In a multicenter study, we determined a prevalence rate of 4% for azole-resistant *Aspergillus fumigatus* in Taiwan. Resistance emerged mainly from the environment (TR\textsubscript{L/G} 1.9/L98H, TR\textsubscript{I/G} /L98S/297/T/F495I, and TR\textsubscript{Y} /Y121F/T289A mutations) but occasionally during azole treatment. A high mortality rate observed for azole-resistant aspergillosis necessitates diagnostic stewardship in healthcare and antifungal stewardship in the environment.

Worldwide emergence of azole-resistant *Aspergillus fumigatus* since the late 2000s threatens human health (1). Azole resistance in *A. fumigatus* might develop during patient therapy with medical azoles or through exposure to azole fungicides in the environment; environmental exposure predominantly involves TR\textsubscript{L/G} 1.9/L98H and TR\textsubscript{I/G} /Y121F/T289A mutations in *cyp51A* (1).

Taiwan is an island country in eastern Asia that is geographically separated from mainland Eurasia and has a long history ofazole fungicide use. To delineate the influence of clinical and environmental use of azoles on resistance, we conducted a multicenter study that investigated 375 *A. fumigatus* sensu stricto isolates collected during August 2011–March 2018 from 297 patients at 11 hospitals in Taiwan (Appendix Table 1, Figure 1, https://wwwnc.cdc.gov/EID/article/26/4/19-0840-App1.pdf).

We confirmed the presence ofazole resistance by using the Clinical Laboratory Standard Institute method (Appendix Table 1) (2). Isolates resistant to ≥1 medical azoles (itraconazole, voriconazole, posaconazole, and isavuconazole) were defined asazole-resistant *A. fumigatus* and examined for resistance mechanisms, microsatellite-based phylogenetic relatedness, and growth rates following previously described methods (3,4).

Overall, 19 isolates from 12 patients were azole-resistant *A. fumigatus*. These isolates had resistance rates of 4.0% / patient and 5.1% / isolate analyses (Appendix Tables 2, 3). Ten (83.3%) patients harbored azole-resistant *A. fumigatus* that had environmental mutations, including TR\textsubscript{L/G} 1.9/L98H (5 isolates, 5 patients), TR\textsubscript{I/G} /L98S/297/T/F495I (7 isolates, 4 patients), and TR\textsubscript{Y} /Y121F/T289A (1 isolate) mutations. This observation

### Multicenter Study of Azole-Resistant *Aspergillus fumigatus* Clinical Isolates, Taiwan

Chi-Jung Wu, Wei-Lun Liu, Chih-Cheng Lai, Chien-Ming Chao, Wen-Chien Ko, Hsuan-Chen Wang, Ching-Tzu Dai, Ming-I Hsieh, Pui-Ching Choi, Jia-Ling Yang, Yee-Chun Chen

1Results from this study were presented in part at the 30th International Congress of Chemotherapy and Infection, November 24–27, 2017, Taipei, Taiwan.

Author affiliations: National Health Research Institutes, Zhunan, Taiwan (C.-J. Wu, H.-C. Wang, M.-I. Hsieh, Y.-C., Chen); National Cheng Kung University Hospital and College of Medicine, Tainan, Taiwan (C.-J. Wu, W.-C. Ko); Fu Jen Catholic University Hospital and College of Medicine, New Taipei, Taiwan (W.-L. Liu), Chi Mei Medical Center, Tainan (C.-C. Lai, C.-M. Chao); National Taiwan University Hospital and College of Medicine, Taipei, Taiwan (C.-T. Dai, P.-C. Choi, J.-L. Yang, Y.-C. Chen)

DOI: https://doi.org/10.3201/eid2604.190840

### References

1. Godfroid J. Brucellosis in wildlife. Rev Sci Tech. 2002; 21:277–86. https://doi.org/10.20506/rst.21.2.1333
2. Garin-Bastuji B, Oudar J, Richard Y, Gasteau J. Isolation of *Brucella melitensis* biovar 3 from a chamois (*Rupicapra rupicapra*) in the southern French Alps. J Wildl Dis. 1990;26:116–8. https://doi.org/10.7589/0090-3558-26.1.116
3. Ferroglio E, Toleri F, Bollo E, Bassano B. Isolation of *Brucella melitensis* from alpine ibex. J Wildl Dis. 1998;34:400–2. https://doi.org/10.7589/0090-3558-34.2.400
4. Muñoz PM, Boadella M, Arnal M, de Miguel MJ, Revilla M, Martinez D, et al. Spatial distribution and risk factors of brucellosis in Iberian wild ungulates. BMC Infect Dis. 2010;10:46. https://doi.org/10.1186/1471-2334-10-46
5. Lindstrom E. The role of medium-sized carnivores in the Nordic boreal forest. Finnish Game Research. 1989;46:53–63.
6. Alton GG, Jones LM, Pietz DE. Laboratory techniques in brucellosis. Monogr Ser World Health Organ. 1975;55:1–163.
7. Wang Q, Zhao S, Wureli H, Xie S, Chen C, Wei Q, et al. *Brucella melitensis* and *B. abortus* in eggs, larvae and engorged females of *Dermacentor marginatus*.Ticks Tick Borne Dis. 2018;9:1045–8. https://doi.org/10.1016/j.ttbdis.2018.03.021
8. Maquart M, Le Flèche P, Foster G, Tryland M, Ramisse F, Djønne B, et al. MLVA-16 typing of 295 marine mammal *Brucella* isolates from different animal and geographic origins identifies 7 major groups within *Brucella ceti* and *Brucella pinnipedialis*. BMC Microbiol. 2009;9:4. https://doi.org/10.1186/1471-2180-9-145
9. Li S, Hu X. Serological investigation on brucellosis of cattle and sheep in Nilka County. Chinese Livestock and Poultry Breeding. 2017;4:41–3.
10. Davis DS, Boeer WJ, Mims JP, Heck FC, Adams LG. *Brucella abortus* in coyotes. I. A serologic and bacteriologic survey in eastern Texas. J Wildl Dis. 1979;15:367–72. https://doi.org/10.7589/0090-3558-15.3.367
Genetic relatedness among *Aspergillus fumigatus* isolates based on microsatellite genotyping, Taiwan. Scale bar indicates percentage relatedness. AF, *A. fumigatus*; C, clinical; E, environmental; R, azole-resistant; S, azole-susceptible; STR, short tandem repeat; TW, Taiwan.

**Figure.** Genetic relatedness among *Aspergillus fumigatus* isolates based on microsatellite genotyping, Taiwan. Scale bar indicates percentage relatedness. AF, *A. fumigatus*; C, clinical; E, environmental; R, azole-resistant; S, azole-susceptible; STR, short tandem repeat; TW, Taiwan.
is consistent with the estimated global prevalence of azole resistance in Aspergillus (3%–6%) and the predominance of environmental resistance mechanisms in azole-resistant A. fumigatus (1,5).

Phylogenetic analysis showed that TR6/L98H/S297T/F495I isolates from 2 patients with pulmonary aspergillosis (isolates B44 and B51 in 2012, isolates E071, E073, and E074 in 2015) (Figure) belonged to a local microsatellite genotype widely distributed in the environment of Taiwan (3), indicating that this clone has locally evolved and adapted to the environment. The TR6/L98H isolates were genetically clustered with local environmental isolates or clinical isolates from China and Europe (Appendix Table 4). The TR6/Y121F/T289A isolate (S05–322) recovered in 2018, which colonized a patient without overseas travel, was genetically identical to a clone prevalent in the Netherlands and Tanzania (6), raising the concern of the intercountry transfer of resistant isolates.

All TR6/L98H/S297T/F495I, TR6/L98H, and TR6/Y121F/T289A isolates exhibited cross-resistance to difenoconazole and tebuconazole (both triazole fungicides) without fitness cost, demonstrated by normal growth rates (Appendix Figure 2). The TR6/L98H/S297T/F495I isolates and TR6/Y121F/T289A isolates were also resistant to prochloraz (an imidazole fungicide) (Appendix Table 2). The prevalence of TR6/L98H/S297T/F495I isolates in Taiwan might be attributed to widespread use of prochloraz over the past 3 decades. Studies have suggested an association between use of imidazole fungicides and emergence ofazole-resistant A. fumigatus with TR6/L98H/S297T/F495I mutations (7,8).

In Taiwan, the annual consumption of difenoconazole and tebuconazole has exceeded that of prochloraz since 2012 (Appendix Figure 3), further creating a favorable environment for maintenance and spread of TR6/L98H, TR6/L98H/S297T/F495I, and TR6/Y121F/T289A isolates. Thus, the One Health approach to implement environmental antifungal stewardship is warranted to minimize ongoing resistance selection in the fields.

Six azole-resistant A. fumigatus isolates with wild-type cyp51A were obtained from 2 patients. Four pan-azole-resistant urinary isolates were sequentially recovered from a patient (no. 11) with A. fumigatus renal abscesses who was receiving voriconazole for >3 months in whom an initial urine isolate was susceptible to azole; all 5 isolates were genetically identical.

Overexpression of cdr1B (a drug efflux transporter) and an S269P mutation in hmg1 (a hydroxymethylglutaryl-CoA reductase) were identified in 4 resistant isolates but not in the initial susceptible isolate (Appendix Table 5, Figure 4), suggesting their roles involved in azole resistance (4,9). Another 2 pan-azole-resistant respiratory isolates were recovered from a patient (no. 12) who had pulmonary aspergillosis and was receiving voriconazole for 4 months. Azole-susceptible and azole-resistant isolates co-existed in this patient, which echoes the international recommendation suggesting testing multiple colonies (>5) from a single culture (1). Cyp51A overexpression and an F262 deletion in hmg1(hmg1F262del) were identified in these 2 resistant isolates. Although hmg1F262del was recently reported in azole-resistant A. fumigatus from a voriconazole-exposed patient (4), whether cyp51A overexpression and hmg1F262del act synergistically to cause resistance warrants further studies.

Finally, reduced colony sizes were observed in all 6 azole-resistant A. fumigatus isolates with wild-type cyp51A (Appendix Figure 2). Thus, attention should be paid to select colonies of various sizes for susceptibility testing from patients with azole exposure.

Overall, 4 patients harboring azole-resistant A. fumigatus with environmental mutations and 2 patients harboring azole-resistant A. fumigatus with wild-type cyp51A showed development of invasive aspergillosis, and all had aspergillosis-related deaths. High mortality rates for azole-resistant aspergillosis we observed (6/6, 100%) and for those from a previous report (10) emphasize the need for a proposed integrated algorithm for management and control of azole-resistant aspergillosis (Appendix Table 6).

In conclusion, we report a health threat that arose from clinical and environmental use of azoles; environmental use contributed at a larger and global scale. These data necessitate diagnostic stewardship in the clinic and antifungal stewardship in the environment.

Acknowledgments
We thank Cheng-Chen Cheng, Li-Fang Chen, Li-Yu Yang, and Ya-Feng Wu for laboratory assistance; Hui-Chun Chang for collecting clinical information; and the following hospitals for sharing Aspergillus spp. clinical isolates: Changhua Christian Hospital, Chi Mei Medical Center-Liouying, Chung Shan Medical University Hospital, Ditmanson Medical Foundation Chia-Yi Christian Hospital, Far Eastern Memorial Hospital, Hualien Tzu Chi Hospital, Kaohsiung Chang Gung Memorial Hospital, Kaohsiung Medical University Chung-Ho Memorial Hospital, Kaohsiung Veterans General Hospital, Mennonite Christian Hospital, National Cheng Kung University Hospital, National Taiwan University Hospital, Show Chwan Memorial Hospital, Taichung Veterans General Hospital, Tainan Sin Lau Hospital, and Taipei City Hospital Heping Fuyou Branch.
This study was supported by the National Health Research Institutes of Taiwan (grants IV-105-PP-08 and IV-106-PP-08), the Ministry of Science and Technology of Taiwan (grants MOST 105-2628-B-400-004-MY2), and Gilead Sciences, Inc. (grant IN-US-131-3918).

Y.-C.C. has received honoraria for speaking and advisory board membership from Pfizer, Gilead, Merck, and Astellas, and was involved as a steering committee member of regional education programs at Gilead (Asia CARE) and Pfizer (Medical Mycology Training Network). W.-LL. has received honoraria for speaking from Pfizer and Gilead.

About the Author
Dr. Wu is an assistant investigator and attending physician at National Health Research Institutes, Zhunan, Taiwan. Her research interests include molecular epidemiology of infectious diseases and antimicrobial drug resistance in bacterial and fungal pathogens.

References
1. Verweij PE, Ananda-Rajah M, Andes D, Arendrup MC, Brüggemann RJ, Chowdhary A, et al. International expert opinion on the management of infection caused by azole-resistant Aspergillus fumigatus. Drug Resist Updat. 2015;21-22:30–40. https://doi.org/10.1016/j.drup.2015.08.001
2. Clinical and Laboratory Standards Institute. Reference method for broth dilution antifungal susceptibility testing of filamentous fungi: approved standard, 2nd ed. M38–A2. Wayne (PA): The Institute; 2008.
3. Wang HC, Huang JC, Lin YH, Chen YH, Hsieh MI, Choi PC, et al. Prevalence, mechanisms and genetic relatedness of the human pathogenic fungus Aspergillus fumigatus exhibiting resistance to medical azoles in the environment of Taiwan. Environ Microbiol. 2018;20:270–80. https://doi.org/10.1111/1462-2920.13988
4. Hagiwara D, Arai T, Takahashi H, Kusuya Y, Watanabe A, Kamei K. Non-cyp51A azole-resistant Aspergillus fumigatus isolates with mutation in HMG-CoA reductase. Emerg Infect Dis. 2018;24:1889–97. https://doi.org/10.3201/eid2410.180730
5. Arendrup MC. Update on antifungal resistance in Aspergillus and Candida. Clin Microbiol Infect. 2014;20(Suppl 6):42–8. https://doi.org/10.1111/1469-0691.12513
6. Chowdhary A, Sharma C, van den Boom M, Yntema JB, Hagen F, Verweij PE, et al. Multi-azole-resistant Aspergillus fumigatus in the environment in Tanzania. J Antimicrob Chemother. 2014;69:2979–83. https://doi.org/10.1093/jac/dkt075
7. Chen YC, Lai MH, Wu CY, Lin TC, Cheng AH, Yang CC, et al. The genetic structure, virulence, and fungicide sensitivity of Fusarium fujikuroi in Taiwan. Phytopathology. 2016;106:624–35. https://doi.org/10.1094/PHYTO-11-15-0285-R
8. Chen Y, Li Z, Han X, Tian S, Zhao J, Chen F, et al. Elevated MIC values of imidazole drugs against Aspergillus fumigatus isolates with TR-/L981I/S297T/F495I mutation. Antimicrob Agents Chemother. 2018;62:e01549-17. https://doi.org/10.1128/AAC.01549-17
9. Fraczek MG, Bromley M, Buied A, Moore CB, Rajendran R, Rautemaa R, et al. The cdr1B efflux transporter is associated with non-cyp51A-mediated itraconazole resistance in Aspergillus fumigatus. J Antimicrob Chemother. 2013;68:1486–96. https://doi.org/10.1093/jac/dkt075
10. van der Linden JW, Snelders E, Kampina GA, Rijnders BJ, Mattsson E, Debets-Ossenkopp YJ, et al. Clinical implications of azole resistance in Aspergillus fumigatus, The Netherlands, 2007–2009. Emerg Infect Dis. 2011;17:1846–54. https://doi.org/10.3201/eid1710.110226

Address for correspondence: Yee-Chun Chen, Department of Internal Medicine, National Taiwan University Hospital, No. 7 Chung-Shan South Rd, Taipei 10002, Taiwan; email: yeechunchen@gmail.com

Knowledge of Infectious Disease Specialists Regarding Aspergillosis Complicating Influenza, United States

Mitsuru Toda, Susan E. Beekmann, Philip M. Polgreen, Tom M. Chiller, Brendan R. Jackson, Karlyn D. Beer

Author affiliations: Centers for Disease Control and Prevention, Atlanta, Georgia, USA (M. Toda, T.M. Chiller, B.R. Jackson, K.D. Beer); University of Iowa, Iowa City, Iowa, USA (S.E. Beekmann, P.M. Polgreen)

DOI: https://doi.org/10.3201/eid2604.190953

In an online survey, we found that nearly one fifth of physicians in the United States who responded had seen or heard about a case of invasive pulmonary aspergillosis after severe influenza at their institution. However, <10% routinely used galactomannan testing to test for this fungus in patients with severe influenza.

Invasive pulmonary aspergillosis (IPA) occurs primarily among immunocompromised patients with a history of organ or stem cell transplantation, chemotherapy, or immunosuppressive medications. However, a multicenter retrospective study in the Netherlands and Belgium suggested that patients...
Multicenter Study of Azole-Resistant Aspergillus fumigatus Clinical Isolates, Taiwan

Appendix

Appendix Table 1. Details of participating hospitals, antifungal susceptibility testing, and isolate collection for analysis of azole-resistant Aspergillus fumigatus clinical isolates, Taiwan*

| Hospital | Location | Period of collection | Susceptibility testing method† | Isolates before June 2017 | Isolates during June 2017–March 2018 |
|----------|----------|----------------------|-------------------------------|--------------------------|-------------------------------------|
| Chi-Mei Medical Center, Luiying (CMMC) | Southern | 2015 Feb–2018 Mar | CLSI M38-A2 | Screening azole agar plates; confirmed by CLSI M38-A2 |
| National Cheng-Kung University Hospital (NCKUH) | Southern | 2011 Aug–2018 Mar | CLSI M38-A2 | Screening azole agar plates; confirmed by CLSI M38-A2 |
| National Taiwan University Hospital (NTUH) | Northern | 2012 Feb–2018 Mar | YeastOne | Screening azole agar plates; confirmed by CLSI M38-A2 |

TSARM hospitals
- Changhua Christian Hospital | Central | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Dihmanson Medical Foundation Chia-Yi Christian Hospital | Southern | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Hualien Tzu Chi Hospital | Eastern | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Kaohsiung Medical University Chung-Ho Memorial Hospital | Southern | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Kaohsiung Veterans General Hospital | Southern | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Show Chwan Memorial Hospital | Central | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Taichung Veterans General Hospital | Central | 2016 Jul–Sep | CLSI M38-A2 | Ni |
- Tainan Sin Lau Hospital | Southern | 2016 Jul–Sep | CLSI M38-A2 | Ni |

This study was approved by the Institutional Review Boards of the National Health Research Institutes (no. EC1040502-E and EC1050307) and participating hospitals: CMMC (10607-L01 and 10711-L03), NCKUH (B-ER-101–342), and NTUH (201605098RIPA). Ni, no isolate; TSARM, Taiwan Surveillance of Antimicrobial Resistance of Molds.

†A. fumigatus sensu stricto was identified on the basis of morphologic characteristics, growth at 50°C, and sequence analyses of the internal transcribed spacer region and calmodulin gene (1). For isolates from CMMC, NCKUH, and TSARM hospitals, MICs of antifungal agents were determined by using the Clinical and Laboratory Standards Institute (CLSI) M38-A2 method. The MICs for the isolates from NTUH were determined by using the Sensititre YeastOne method (Trek Diagnostic Systems, http://www.trekds.com); isolates with any of the following MIC values (i.e., ≥2, ≥2, and ≥0.25 μg/mL of itraconazole, voriconazole, and posaconazole, respectively) were reexamined by using the CLSI method. For isolates collected during June 2017–March 2018, azole resistance was screened by using azole-containing agar plates. In brief, the conidia of these isolates were directly inoculated onto 3 Sabouraud dextrose agar plates supplemented with itraconazole (2 μg/mL), voriconazole (1 μg/mL), or posaconazole (0.25 μg/mL), and incubated at 37°C. Colonies that grew after 2–4 d on any of the azole-containing agar plates were selected for the MIC determination by using the CLSI method.
Table 2. Laboratory characteristics of *Aspergillus fumigatus* clinical isolates from 12 patients with aspergillosis, Taiwan*

| Patient no. | Strain no. | Year | Cyp51A mutation | MIC or MIC\(50/\text{MIC}_{\text{iso}}\) (range), μg/mL | AMB | ITR | VRC | POS | ISA | DFC | TBC | PRC |
|-------------|------------|------|----------------|--------------------------------------------------|-----|-----|-----|-----|-----|-----|-----|-----|
| 1           | B44        | 2012 (d0) | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 16 >32 |     |     |     |     |     |     |     |     |
| 2           | B51        | 2012 (d2) | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 16 >32 |     |     |     |     |     |     |     |     |
| 3           | D007       | 2014   | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 4           | E071       | 2015 (d0) | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 5           | E073       | 2015 (d3) | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 6           | E074       | 2015 (d4) | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 7           | S05–11     | 2016   | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 8           | S05–12     | 2016   | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 9           | S05–13     | 2016   | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 10          | S05–14     | 2016   | TR\(^{u}/L98H/S297T/F495l\) | 0.5 >16 2 1 >16 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 11          | YL1        | 2014 (d0) | Polymorphisms‡ | 0.5 0.25 0.25 1 4 4 0.25 |     |     |     |     |     |     |     |     |
| 12          | YL3        | 2014 (d10) | Polymorphisms‡ | 0.5 >16 8 1 8 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 13          | YL4        | 2015 (d134) | Polymorphisms‡ | 0.5 >16 8 1 8 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 14          | YL5        | 2015 (d165) | Polymorphisms‡ | 0.5 >16 2–4 0.5 2 16 16 1 |     |     |     |     |     |     |     |     |
| 15          | YL6        | 2015 (d185) | Polymorphisms‡ | 0.5 >16 8 1 8 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 16          | g054m      | 2016 (d0) | Wild-type | 0.12 >16 8 1 8 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 17          | g054L      | 2016 (d0) | Wild-type | 0.5 0.5 0.5 0.06 0.5 1 4 0.25 |     |     |     |     |     |     |     |     |
| 18          | g057m      | 2016 (d2) | Wild-type | 0.12 >16 4 1 8 >32 >32 >32 |     |     |     |     |     |     |     |     |
| 19          | g057L      | 2016 (d2) | Wild-type | 0.5 0.5 0.5 0.06 0.5 1 4 0.25 |     |     |     |     |     |     |     |     |

Azole-resistant isolates§ except YL1, g054L, and g057L

*AMB, amphotericin B; ATCC, American Type Culture Collection; DFC, difenoconazole; ISA, isavuconazole; ITR, itraconazole; NT, not done; POS, posaconazole; PRC, prochloraz; TBC, tebuconazole; VRC, voriconazole.*

†d0 and d1 indicate day of and n days after the first isolation of *A. fumigatus*, respectively.

‡*Candida parapsilosis* ATCC 22019 and *A. fumigatus* ATCC MYA 3626 were used as quality control and reference strains.

§Because of good essential agreement in the azole MIC values between the CLSI and European Committee on Antimicrobial Susceptibility Testing (EUCAST) methods (2) and the lack of established CLSI clinical breakpoints for *A. fumigatus*, the MIC interpretive criteria for resistance in this study followed the EUCAST clinical breakpoints (i.e., >2, >2, >0.25, and >1 μg/mL) for itraconazole, voriconazole, posaconazole, and isavuconazole, respectively (3). The drugs for susceptibility testing were obtained from Sigma-Aldrich (https://www.sigmaaldrich.com) (AMB, ITR, VRC, POS, DFC, and PRC), Toronto Research Chemicals (https://www.tr-canada.com) (ISA), and Chem Service (https://www.chemservice.com) (TBC).

¶These isolates have F46Y/G89S/L136A/D137N/L140F/L141F/L143W/L281F/L284F/L285F/L350F/L351F/L357F/L359F/L427K/C454C polymorphisms.

#The MICs of 200 azole-susceptible isolates were used. Only 62 isolates were tested for MICs for DFC and TBC.
### Appendix Table 3. Clinical characteristics of 12 patients harboring azole-resistant *Aspergillus fumigatus* isolates, Taiwan*

| Patient no. | Strain no. | Age, y/sex | Concurrent condition | Sample | Aspergillus disease† | Previousazole exposure | Sequential antifungal treatment (d) | Outcome | IA-related death |
|-------------|------------|------------|----------------------|--------|----------------------|------------------------|------------------------------------|---------|-----------------|
| 1‡‡         | B44, B51   | 66/F       | DM, HCV/cirrhosis,  | Sputum | IPA, unclassified     | No                     | VRC (1), AMB (3)                | Died    | Yes             |
| 2           | D007       | 49/M       | AML, peritonitis     | Nasal swab | Colonization         | POS/VR C (2 wk)        | MCF (33)                          | No      |                 |
| 3           | E071, E074; E073 | 64/F | SLE, ESRD, bacterial septicemia, meningocencephalitis | Pleural effusion; sputum | Proven IPA with empyema | No                     | VRC (1), LAMB (3)                | Died    | Yes             |
| 4           | S05–319 A31 | 88/F       | Lung cancer, COPD, bronchiectasis | Sputum | Colonization         | No                     | No                                 | Died    | No              |
| 5‡‡         | S05–122    | 90/F       | COPD, steroid use, bronchiectasis | Sputum | Probable IPA         | No                     | AMB (3), POS (10), VRC (11)       | Alive   | No              |
| 7           | C03–004    | –          | –                     | Sputum | –                   | –                      | –                                 | Died    | No              |
| 8           | S07–008    | –          | –                     | Ear    | –                   | –                      | –                                 | Died    | No              |
| 9           | S05–205    | 76/F       | COPD, steroid use, bronchiectasis, DM B cell lymphoma, COPD, HCV, CAD HIV/AIDS | Sputum | Colonization         | No                     | No                                 | Alive   | No              |
| 10          | S05–322    | 74/M       | –                     | Sputum | Colonization         | No                     | No                                 | Alive   | No              |
| 11          | YL1;§ YL3, YL4, YL5, YL6 | 36/M | –                     | Urine; PCN | Proven IA, (renal abscess) | VRC (3 mo)                          | VRC (93), VRC/CAS (43), LAMB (44), LAMB/SFC (25), LAMB/AND (24), AMB (18), POS (21), AMB (13), AMB/AND (8), AMB (5), AMB/AND (1), AND (4), AMB/AND (2) | Died    | Yes             |
| 12          | g054m, g054L§, g057m, g057L§ | 39/M | MDS with RAEB-T, status post-HSCT with GVHD, bacterial septicemia | Sputum; BAL | Probable IPA         | VRC (4 mo)                          | Died    | Yes             |

*AMB, conventional amphotericin B; AML, acute myeloid leukemia; AND, anidulafungin; BAL, bronchoalveolar lavage; CAD, coronary artery disease; CAS, caspofungin; COPD, chronic obstructive pulmonary disease; DM, diabetes mellitus; ESRD, end-stage renal disease; 5FC, flucytosine; GVHD, graft versus host disease; HCV, hepatitis C virus infection; HSCT, allogeneic hematopoietic stem cell transplantation; IA, invasive aspergillosis; IPA, invasive pulmonary aspergillosis; LAmB, liposomal amphotericin B; MCF, micafungin; MDS with RAEB-T, myelodysplastic syndrome with refractory anemia and excess blast in transformation; PCN, percutaneous nephrostomy; POS, posaconazole; SLE, systemic lupus erythematosus; VRC, voriconazole; –, data not available.

†Clinical data for patients harboring azole-resistant *A. fumigatus* were reviewed, and IA was classified according to the EORTC/MSG definition (4).

‡Three isolates (A31, B44, and B51) from 2 patients have been described in our previous report (5).

§YL1, g054L, and g057L were azole susceptible.
## Appendix Table 4. References for 38 oversea *Aspergillus fumigatus* strains included in microsatellite-based phylogenetic analysis during a multicenter study of azole-resistant *Aspergillus fumigatus* clinical isolates, Taiwan

| Strain     | Reference |
|------------|-----------|
| C485       | (6)       |
| C96        | (6)       |
| E2619      | (6)       |
| 20643.023  | (7)       |
| 20684.007  | (7)       |
| E1001      | (6)       |
| 20677.079  | (7)       |
| 2087 m1341.17–06–2012 | (8) |
| 2091 m1428.01–07–2012 | (8) |
| C94        | (6)       |
| 09441/7/50 | (8)       |
| Case 2–90d | (9)       |
| Japan Dec 2013 | (10) |
| The Netherlands 7 | (10,11) |
| 20643.017  | (7)       |
| 2005–456307L | (6) |
| OKH50      | (12)      |
| 04–202165  | (13)      |
| F2126      | (6)       |
| 1042/09    | (14)      |
| 14–148–2460| (6)       |
| 2107m1974.23–09–2012 | (8) |
| A12519     | (8)       |
| R2–07–1_R  | (6)       |
| E454       | (15)      |
| Case 1–7d  | (9)       |
| Myc-2008–002 nr.42 | (14) |
| Case 3–6d  | (9)       |
| CF/NL2992  | (8)       |
| Tanzania   | (11)      |
| CF/NL0682  | (8)       |
| VPCI851/El/12/2/a3 | (8) |
| CF/NL0645  | (11)      |
| The Netherlands 2 | (10,11) |
| The Netherlands 3 | (10,11) |
| C195       | (7)       |
| 12–90032258| (13)      |
| Case 4–36d | (9)       |

## Appendix Table 5. Gene substitutions in azole-resistant *Aspergillus fumigatus* isolates and control strains, Taiwan*

| Strain no. | Azole susceptibility (mechanism) | hapE | srbA | hmg1 | erg6 |
|------------|----------------------------------|------|------|------|------|
| ATCC MYA-3626 | S | – | V37D | – | – |
| ATCC 16903   | S | – | V37D, S82P | H564Y | – |
| F2509        | S | – | V37D | – | – |
| F02411       | S | – | A865V | P212S, H564Y | – |
| YL1          | S | – | E957D | P212S, S269P, H564Y | F186V |
| YL3          | R | – | E957D | P212S, S269P, H564Y | F186V |
| YL4          | R | – | E957D | P212S, S269P, H564Y | F186V |
| YL5          | R | – | E957D | P212S, S269P, H564Y | – |
| YL6          | R | – | E957D | P212S, S269P, H564Y | F186V |
| g054m        | R | – | A865V | F262_del, H564Y | – |
| g057m        | R | – | A865V | F262_del, H564Y | – |
| g054L        | S | – | V37D | – | – |
| g057L        | S | – | V37D | – | – |
| B44          | R (TRu/L98H/S297T/F495I) | – | V37D | H564Y | – |
| A31          | R (TRu/L98H) | – | V37D | – | – |

*ATCC, American Type Culture Collection; del, deletion; R, resistant; S, susceptible; –, wild type.
†The hapE, hmg1, erg6 and srbA genes were sequenced as described (16–18). The hapE and srbA sequences were compared with those of *A. fumigatus* f293; the hmg1 and erg6 sequences were compared with those of *A. fumigatus* A1163.
### Appendix Table 6. Proposed integrated algorithm for management and control of azole-resistant aspergillosis, Taiwan*

| Category                      | Response                                                                                                                                                                                                                                                                                                                                 |
|-------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| **Academic society**          | Incorporate the issue of azole resistance into national guidelines for management of aspergillosis; hold educational programs to improve diagnosis and treatment for azole-resistant aspergillosis.                                                                                                                                          |
| **Laboratory personnel**      | Identify clinically relevant *Aspergillus* isolates at the species complex level and confirm *A. fumigatus* by thermotolerance test (growth at 50°C) (19); screen for azole resistance with azole agar plates for clinically relevant *A. fumigatus* isolates and screen multiple colonies (<5 colonies) from a single specimen (19); for colonies grown on any azole agar plate, perform azole MIC testing by using reference CLSI or EUCAST methods or an alternative Sensititre YeastOne assay (19,20); If MIC testing is not available, refer isolates to a mycology reference laboratory (19); prompt notification of the clinical team if azole-resistant *A. fumigatus* is suspected and identified. |
| **Physicians**                | Be familiar with patient risk factors for invasive aspergillosis; obtain clinical specimens for fungal culture as possible; select empirical antifungal agents according to the updated local prevalence rate of azole resistance (21); antifungal susceptibility testing is recommended for *A. fumigatus* isolates from invasive diseases and should be repeated on later isolates if infection persists despite treatment (21); be aware of the possibility of azole-resistance in patients unresponsive to azole treatment; consider amphotericin B–based or azole/echinocandin combination therapy for azole-resistant aspergillosis (19,21). |
| **Hospital environment**      | Segregate patients from construction or renovation, potted plants, and flowers in wards and patients’ room (22); control the airborne dissemination of fungal spores, (e.g., barriers, containment, air handling, HEPA filters, sealed windows, sealing the area of construction or renovation activities if possible) (23). |
| **Reference mycology laboratory** | Identify *Aspergillus* isolates to the species level by molecular methods; confirm antifungal susceptibility of *Aspergillus* isolates with reference CLSI or EUCAST methods; perform periodic reference MIC testing of isolates of *A. fumigatus* complex (≥100 isolates) (19); sequence cyp51A genes in resistant isolates to determine the nature and trends in cyp51A mutation distribution (19); establish molecular typing methods; collect strains. |
| **Scientists and plant pathologists** | Identify the key azole fungicides that select azole-resistant *Aspergillus*; propose better fungicide application strategies to minimize resistance development (24).                                                                                                                                 |
| **Agricultural authority**    | Include azole fungicides that select azole-resistant isolates into national pesticide risk reduction programs; advise farmers to reduce culprit azole fungicide use by in rotation with alternative fungicides with different modes of action.                                                                                       |
| **Governance**                | Update and evaluate the global situation; accredit national mycology reference laboratories; implement antifungal stewardship programs in agriculture in addition to hospitals and animal husbandry to achieve the One Health goal (24).                                                                                                                      |

*CLSI, Clinical and Laboratory Standards Institute; EUCAST, European Committee on Antimicrobial Susceptibility Testing.
Appendix Figure 1. Distribution of participating hospitals and isolate collection areas for *Aspergillus fumigatus*, Taiwan. Values in parentheses indicate no. *A. fumigatus* isolates/no. patients and no. azole-resistant *A. fumigatus* isolates/no. patients.
Appendix Figure 2. A) Radial growth of *Aspergillus fumigatus* isolates on Sabouraud dextrose agar plates at 35°C, Taiwan. The radius of the growing colony was measured after 72 hours of incubation. Values are the mean diameter of triplicate samples. Error bars indicate SD. Colonies of B) YL1, YL3, YL4, YL5, and YL6 from patient 11 and C) g054m, g054L, g057m, and g057L from patient 12 observed at 72 hours. C, clinical isolate; E, environmental isolate; R, azole-resistant; S, azole-susceptible; ↑, overexpression.
Appendix Figure 3. Annual sales of azole fungicides in Taiwan, 2003–2016. Annual sales of imidazole fungicides (imazalil and prochloraz) and triazole fungicides (bromuconazole, difenoconazole, epoxiconazole, propiconazole, and tebuconazole) are shown according to data derived from Domestic Manufacturers Production and Sale of Pesticides published by the Taiwan Crop Protection Industry Association (25).
Appendix Figure 4. mRNA expression levels of A) a drug efflux transporter gene, *cdr1B*, and B) *cyp51A* in *Aspergillus fumigatus* isolates, Taiwan. Expression levels were normalized to β-tubulin levels and compared with those in *A. fumigatus* ATCC MYA-3626. Error bars indicate SD. Results for the *cyp51B* gene and other transporter genes (*AfuMDR1, AfuMDR2, AfuMDR3, AfuMDR4, atrF, and MFS56*) were inconclusive and are not shown.

ATCC, American Type Culture Collection; ITR, itraconazole; VRC, voriconazole.
1. Balajee SA, Houbraken J, Verweij PE, Hong SB, Yaghuchi T, Varga J, et al. *Aspergillus* species identification in the clinical setting. Stud Mycol. 2007;59:39–46. PubMed [https://doi.org/10.3114/sim.2007.59.05](https://doi.org/10.3114/sim.2007.59.05)

2. Pfaller M, Boyken L, Hollis R, Kroeger J, Messer S, Tendolkar S, et al. Comparison of the broth microdilution methods of the European Committee on Antimicrobial Susceptibility Testing and the Clinical and Laboratory Standards Institute for testing itraconazole, posaconazole, and voriconazole against *Aspergillus* isolates. J Clin Microbiol. 2011;49:1110–2. PubMed [https://doi.org/10.1128/JCM.02432-10](https://doi.org/10.1128/JCM.02432-10)

3. EUCAST. Subcommitte on Antimicrobial Susceptibility Testing (EUCAST-AFST). Clinical breakpoints for fungi, version 9 [cited 2020 Jan 25]. http://www.eucast.org/astoffungi клиничных границ для грибов

4. De Pauw B, Walsh TJ, Donnelly JP, Stevens DA, Edwards JE, Calandra T, et al.; European Organization for Research and Treatment of Cancer/Invasive Fungal Infections Cooperative Group; National Institute of Allergy and Infectious Diseases Mycoses Study Group (EORTC/MSG) Consensus Group. Revised definitions of invasive fungal disease from the European Organization for Research and Treatment of Cancer/Invasive Fungal Infections Cooperative Group and the National Institute of Allergy and Infectious Diseases Mycoses Study Group (EORTC/MSG) Consensus Group. Clin Infect Dis. 2008;46:1813–21. PubMed [https://doi.org/10.1086/588660](https://doi.org/10.1086/588660)

5. Wu CJ, Wang HC, Lee JC, Lo HJ, Dai CT, Chou PH, et al. Azole-resistant *Aspergillus fumigatus* isolates carrying TR34/L98H mutations in Taiwan. Mycoses. 2015;58:544–9. PubMed [https://doi.org/10.1111/myc.12354](https://doi.org/10.1111/myc.12354)

6. Chen Y, Lu Z, Zhao J, Zou Z, Gong Y, Qu F, et al. Epidemiology and molecular characterizations of azole resistance in clinical and environmental *Aspergillus fumigatus* isolates from China. Antimicrob Agents Chemother. 2016;60:5878–84. PubMed [https://doi.org/10.1128/AAC.01005-16](https://doi.org/10.1128/AAC.01005-16)

7. Deng S, Zhang L, Ji Y, Verweij PE, Tsui KM, Hagen F, et al. Triazole phenotypes and genotypic characterization of clinical *Aspergillus fumigatus* isolates in China. Emerg Microbes Infect. 2017;6:e109. PubMed [https://doi.org/10.1038/emi.2017.97](https://doi.org/10.1038/emi.2017.97)

8. Steinmann J, Hamprecht A, Vehreschild MJ, Cornely OA, Buchheidt D, Spiess B, et al. Emergence of azole-resistant invasive aspergillosis in HSCT recipients in Germany. J Antimicrob Chemother. 2015;70:1522–6. PubMed [https://doi.org/10.1093/jac/dku566](https://doi.org/10.1093/jac/dku566)

9. Astvad KM, Jensen RH, Hassan TM, Mathiasen EG, Thomsen GM, Pedersen UG, et al. First detection of TR46/Y121F/T289A and TR34/L98H alterations in *Aspergillus fumigatus* isolates from azole-naive patients in Denmark despite negative findings in the environment. Antimicrob Agents Chemother. 2014;58:5096–101. PubMed [https://doi.org/10.1128/AAC.02855-14](https://doi.org/10.1128/AAC.02855-14)
10. Hagiwara D, Takahashi H, Fujimoto M, Sugahara M, Misawa Y, Gono T, et al. Multi-azole resistant *Aspergillus fumigatus* harboring Cyp51A TR46/Y121F/T289A isolated in Japan. J Infect Chemother. 2016;22:577–9. PubMed https://doi.org/10.1016/j.jiac.2016.01.015

11. Chowdhary A, Sharma C, van den Boom M, Yntema JB, Hagen F, Verweij PE, et al. Multi-azole-resistant *Aspergillus fumigatus* in the environment in Tanzania. J Antimicrob Chemother. 2014;69:2979–83. PubMed https://doi.org/10.1093/jac/dku259

12. Toyotome T, Hagiwara D, Kida H, Ogi T, Watanabe A, Wada T, et al. First clinical isolation report of azole-resistant *Aspergillus fumigatus* with TR34/L98H-type mutation in Japan. J Infect Chemother. 2017;23:579–81. PubMed https://doi.org/10.1016/j.jiac.2016.12.004

13. Kidd SE, Goeman E, Meis JF, Slavin MA, Verweij PE. Multi-triazole-resistant *Aspergillus fumigatus* infections in Australia. Mycoses. 2015;58:350–5. PubMed https://doi.org/10.1111/myc.12324

14. Chowdhary A, Kathuria S, Xu J, Sharma C, Sundar G, Singh PK, et al. Clonal expansion and emergence of environmental multiple-triazole-resistant *Aspergillus fumigatus* strains carrying the TR34/L98H mutations in the cyp51A gene in India. PLoS One. 2012;7:e52871. PubMed https://doi.org/10.1371/journal.pone.0052871

15. Ahmad S, Khan Z, Hagen F, Meis JF. Occurrence of triazole-resistant *Aspergillus fumigatus* with TR34/L98H mutations in outdoor and hospital environment in Kuwait. Environ Res. 2014;133:20–6. PubMed https://doi.org/10.1016/j.envres.2014.05.009

16. Camps SM, Dutilh BE, Arendrup MC, Rijs AJ, Snelders E, Huynen MA, et al. Discovery of a *hapE* mutation that causes azole resistance in *Aspergillus fumigatus* through whole genome sequencing and sexual crossing. PLoS One. 2012;7:e50034. PubMed https://doi.org/10.1371/journal.pone.0050034

17. Hagiwara D, Arai T, Takahashi H, Kusuya Y, Watanabe A, Kamei K. Non-cyp51A azole-resistant *Aspergillus fumigatus* isolates with mutation in HMG-CoA reductase. Emerg Infect Dis. 2018;24:1889–97. PubMed https://doi.org/10.3201/eid2410.180730

18. Wang HC, Huang JC, Lin YH, Chen YH, Hsieh MI, Choi PC, et al. Prevalence, mechanisms and genetic relatedness of the human pathogenic fungus *Aspergillus fumigatus* exhibiting resistance to medical azoles in the environment of Taiwan. Environ Microbiol. 2018;20:270–80. PubMed https://doi.org/10.1111/1462-2920.13988

19. Ullmann AJ, Aguado JM, Arikante-Ladaghi S, Denning DW, Groll AH, Lagrou K, et al. Diagnosis and management of *Aspergillus* diseases: executive summary of the 2017 ESCMID-ECMM-ERS guideline. Clin Microbiol Infect. 2018;24(Suppl 1):e1–38. PubMed https://doi.org/10.1016/j.cmi.2018.01.002
20. Wang HC, Hsieh MI, Choi PC, Wu CJ. Comparison of the Sensititre YeastOne and CLSI M38–A2 microdilution methods in determining the activity of amphotericin B, itraconazole, voriconazole, and posaconazole against *Aspergillus* species. J Clin Microbiol. 2018;56:e00780–18. PubMed https://doi.org/10.1128/JCM.00780-18

21. Verweij PE, Ananda-Rajah M, Andes D, Arendrup MC, Brüggemann RJ, Chowdhary A, et al. International expert opinion on the management of infection caused by azole-resistant *Aspergillus fumigatus*. Drug Resist Updat. 2015;21-22:30–40. PubMed https://doi.org/10.1016/j.drup.2015.08.001

22. Freifeld AG, Bow EJ, Sepkowitz KA, Boeckh MJ, Ito JI, Mullen CA, et al.; Infectious Diseases Society of America. Clinical practice guideline for the use of antimicrobial agents in neutropenic patients with cancer: 2010 update by the Infectious Diseases Society of America. Clin Infect Dis. 2011;52:e56–93. PubMed https://doi.org/10.1093/cid/cir073

23. Kanamori H, Rutala WA, Sickbert-Bennett EE, Weber DJ. Review of fungal outbreaks and infection prevention in healthcare settings during construction and renovation. Clin Infect Dis. 2015;61:433–44. PubMed https://doi.org/10.1093/cid/civ297

24. Chowdhary A, Meis JF. Emergence of azole resistant *Aspergillus fumigatus* and One Health: time to implement environmental stewardship. Environ Microbiol. 2018;20:1299–301. PubMed https://doi.org/10.1111/1462-2920.14055

25. Taiwan Crop Protection Industry Association. Domestic manufacturers production and sale of pesticides [in Chinese]. Taipei: Taiwan Crop Protection Industry Association, 1962–2016.