Diversity of endophytic bacteria and microfungi in *Syzygium cumini* fruit from West Java, Indonesia

IDA INDRAWATI *, NIA ROSSIANA**, MUHAMMAD FAIZAL FATHURROHIM ***

Department of Biology, Faculty of Mathematics and Natural Science, Universitas Padjadjaran, Jl. Raya Bandung-Sumedang Km. 21, Sumedang 45363, West Java, Indonesia. Tel./fax: +62-22-7796412, *email: ida.indrawati81@gmail.com, **niarossiana@yahoo.com, ***faizalmaret26@gmail.com

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Abstract. Rossiana N, Fathurrohim MF, Indrawati I. 2021. Diversity of endophytic bacteria and microfungi in *Syzygium cumini* fruit from West Java, Indonesia. Biodiversitas 22: 3943-3948. *Syzygium cumini* L. Skells is a native evergreen tropical tree in Southeast Asia belonging to the Myrtaceae family, known as the Java plum, jambul, jambolan, jamblang, or jaman. The bacterial and fungal endophytes associated with fruit have not been determined and functionally characterized. The endophytic microbes live inside the surface-sterilized fruits and have no visibly harmful effects on the plants. The purposes of the study were to isolate, characterize, and determine the diversity of endophytic bacteria and fungi in *S. cumini* fruit. The endophytes from *S. cumini* fruit were observed morphologically for identification. The result of isolation and identification showed there are four bacterial isolate endophytes (*B. cereus*, *B. subtilis*, *B. megaterium*, and Bacillus sp.) and four fungal endophytes (Candida guilliermondii, Penicillium sp., Mycelia sterilia, and Aspergillus sp.) isolated from *S. cumini* fruit.

Keywords: Bacterial endophyte, biodiversity, fungal endophyte, jamblang, *Syzygium cumini*

INTRODUCTION

With its thousands of islands, Indonesia has a myriad wealth of biological resources, especially tropical fruits. Approximately 329 types of fruits, both indigenous and introduced, could be found in Indonesia. Two hundred and sixty-six species of Indonesian indigenous fruits mostly grow wild in the forest and only a small portion has been cultivated (Hermanto et al. 2013).

*Syzygium cumini* L. Skells (Java plum, jambul, jambolan, jamblang, or jaman) is one of the plants that have many benefits in Indonesia. However, Indonesian people do not know much about the benefits and nutritional value of *S. cumini* fruit. *S. cumini* fruit contains flavonoids, quinones, steroids, and polyphenols (Marliani et al. 2014). So, it might be potential as a medicinal plant.

The plant is the host for various types of endophytic microbes. Natural products produced by endophytic microbes were reported to exhibit a wide range of biological activities and are categorized into various categories of chemical compounds including alkaloids, terpenoids, steroids, lactones, phenolic compounds, quinones, and lignans (Anjum and Chandra 2015).

The internal non-sterile plant tissue is inhabited by various types of fungi and bacteria known as endophytes (Goryluk et al. 2009). Endophytic microbes in plant tissues do not cause disease symptoms in their hosts (Anjum and Chandra 2015). Endophytic microbes can obtain nutrients to complete their life cycle from the host plant, while the host gets protection against plant pathogens from compounds produced by endophytic microbes (Ariyanto et al. 2013).

Endophytic microbes can have a mutualistic symbiosis with their host plant. The search for endophytic microbes from different plant species in different ecosystems will be beneficial. The diversity of endophytic microbes is under-explored although they are a great resource for application in the fields of agriculture, medicine, pharmacy, farm, and industry (Zhang et al. 2016). The diversity of microorganisms i.e., the diversity of endophytic microbes in *S. cumini* fruit is very important, to be studied.

Endophytic bacteria generally enter the plant tissues through roots and plant parts exposed to direct air such as flowers, stems, and cotyledons (Desriani et al. 2014). Endophytic bacteria can be isolated through internal plant tissue in which its outer surface has been sterilized (Munif et al. 2012).

Endophytic fungi are a group of fungi that colonize living and internal tissues of plants without causing any immediate, overt negative effects (Padhi et al. 2013). Endophytic fungi have an important role in their dead hosts to initiate biological degradation to start the nutrient recycling process (Shekhawat et al. 2010).

Studies on the diversity of endophytic bacteria and fungi in *S. cumini* fruit in Indonesia have not been conducted. This study was conducted to determine the diversity of bacteria and endophytic fungi in *S. cumini* fruit.

MATERIALS AND METHODS

Plant material

*S. cumini* fruit was collected from the plant grown in Padjadjaran University Campus, Sumedang, West Java, Indonesia.
Isolation of endophytic bacteria

Syzygium cumini fruits were washed under running water for 5 minutes and then dipped in 96% ethanol for 2 seconds, followed by rinsing with sterile distilled water for 30 seconds and then rinsed using 1% NaOCl solution for 5 minutes. As much as 100 grams of the fruits, added with 9 ml of sterile physiological NaCl, and then finely crushed. As much as 1 ml of finely crushed fruits was diluted to 10^2 dilutions. One ml of diluted crushed fruit was inoculated on nutrient jell for the last three dilutions (10^5, 10^6 and 10^7) and incubated for 24 hours at 37°C. After incubation, one of each species of different morphological bacterial colonies were subcultured onto a nutrient agar slope. Pure bacterial isolate was identified.

Identification of endophytic bacteria

Pure bacterial cultures were identified microscopically, i.e., colony color, elevation, surface, and bacterial colonies. Gram-staining carried out microscopic observation to determine bacterial cell shape and biochemical characterization was performed using VITEK 2.0 compact system.

Isolation of endophyte fungi

The fruit was washed under running water for 5 minutes, then rinsed with 70% alcohol for 5 minutes, followed by soaking in 1% NaOCl solution for 5 minutes and then drained. The fruit was then rinsed with sterile distilled water for one minute twice. Then the fruit was crushed and was diluted with series from 10^{-1} to 10^{-2} in medium Potato Dextrose Agar. The last three dilutions (1ml) were added into the petri dish, as much as 1 ml and 3 last dilutions were inserted into a petri dish containing the PDA medium and were then homogenized, followed by incubation at room temperature for 48-72 hours. Emerging fungi on Petri dishes were subcultured on PDA to obtain a pure culture.

Identification of endophyte fungi

The moist chamber performed fungal identification. Macroscopic observations were carried out including colony color, colony surface, concentric circle, and radial surface lines of the colony. Microscopic observations included the morphology of spore, sporangium, and conidia.

RESULTS AND DISCUSSION

Isolation of endophytic bacteria and fungi from S. cumini fruit resulted in 4 isolates of bacteria (FIN 2, FIN 3, FIN 4, and FIN 11) and 4 isolates of fungi. Based on Gram-staining, the four endophytic bacteria were Gram-positive bacilli. Macroscopic characters of endophytic bacteria FIN 2 were a white colony, round shape, flat colony elevation, flat edge, and smooth surface (Figure 1.A). Microscopically, endophytic bacteria FIN 2 is a purple Gram-positive bacteria bacilli (Figure 1.B). Endophytic bacteria FIN 3 has macroscopic characteristics of spherical shaped colonies, jagged edges, flat elevation, rough surface, and slightly greenish-white (Figure 2.A), and microscopic characteristics as Gram-positive bacilli (Figure 2.B). Endophytic bacterial FIN 4 has macroscopic characteristics of spherical shaped colonies, flat edges, flat elevations, smooth, and white surface (Figure 3.A) with microscopic characteristics as Gram-positive bacilli (Figure 3.B). Endophytic bacteria FIN 11 has macroscopic characteristics of spherical-shaped colonies, jagged edges, raised elevations, rough surface with white and greenish margins in the middle (Figure 4.A), and microscopic characteristics as Gram-positive (Figure 4.B).

The results of the biochemical character of four endophytic bacteria from S. cumini fruit using VITEK Compact 2.0 were prested Table 1.

The results of the identification of endophytic fungi from S. cumini fruit (MFFJB, MFFJE, MFFJF, and MFFJG) were presented in Table 2.

Discussion

Bacillus cereus (FIN 3)

Bacillus cereus is a Gram-positive, aerobic, facultatively anaerobic, spore-forming, and mesophilic bacterium, with growth temperatures from 10°C to 48°C with optimal growth between 28°C and 35°C. B. cereus measures 1 x 3 – 4 μm. Most B. cereus strains are motile via peritrichous flagella, grow on solid growth media as irregular colonies, use glucose as a source of carbon (but not mannitol, arabinose, or xylose), hydrolyze starch and gelatin, show hemolytic activity, are resistant to ampicillin, and display pronounced lecithinase activity (Vilas-Boas et al. 2007). The endophytic B. cereus isolated from Garcinia xanthochymus based nanoparticles has the potential to be developed as promising antibacterial and antioxidant agents (Mujaddidi et al. 2021).

Bacillus megaterium (FIN 2)

Bacillus megaterium is a Gram-positive spore-forming bacteria. As most of the spore-formers, it is usually found in the soil, from which it can easily be transmitted to the foods we consume (Periago et al. 2006). From macroscopic and microscopic observations and biochemical tests, it is shown that B. megaterium is concave, smooth, and milky-white. The cell morphology shows that the cell is rod-shaped, Gram-positive, and sporous (Andriani et al. 2017). Some B. megaterium proteins are very important in the food industry and pharmacy. B. megaterium secretes a variety of enzymes, ranging from amylases used in the bakery industry to penicillin amidase used to manufacture new synthetic antibiotics (Mobitec 2008). B. megaterium can also produce several other enzymes, such as mutarotase, glucose dehydrogenase, β-galactosidase, and cellulose (Andriani et al. 2017).

Bacillus subtilis (FIN 4)

Bacillus subtilis is a Gram-positive, aerobic, non-encapsulated, mobile, and spore-bearing bacterium, a basil chain with a size of 0.8-0.7 or 2-3 um, commonly found in nature. Bacterial colonies have a white or slightly yellowish, rough, and opaque surface. Grow in the mesophilic temperature range of 25-35°C. Its active form is
usually spore-shaped. Therefore, it can survive in difficult conditions (Saleh et al. 2014). 

\textit{B. subtilis} is useful for biotechnology as well as industrial and agricultural applications. \textit{B. subtilis} can directly fight pathogens by producing a secondary metabolite of extracellular lytic enzymes to inhibit growth by quorum quenching to disrupt the communication of cell-to-cell expression from infectious expression in pathogenic bacteria (Alina et al. 2015).

\textbf{Bacillus sp. (FIN 11)}

The genus \textit{Bacillus} is Gram-positive bacteria, a stem cell shape with a cell size of 0.3-2.2 to 1.2-7.0 μm (Alina et al. 2015). It can also move freely and has good competence and survivability on rhizosphere and facultative anaerobes so that it can adapt to living in the soil under various environmental conditions (Yanti et al. 2018). \textit{Bacillus} sp. can produce digestive enzymes such as proteases and amylases that can help digestion and produce short-chain organic acids with antimicrobial properties (Sumardi et al. 2012). \textit{Bacillus} can also be used as potential biofertilizers, biopesticides, and non-pathogenic to plants.

\textbf{Candida guiliermondii (MFFJB)}

\textit{Candida guiliermondii} is the most common opportunistic fungus, a normal flora present in human skin and mucosal surfaces, but occasionally the cause of chronic onychomycosis, acutely osteomyelitis, septic arthritis, endocarditis, fungemia, and invasive infections (Girmenia et al. 2006). The yeast cells of \textit{C. guiliermondii} are both rounded and ellipsoidal, the length is 4.21 ± 0.3 μm and the diameter is 2.29 ± 0.1 μm (Hovnanyan et al. 2019). Cells of \textit{C. guiliermondii} are mostly heterogenous, mainly elongated in shape (approximately 2, 9, 10 cm). In contrast to \textit{Candida albicans}, \textit{C. guiliermondii} is unable to produce true hyphae. Nevertheless, under certain conditions, such as nitrogen or carbon deficiency, these budding yeast can efficiently switch to pseudohyphal structure harboring blast conidia circlet (Koehler et al. 1999; Papon et al. 2013).

\textbf{Penicillium sp. (MFFJE)}

\textit{Penicillium} is a genus of one of the most fungi found in different environmental and suitable environmental (temperature, humidity, pH). According to Subowo (2015) \textit{Penicillium sp.} can decompose cellulose and lignin compounds into simple carbon compounds required by microbes as an energy source (Carbon source). Penicillium species produce various secondary metabolites such as antibacterial (Petit et al. 2009). Many \textit{Penicillium} produced the different chemical types of secondary metabolite, while some of them are important in the field of medicine. Others are used for the production of mycotoxins, important drugs, and some of the \textit{Penicillium} species are used in industry, especially penicillin production (Garca-Estrada et al. 2011).

\begin{table}
| Name of compound biochemical test | Endophytic bacterial isolate |
|-----------------------------------|-----------------------------|
|                                   | FIN 2 | FIN 3 | FIN 4 | FIN 11 |
| Beta-xylosidase                   | -     | -     | -     | -      |
| L-lysine-arylamiadase             | -     | -     | -     | -      |
| L-aspartate- arylamiadase         | +     | -     | -     | -      |
| Leucine arylamiadase              | +     | +     | +     | +      |
| Phenylalanine arylamiadase        | +     | +     | +     | +      |
| L-proline arylamiadase            | -     | +     | -     | -      |
| Beta-galactosidase                | +     | -     | +     | +      |
| L-tryptophol- aryldiamide         | -     | -     | +     | +      |
| Alpha-galactosidase               | +     | -     | -     | +      |
| Alanine arylamiadase              | +     | +     | -     | -      |
| Tyrosine arylamiadase             | +     | -     | -     | -      |
| Beta-n-acetyl-glucosaminidase      | -     | +     | -     | -      |
| Ala-pha-pro arylamiadase          | +     | +     | +     | -      |
| Cyclodextrin                      | -     | -     | +     | +      |
| D-galactose                       | -     | -     | -     | -      |
| Glycogen                          | +     | -     | -     | -      |
| Myo-inositol                      | -     | -     | -     | -      |
| Methyl-A-D-glucopyranoside        | -     | -     | +     | -      |
| acidification                     | -     | -     | -     | +      |
| Ellman                             | +     | +     | -     | +      |
| Methyl-D-xyloside                 | -     | -     | -     | -      |
| Alpha-mannosidase                 | -     | -     | -     | -      |
| Maltotriose                       | -     | -     | -     | -      |
| Glycine arylamiadise              | +     | -     | -     | -      |
| D-Mannitol                        | +     | +     | +     | -      |
| D-Mannose                         | +     | -     | +     | -      |
| D-Melezitose                      | -     | -     | -     | -      |
| N-Acetyl-D-glucosamine            | +     | +     | -     | -      |
| Palatinose                        | +     | -     | -     | -      |
| L-Rhamnose                        | -     | -     | -     | -      |
| Beta-glucosidase                  | +     | +     | +     | -      |
| Beta-mannosidase                  | -     | -     | -     | -      |
| Phosphoryl choline                 | -     | -     | -     | -      |
| Pyruvate                          | +     | +     | +     | -      |
| Alpha-glucosidase                 | +     | -     | -     | -      |
| D-tagatose                        | -     | -     | -     | -      |
| D-trehalose                       | +     | +     | +     | +      |
| Inulin                           | -     | -     | +     | +      |
| D-glucose                         | +     | +     | -     | -      |
| D-ribose                         | +     | +     | -     | -      |
| Putrescine assimilation           | -     | -     | -     | -      |
| Growth in 6.5% NaCl               | +     | +     | +     | -      |
| Kanamycin resistance              | -     | +     | -     | -      |
| Oleandomycin resistance           | +     | -     | -     | -      |
| Esculin hydrolyze                  | +     | -     | -     | -      |
| Tetrazolium RED                   | -     | -     | +     | +      |
| Polymixin_B resistance            | -     | -     | -     | -      |
| Probabilitas                      | 93%   | 94%   | 87%   | -      |
\end{table}
| Isolate code | Characteristics of colonies | Characteristics of fungal cells | Identification |
|-------------|-----------------------------|---------------------------------|----------------|
| MFFJB       | Milk white colony, smooth rounded shape colony, corrugated, concave in the middle | Large coccus cell shape | Candida guilliermondii |
| MFFJE       | Green to grayish colonies, uneven colony edges, look like velvet | Broom-like structure | Penicillium sp. |
| MFFJF       | White colonies, furry and wrinkled in the middle, on the white colony-like edges of the powder | The fungal cell has the only mycelium | Mycelia sterilia |
| MFFJG       | Colonies of green colonies, uneven colony edges, colonies shaped like cotton and growing upward. | Fungal cell resembles to fan | Aspergillus sp. |
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Figure 1. The bacterial isolate of FIN 2 (*Bacillus cereus*) on TSA (A) and Gram Stained FIN 2 (B)

Figure 2. The bacterial isolate of FIN 3 (*Bacillus megaterium*) on TSA (A) and Gram Stained FIN 3 (B)

Figure 3. The bacterial isolate of FIN 4 (*Bacillus subtilis*) on TSA (A) and Gram Stained FIN 4 (B)

Figure 4. The bacterial isolate of FIN 11 (*Bacillus sp.*) on TSA (A) and Gram Stained FIN 11 (B)

*Mycelia sterilia* (MFFJF)

*Mycelia sterilia* is a non-forming spore fungus. Endophytic fungus *Mycelia sterilia* is often found in the bifurcated mycelium assemblies, erect conidia, globose vesicles with uniseriate stigmates with conidial chain basipetal succession (Srinivas et al. 2015). Based on research Carbuncgo et al. (2017), *Mycelia sterilia* has a floccose texture, white greenish peripheral with a circular filamentous zone. *Mycelia sterilia* has hyphae with a length of 40 µ and a diameter of 8 µ and has absent spores. *Mycelia sterilia* is also found in vegetables.

*Aspergillus sp.* (MFFJG)

*Aspergillus* is probably the most widespread group of fungi in the human environment. Many species are found in a variety of substrates, including soil, forage products, various types of food products, dust, organic debris, and decomposing matter. *Aspergillus* species play an essential role in the recycling of carbon and nitrogen sources. Many *Aspergillus* species, including important pathogenic species, do not have special nutritional requirements and can grow in simple media, such as glucose-asparagine-phosphate broth, which contains a single protein hydrolysate (Gugnani 2003). *Aspergillus* has the characteristics of a spore-bearing structure called a conidial head, a basal foot of a bicep. The leg cells have more or less perpendicular hyphae and a vesicular conidiophore, having one or two synchronous cell layers and asexually formed spores known as conidia produced by phialides. The head of conidia of *Aspergillus* may be uniseriate or biseriate (Nyongesa et al. 2015).

In conclusion, the isolation of endophytic bacteria and fungus from *S. cumini* fruit successfully obtained 4 species of bacteria and 4 species of fungi. Endophytic bacteria from *S. cumini* fruit were identified as *Bacillus cereus*, *B. megaterium*, *B. subtilis*, and *Bacillus sp.* Identified endophytic fungi from *S. cumini* fruit consisted of *Candida guilliermondii*, *Penicillium sp.*, *Mycelia sterilia*, and *Aspergillus sp.*

REFERENCES

Alina SO, Constantinscu F, Petruta CC. 2015. Biodiversity of *Bacillus subtilis* group and beneficial traits of *Bacillus species* useful in plant protection. Romanian Biotechnol Lett 20 (5): 10737-10750.

Andriani Y, Rochima E, Safitri R, and Rahayu SR. 2017. Characterization of *Bacillus megaterium* and *Bacillus mycoides* bacteria as probiotic bacteria in fish and shrimp feed. ICSAFS Conf Proc 2017: 127-135. DOI: 10.18502/kls.v2i6.1029

Anjum N, Chandra N. 2015. Endophytic bacteria: Optimization of isolation procedure from various medicinal plants and their preliminary characterization. Asian J Pharm Clin Res 8(4): 233-238.

Ariyanto EF, Abadi AL, Djauhari S. 2013. Keanekaragaman jamur endofit pada daun tanaman padi (*Oryza sativa L*) dengan sistem Pengelolaan Hama Terpadu (PHT) dan konvensional di Desa Bayem, Kecamatan

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Kasembron, Kabupaten Malang. Jurnal Hama dan Penyakit Tumbuhan 1(2): 37-51. [Indonesian]

Carbungco ES, Pedrero NB, Panes VA, De la Cruz TE. 2015. Identification and characterization of endophytic fungi associated with the leaves of Moringa oleifera Lam. In: I Int Symp Moringa 1158: 373-380. DOI: 10.17660/ActaHortic.2017.1158.42

Desriani, Safira PUM, Bintang M, Rivai A, Lisdiansyanti P. 2014. Isolasi dan karakterisasi bakteri endofit dari tanaman binahong dan katapeng. Jurnal FK Unand 3(2): 89-93. DOI: 10.25077/fka.v3i2.33

García-Estrada C, Ullán RV, Albillos SM, Fernández-Bodega MA, Derek P, von Dohren H, Martin JP. 2011. A single cluster of coregulated genes encodes the biosynthesis of the mycotoxins Roquefortine C and Melgerin in Penicillium chrysogenum. J Chem Biol 18: 1499-1512. DOI: 10.1016/j.chembiol.2011.08.012

Gugnani, H C. 2003. Ecology and taxonomy of pathogenic Aspergilli. Front Biosci 8: 346-357. DOI: 10.2741/1002

Girmenia C, Pizzarelli G, Cristini F, Barchiesi F, Sperghini E, Scalise Martino P. 2006. Candida guilliermondii fungemia in patients with hematologic malignancies. J Clin Microbiol 44 (07): 2458-2464. DOI: 10.1128/JCM.00356-06

Goryluk A, Burlaga HB, Blaszczyk M. 2009. Isolation and characterization of bacterial endophytes of Chelidonium majus L. Polish J Microbiol 58 (4): 355-361.

Hovnanyan KO, Gasparyan HV, Marutyan SV, Navasardyan LH, Tschoumian AH. 2019. Comparative structural analysis of yeast Candida guilliermondii NP-4 Cultivated with and without nitrogen source. Proc Yerevan State Univ Chem Biol 53 (1): 53-58.

Hermanto C, Indrianti NLP, Hardiati S. 2013. Keragaman dan Kekayaan Buah Tropika Nusantara. Badan Penelitian dan Pengembangan Pertanian. Kementrian Pertanian. IAARD Press.

Husametto, H, Takahashi JA. 2009. Novel antifungal secondary metabolites from Penicillium sp. isolated from Brazilian cerrado soil. Electr J Biotechnol 12 (4): 8-9. DOI: 10.2225/vol12-issue4-fulltext-9

Sahl G, Kheirandish F, Azizi H, Azizi M. 2014. Molecular diagnosis and characterization of Bacillus subtilis isolated from Burn Wound in Iran. Res Mol Med 2 (2): 40-44. DOI: 10.18809/acadpub.rmm.2.2.40

Shekawat KK, Rao DV, Batra A. 2010. Morphological study of endophytic fungi inhabiting leaves of Melia azederach L. Intl J Pharm Sci Res 5 (3): 117-180.

Subowo YB. 2015. Pengujian aktivitas jamur Penicillium sp. R7.5 dan Aspergillus niger NK pada media tumbuh untuk mendukung pertumbuhan tanaman padi di lahan salin. Pros Sem Nas Masy Biodiv Indon 1 (5): 1136-1141. DOI: 10.13057/psmb/mi010529.

Srinivas RP, Nigam A, Aruna J, Silva WCD, Chikkaswanny BK. 2015. An investigation of biodiversity of endophytic fungi associated with some medical plants. Intl J Adv Res Eng Appl Sci 4 (2): 27-44

Sumardi S, Ekowat CN, Handayani K, Nurhayati N. 2012. Isolasi dan karakteristik Bacillus sp. penghasil antimikroba dari saluran pencernaan ayam kampung (Gallus domesticus). Prosiding Seminar Nasional Sains, Matematika, Informatika dan Aplikasi 3 (3): 306-311. [Indonesian]

Vilas-Boas GT, Perusa AP, Arantes OMN. 2007. Biology and taxonomy of Bacillus cereus, Bacillus anthracis, and Bacillus thuringiensis. Can J Microbiol 53: 673-687. DOI: 10.1139/w07-029.

Yanti W, Warnita, Reflin, Nasution CR. 2018. Caracterizacion de endofiticas Bacillus strain from tomato roots as growth promoter and biocontrol of Ralstonia solanacearum. Biodiversitas 19 (3): 906-911. DOI: 10.13057/biodiv/d190320.

Zhang HS. 2016. Two new secondary metabolites from the endophytic fungus Endomelanconiosis endophytica. Molecules 21 (943): 1-6. DOI: 10.3390/molecules21070943.