, rice planlet (upper) and root (lower) of wildtype (wt) and transgenic (lane 1–5); B. Transgenic plants after acclimatization in green house; C. PCR analysis of the T0 transgenic plants (line 1–5), *wildtype* (WT), pRI101ON. *OsWee1* (P) as positive control, and DNA marker (M).

ing *OsWee1* were higher compared to the *wildtype*. The data indicates that *OsWee1* driven by CaMV35S was expressed in rice.

Southern blot analysis of transgenic rice was performed to prove integration of the transgene into the plant genome and to determine copy number of the T-DNA. Southern blot analysis was conducted using a gDNA isolated from leaves of four transgenic rice lines. The Southern blot analysis showed that the transgenic rice displayed one to three hybridized DNA copy with a difference in molecular size suggesting the independent transformation events. The hybridized DNA was not found in the genome of the WT plant (Fig. 5B). These results confirmed that the copy of the *OsWee1* gene was integrated into the genome of the transgenic rice.

The transgenic plants required stability of expression to be used in seed production [33], as well as *OsWee1* overexpression plants. In the present study, four among five independent transformation events have been shown to carry multiple copies of the T-DNA. Multiple copies of the transgene are prone for transgene inactivation, silencing, and likely to cause a high frequency of insertional mutagenesis [34]. To obtained stability of expression, the selection will be carried out through anther culture in the second generation

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579

578

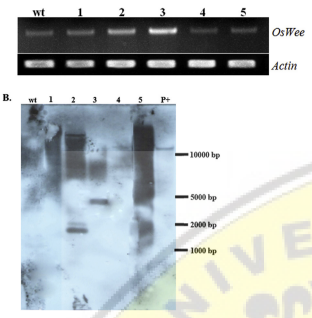


Fig. 5. Analysis of the first generation of transgenic plants (TO). A. RT-PCR analysis of the TO transgenic plants overexpressing *OsWee1* using full length *OsWee1* primer (upper) and *Actin* gene as a control (lower); B. Southern blot analysis of transgenic plants (line 1–5), wildtype (WT), pR1010N. *OsWee1* (+P), and DNA marker (M).

of transgenic rice. Regeneration of haploid plants from anther culture followed by chromosomal doubling can produce double haploid or pure line of plants. This result will provide an opportunity to accelerate the time for the formation of inbreed line which is normally through several inbreeding cycles [35].

Acknowledgements

We thank to Dr. M. Su'udi (Jember University) for helpful comments on this work. This research was supported by Competency-Based Grant (Hibah Berbasis Kompetensi 2018) No. 023/SP2H/LT/ DRPM/2018 from The Ministry of Research, Technology, and Higher Education of Indonesia to Dr. Netty Ermawati.

References

- Dewitte W, Murray JAH. The plant cell cycle. Annu. Rev. Plant Biol. 2018;54 (2003):235–64. doi: <u>https://doi.org/10.1146/annurev.arplant.54.031902.134836</u> [May 29, 2018].
- [2] Harashima H, Dissmeyer N, Schnittger A. Cell cycle control across the eukaryotic kingdom. Trends Cell Biol. 2013;23(7):345–56. doi: <u>https://doi. org/10.1016/j.tcb.2013.03.002</u> [May 29, 2018].
- [3] Nurse P, Thuriaux P. Regulatory genes controlling mitosis in the fission yeast Schizosaccharomyces pombe. Genetics 1980;96(3):627–37 [September 21, 2018] https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1214365/.
- [4] Shiozaki K, Russell P. Cell-cycle control linked to extracellular environment by MAP kinase pathway in fission yeast. Nature 1995;378(6558):739–43. doi: https://doi.org/10.1038/378739a0 [September 21, 2018].
- [5] Nurse P. Genetic control of cell size at cell division in yeast. Nature 2018;256 (1975):547–51. doi: <u>https://doi.org/10.1038/256547a0</u> [May 29, 2018].
- [6] Do K, Doroshow JH, Kummar S. Wee1 kinase as a target for cancer therapy. Cell Cycle 2013;12(19):3159–64. doi: <u>https://doi.org/10.4161/cc.26062</u> [May 29, 2018].
- [7] Kalhorzadeh P, Hu Z, Cools T, Amiard S, Willing E, De Winne N, et al. Arabidopsis thaliana RNase H2 deficiency counteracts the needs for the WEE1 checkpoint kinase but triggers genome instability. Plant Cell 2018;26 (2014):3680–92. doi: <u>https://doi.org/10.1105/tpc.114.128108</u> [May 29, 2018].
- [8] Sørensen CS, Syljua RG. Safeguarding genome integrity: the checkpoint kinases ATR, CHK1 and WEE1 restrain CDK activity during normal DNA replication. Nucleic Acids Res 2012;40(2):477–86. doi: <u>https://doi.org/10.1093/nar/gkr697</u> [May 29, 2018].

- [9] Caparelli ML, O'Connell MJ. Regulatory motifs in Chk1. Cell Cycle 2013;12 (6):916–22. doi: <u>https://doi.org/10.4161/cc.23881</u> [May 29, 2018].
 [10] Su C, Zhao H, Zhao Y, Ji H, Wang Y, Zhi L, et al. RUG3 and ATM synergistically
- [10] Su C, Zhao H, Zhao Y, Ji H, Wang Y, Zhi L, et al. RUG3 and ATM synergistically regulate the alternative splicing of mitochondrial nad2 and the DNA damage response in Arabidopsis thaliana. Sci Rep 2018;7(2017):1–14. doi: <u>https://doi. org/10.1038/srep43897</u> [May 29, 2018].
- [11] Yu Z, Zhang M, Wang G, Xu D, Keifenheim D, Franco A, Cansado J, Masuda H, Rhind N, Wang Y, Jin Q. Fission yeast nucleolar protein Dnt1 regulates G2/M transition and cytokinesis by downregulating Wee1 kinase. J Cell Sci 2014;126 (21):4995–5004. doi: <u>https://doi.org/10.1242/jcs.132845</u> [May 29, 2018].
- [12] Russell P, Nurse P. Negative regulation of mitosis by weel⁺, a gene encoding a protein kinase homolog. Cell 2018;49(1987):559–67. doi: <u>https://doi.org/ 10.1016/0092-8674(87)90458-2</u> [May 29, 2018].
- [13] Sun Y, Dilkes BP, Zhang C, Dante RA, Carneiro NP, Lowe KS, Jung R, Gordon-Kamm WJ, Larkins B. Characterization of maize (Zea mays L.) Wee1 and its activity in developing endosperm. Proc Natl Acad Sci 1999;96:4180–5. doi: https://doi.org/10.1073/pnas.96.7.4180 [May 29, 2018].
- [14] Gonzalez N, Gevaudant F, Hernould M, Chevalier C, Mouras A. The cell cycleassociated protein kinase WEE1 regulates cell size in relation to endoreduplication in developing tomato fruit. Plant J 2007;51(4):642–55. doi: https://doi.org/10.1111/j.1365-313X.2007.03167.x [May 29, 2018].
- [15] Sorrell DA, Marchbank A, McMahon K, Dickinson JR, Rogers HJ, Francis D. A WEE1 homologue from Arabidopsis thaliana. Planta 2002;215(3):518–22. doi: https://doi.org/10.1007/s00425-002-0815-4 [May 29, 2018].
- [16] Ermawati N, Wibisono Y. Early isolation of cell cycle-associated protein kinase (OsWee) gene in rice (Oryza sativa L.). Pak. J. Biotechnol. 2017;14(1):71-6 [May 29, 2018] http://www.pjbt.org/uploads/PJBT-VOL-14-NO-1-OF-YEAR-2017%20%2811%29.
- [17] Sambrook J, Russell. Molecular cloning a laboratory manual. third ed. Cold Spring Harbor Laboratory Press; 2001.
- [18] Dellaporta SL, Wood J, Hicks JB. A plant DNA minipreparation: version II. Plant Mol Biol Rep 1983;1:19–21. doi: <u>https://doi.org/10.1007/BF02712670</u> (September 19, 2018).
- [19] Apriasti R, Widyaningrum S, Hidayati WN, Sawitri WD, Darsono N, Hase T, et al. Full sequence of the coat protein gene is required for the induction of pathogen-derived resistance against sugarcane mosaic virus in transgenic sugarcane. Mol Biol Rep 2018;1(2018):1–10. doi: https://doi.org/10.1007/s11033-018-4326-1 [September 19, 2018].
- [20] Aldemita RR, Hodges TK. Agrobacterium tumefaciens-mediated transformation of *japonica* and *indica* rice varieties. Planta 2018;199 (1996):612–7. doi: https://doi.org/10.1007/BF00195194 [May 29, 2018].
- [21] Balaji V, Rajamuni P, Sridevi G, Veluthambi K. Agrobacterium-mediated Transformation Efficiency in Blackgram and Rice Enhanced by Multiple Copies of pTiBo542 virB and virG. Indian J Biotechnol 2003;2:138–46 [May 29, 2018] http://nopr.niscair.res.in/handle/123456789/11283.
- [22] Ratanasut K, Rod-In W, Sajipuli K. In planta agrobacterium-mediated transformation of rice. Rice Sci 2017;24(3):181–6. doi: <u>https://doi.org/</u> 10.1016/j.rsci.2016.11.001 [May 29, 2018].
- [23] Lin J, Zhou B, Yang Y, Mei J, Zhao X, Guo X, Huang X, Tang D, Liu X. Piercing and vacuum infiltration of the mature embryo: a simplified method for Agrobacterium-mediated transformation of indica rice. Plant Cell Rep 2009;28:1065–74. doi: <u>https://doi.org/10.1007/s00299-009-0706-2</u> [May 29, 2018].
- [24] Saika H, Toki S. Mature seed-derived callus of the model indica rice variety Kasalath is highly competent in *Agrobacterium*-mediated transformation. Plant Cell Rep 2010;29(12):1351–64. doi: https://doi.org/10.1007/s00299-010-0921-x [May 29, 2018].
- [25] Manimaran P, Kumar GR, Reddy MR, Jain S, Rao TB, Mangrauthia SK, et al. Infection of early and young callus tissues of indica rice BPT 5204 enhances regeneration and transformation efficiency. Rice Sci 2018;20(2013):415–26. doi: https://doi.org/10.1016/S1672-6308 (13)60153-5 [May 29, 2018].
- [26] Dey M, Bakshi S, Galiba G, Sahoo L, Panda SK. Development of a genotype independent and transformation amenable regeneration system from shoot apex in rice (*Oryza sativa* spp. *indica*) using TDZ. Biotech 2012;2(3):233–40. doi: <u>https://doi.org/10.1007/s13205-012-0051-y</u> [May 29, 2018].
- [27] Hiei Y, Ohta S, Komari T, Kumashiro T. Efficient transformation of rice (Oryza sativa L.) mediated by Agrobacterium and sequence analysis of the boundaries of the T-DNA. Plant J 1994;6(2):271–82. doi: <u>https://doi.org/10.1046/j.1365-313X.1994.6020271.x</u> [May 29, 2018].
- [28] Nishimura A, Ashikari M, Lin S, Takashi T, Angeles ER, Yamamoto T, Matsuoka M. Isolation of a rice regeneration quantitative trait loci gene and its application to transformation systems. PNAS 2005;102(33):11940-4. doi: <u>https://doi.org/10.1073/pnas.0504220102</u> [May 29, 2018].
- [29] Sahoo RK, Tuteja N. Development of Agrobacterium-mediated transformation technology for mature seed-derived callus tissues of indica rice cultivar IR64. GM Crops Food: Biotechnol Agric Food Chain 2012;3(2):123–8. doi: <u>https:// doi.org/10.4161/gmcr.20032</u> [May 29, 2018].
- [30] Rashid H, Afzal A, Khan MH, Chaudhry Z, Malik SA. Effect of bacterial culture density and acetosyringone concentration on *Agrobacterium* mediated transformation in wheat. Pak J Bot 2010;42(6):4183–9 [May 29, 2018] https://www.researchgate.net/publication/260156533_Effect_of_ bacterial_culture_density_and_acetosyringone_concentration_on_ Agrobacterium_mediated_transformation_in_wheat.
- [31] Nap JP, Bijvoet J, Stiekema WJ. Biosafety of kanamycin-resistant transgenic plants. Transgenic Res 1992;1(6):239–49 [May 29, 2018] https://link. springer.com/article/10.1007/BF02525165.

A.

F.H.H. Prasetyo et al./Journal of Genetic Engineering and Biotechnology 16 (2018) 573-579

- [32] Bibi N, Fan K, Yuan S, Ni M, Ahmed IM, Malik W, Wang X. An efficient and highly reproducible approach for the selection of upland transgenic cotton produced by pollen tube pathway method. Aust J Crop Sci 2013;7 (11):1714–22 [May 29, 2018] https://search.informit.com. au/documentSummary;dn=644871058289920;res=IELHSS.
- [33] Takaiwa F, Wakasa Y, Takagi H, Hiroi T. Rice seed for delivery of vaccines to gut mucosal immune tissues. Plant Biotechnol J 2015;13(8):1041–55. doi: <u>https:// doi.org/10.1111/pbi.12423</u> [May 29, 2018].
- [34] Finn TE, Wang L, Smolilo D, Smith NA, White R, Chaudhury A, Dennis ES, Wang M. Transgene expression and transgene-induced silencing in diploid and autotetraploid arabidopsis. Genetic 2011;187(2):409–23. doi: <u>https://doi.org/ 10.1534/genetics.110.124370</u> [September 22, 2018].
- [35] Silva TD. Indica rice anther culture: can the impasse be surpassed? Plant Cell Tissue Organ Culture 2010;100(1):1–11. doi: <u>https://doi.org/10.1007/s11240-009-9616-9</u> [May 29, 2018].



Research article Open access <u>Agrobacterium tumefaciens-mediated transformation of Dendrobium lasianthera J.J.Sm: An</u> <u>important medicinal orchid</u> Edy Setiti Wida Utami, Sucipto Hariyanto, Yosephine Sri Wulan Manuhara

Pages 703-709

In Silico Biotechnology

Short communication Open access Screening of anti-inflammatory phytocompounds from *Crateva adansonii* leaf extracts and its validation by *in silico* modeling

Rathinavel Thirumalaisamy, Subramanian Ammashi, Govarthanan Muthusamy Pages 711-719

Research articleOpen access

In silico structural and functional modelling of Antifreeze protein (AFP) sequences of Ocean pout (*Zoarces americanus*, Bloch & Schneider 1801)

Manojit Bhattacharya, Arpita Hota, Avijit Kar, Deep Sankar Chini, ... Basanta Kumar Das Pages 721-730

Research article Open access

In silico structural homology modeling of nif A protein of rhizobial strains in selective legume plants

Sadam D.V. Satyanarayana, M.S.R. Krishna, Pindi Pavan Kumar, Sirisha Jeereddy Pages 731-737

Research article Open access

<u>In silico analysis of squalene synthase in Fabaceae family using bioinformatics tools</u> Zahra Aminfar, Masoud Tohidfar Pages 739-747

Pages /39-/4/

Research article Open access

In silico studies on bacterial xylanase enzyme: Structural and functional insight Bhramar Dutta, Aparna Banerjee, Priyanka Chakraborty, Rajib Bandopadhyay Pages 749-756

Research article Open access

In silico thermodynamic stability of mammalian adaptation and virulence determinants in polymerase complex proteins of H9N2 virus

Zienab Mosaad, Abdelsatar Arafa, Hussein A. Hussein, Mohamed A. Shalaby Pages 757-767

Research article Open access Interaction of rs316019 variants of SLC22A2 with metformin and other drugs- an *in silico* analysis Also Achforen Soiith, Teamia Islam, Nilaniana Baul, Solina Vacamin

Abu Ashfaqur Sajib, Tasmia Islam, Nilanjana Paul, Sabina Yeasmin