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Azole resistance in *Aspergillus fumigatus*. The first 2-year's Data from the Danish National Surveillance Study, 2018–2020

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Abstract

**Background:** Azole resistance complicates treatment of patients with invasive aspergillosis with an increased mortality. Azole resistance in *Aspergillus fumigatus* is a growing problem and associated with human and environmental azole use. Denmark has a considerable and highly efficient agricultural sector. Following reports on environmental azole resistance in *A. fumigatus* from Danish patients, the ministry of health requested a prospective national surveillance of azole-resistant *A. fumigatus* and particularly that of environmental origin.

**Objectives:** To present the data from the first 2 years of the surveillance programme.

**Methods:** Unique isolates regarded as clinically relevant and any *A. fumigatus* isolated on a preferred weekday (background samples) were included. EUCAST susceptibility testing was performed and azole-resistant isolates underwent cyp51A gene sequencing.

**Results:** The azole resistance prevalence was 6.1% (66/1083) at patient level. The TR\(_{34}/L98H\) prevalence was 3.6% (39/1083) and included the variants TR\(_{34}/L98H\),...
1 | INTRODUCTION

Azole resistance in *Aspergillus fumigatus* due to the specific molecular mechanisms TR\textsubscript{34}/L98H or TR\textsubscript{34}/Y121F/T289A has been reported from all seven continents except Antarctica.\textsuperscript{1} These mechanisms are found in environmental *A. fumigatus* isolates and in isolates from azole naïve as well as from exposed patients. Azole resistance can also arise in *A. fumigatus* in patients receiving long-term azole treatment.\textsuperscript{2} Most resistant isolates harbour mutations in cyp51A, which encodes the azole target 14α-sterol-demethylase, essential for ergosterol biosynthesis.\textsuperscript{2} However, azole resistance has also been ascribed to efflux pumps and other non-cyp51A-mediated resistance mutations.\textsuperscript{2,3}

In Denmark, the first isolation of TR\textsubscript{34}/L98H and TR\textsubscript{34}/Y121F/T289A were from clinical samples in 2007 and 2014, respectively.\textsuperscript{4,5} Subsequently, TR\textsubscript{34}/L98H and TR\textsubscript{34}/Y121F/T289A have also been found in environmental samples since 2009 and 2019, respectively.\textsuperscript{6,7} Moreover, an increase in the prevalence of azole resistance among Danish cystic fibrosis (CF) patients was found over a 10-year period.\textsuperscript{8}

Clinical manifestations with *Aspergillus* vary according to patient group. In the CF population, *Aspergillus* occurs most often as part of colonisation, allergic bronchopulmonary aspergillosis (ABPA) and bronchitis.\textsuperscript{9} ABPA is also a well-known condition in patients with asthma.\textsuperscript{10} Invasive aspergillosis mainly occurs in patients who are immunosuppressed and chronic aspergillosis in patients with impaired lung tissue architecture. Azoles are the drugs of choice in the management of aspergillosis.\textsuperscript{10} Voriconazole and isavuconazole are first choice in invasive aspergillosis,\textsuperscript{11,12} itraconazole (and voriconazole) in chronic aspergillosis\textsuperscript{13} and posaconazole as prophylaxis or salvage treatment, but with potential future broadening of its licensed indication due to non-inferior to voriconazole for primary therapy.\textsuperscript{10,12,14} At this point, azoles are the only antifungal agents against aspergillosis for oral administration. The emergence of azole resistance complicates patient treatment, and invasive aspergillosis with azole resistance is associated with an inferior outcome compared to invasive aspergillosis with a susceptible strain.\textsuperscript{15,16}

An international expert opinion suggested that when the environmental resistance rate exceeds 10% in a region, the initial treatment for invasive aspergillosis should be either liposomal amphotericin B or voriconazole combined with an echinocandin.\textsuperscript{17} This recommendation was based on two observations. First, the significantly increased mortality found in patients who received voriconazole initially for invasive aspergillosis due to resistant *A. fumigatus*\textsuperscript{15,16} and second, the superior activity of voriconazole for those with susceptible *A. fumigatus* (~70% survival vs. 55% for amphotericin B and 50% for echinocandins).\textsuperscript{17,18} This approach requires reliable epidemiological data on the prevalence of azole resistance in *A. fumigatus* due to the environmental route of acquisition (which may occur even in azole naïve patients) and medical route (which is limited to the azole exposed patient population).

The Danish national surveillance programme on azole resistance was established in 2018 upon request from the ministry of health due to the rising concerns for azole resistance of environmental origin. The objective was to determine the prevalence of azole-resistant *A. fumigatus* isolates among *A. fumigatus* colonised and infected patients in Denmark and determine the underlying resistance mechanism. We present data from the first 2 years of the surveillance.

2 | METHODS

2.1 | Organisation of the national surveillance programme of azole-resistant *A. fumigatus*

The surveillance programme was initiated on October 1st 2018 with participation from all 10 Danish clinical microbiological departments. Inclusion criteria were as follows: (a) unique
A. fumigatus isolates that were regarded clinically significant and (b) any A. fumigatus isolated on a preferred weekday (regardless of clinical significance) were included when marked ‘Background’. The adherence to the inclusion criteria varied. Six departments followed the instructions with ‘Background’ samples with a potential uncertainty of whether the isolate represented a clinical condition with aspergillosis or a contamination. Two departments sent all isolates, and two departments sent only clinically relevant isolates. The centres are quite in-homogeneous in patient up-take and size of uptake area. For example, three are district hospitals (Veje, Sønderborg and Esbjerg), two hold CF-centres (AUH and RH) and one has a centre for chronic pulmonary aspergillosis (OUH). Isolates from the same patients were deemed unique if one of the following conditions were met: (1) when sampled more than 30 days apart, (2) if the isolate had a different susceptibility or (3) a different molecular resistance mechanism.

The clinical microbiological departments referred isolates to the reference mycological laboratory at Statens Serum Institut prospectively. Some departments performed species identification of moulds to the species level and only referred A. fumigatus while others referred all Aspergillus isolates or all mould isolates for species identification and susceptibility testing. One department at Aarhus University Hospital (AUH) performed EUCAST susceptibility testing (E. Def 10.1 and E. Def 9.3.1 as described below) of most A. fumigatus isolates locally and referred the MIC data and all resistant isolates for cyp51A sequencing (and confirmatory MIC determination) thus ensuring that all A. fumigatus isolates from AUH were included in the data analysis. Monthly reports on referred isolates were communicated to the participating laboratories to motivate and ensure adherence to the surveillance programme.

2.2 | Culturing and species identification

Primary cultures were performed using Sabouraud glucose agar (SSI Diagnostika or bioMérieux) or YGC agar (yeast glucose agar; SSI Diagnostika) with incubation at 35–37°C for 5 days. Species identification included classical techniques including macro- and micromorphology and thermotolerance testing supplemented with MALDI-TOF MS and β-tubulin sequencing as needed as previously described.8 Only A. fumigatus sensu stricto isolates were included in the surveillance.

2.3 | Susceptibility testing and target gene sequencing

A. fumigatus isolates underwent screening for azole resistance following the EUCAST E. Def 10.1 method using VIPcheck azole agar plates (Mediaproducts BV).19 Screening positive isolates underwent EUCAST E. Def 9.3.1 susceptibility testing.20 For consistency, the MIC values from the reference laboratory were used throughout. The applied antifungal concentration ranges for the MIC testing varied slightly during the study period. Susceptibility classification was performed according to the current EUCAST breakpoints v. 10.0.21 Cyp51A sequencing was performed for isolates classified as azole resistant to at least one azole. The promoter and full coding region of the cyp51A gene were sequenced as previously described,5 with the exception that for Sanger sequencing, OF was replaced with a new primer 1F (5’TGGCGTAGCAAGGAGAAGG-3’) for improved results.

2.4 | Data management

The azole resistance prevalence was determined at patient level and compared to the Dutch national surveillance, 2013–2018.22 Azole resistance was divided into environmentally driven resistance (presence of Trs46/L98H or Trs46/Y121F/T289A), other cyp51A mutations and non-cyp51A-mediated resistance (when resistant, but no cyp51A mutations were identified).


The surveillance was requested by the Danish Ministry of Health and the scientific study approved by the QA & Compliance at Statens Serum Institut (journal number 21/00765).

Preliminary results have previously been presented in part at the Trends in Medical Mycology 2019 and at European Congress on Clinical Microbiology and Infectious Disease (ECCMID) in 2020 and 2021 conferences and briefly summarised as part of the national Danish Integrated Antimicrobial Resistance Monitoring and Research Programme (DANMAP) established by the Danish Ministry of Food, Agriculture and Fisheries and the Danish Ministry of Health in annual reports 2018, 2019 and 2020 reports.23,24 The results presented here are updated since then.

3 | RESULTS

A total of 1820 susceptibility-tested A. fumigatus isolates from 1083 patients were included in the analysis. The vast majority originated from airways including nose/sinus (1609/1820) and ear samples (182/1820) (Table 1).

Itraconazole resistance was found in 5.9% (108/1820) of isolates and voriconazole resistance in 5.6% (102/1820) (Figure 1). Posaconazole resistance was detected in 103 isolates due to MICs of ≥0.5 mg/L, and 85 had MIC 0.25 mg/L (defined as area of technical uncertainty [ATU]) of which 6 were classified as resistant due to an itraconazole MIC >1 mg/L. Isavuconazole resistance with MICs of ≥4 mg/L was detected in 90 isolates, and 235 isolates had MIC 2 mg/L (ATU) of which 13 were classified as resistant due to a voriconazole MIC >1 mg/L. Overall, susceptibility testing identified 119 isolates...
resistant to at least one azole from 66 patients leading to a resistance prevalence among patients of 6.1% (66/1083, 95% CI 4.8%–7.7%). The proportion of isolates that were azole resistant was 6.5% (119/1820). From lower airways, the proportion of resistant isolates were 4.3% (12/278) compared to 3.6% (6/166) of isolates from tracheal aspirates and 8.5% (99/1165) of isolates in the upper airways (Table 1).

**TABLE 1** Number of patients and sample types with *Aspergillus fumigatus* isolates

<table>
<thead>
<tr>
<th>Sample type (n)</th>
<th>Clinical samples</th>
<th>Background samples</th>
<th>Total</th>
<th>Proportion of resistant isolates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male/Female (n)</td>
<td>527/462</td>
<td>48/46</td>
<td>575/508</td>
<td>—</td>
</tr>
<tr>
<td>Isolates (n)</td>
<td>1721</td>
<td>99</td>
<td>1820</td>
<td>—</td>
</tr>
<tr>
<td>Sputum samples/sinus/nose&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1113</td>
<td>52</td>
<td>1165</td>
<td>8.5% (99/1165)</td>
</tr>
<tr>
<td>Tracheal aspirate</td>
<td>155</td>
<td>21</td>
<td>166</td>
<td>3.6% (6/166)</td>
</tr>
<tr>
<td>BAL/Pleura fluid/Lung/Lung biopsy&lt;sup&gt;b&lt;/sup&gt;</td>
<td>167</td>
<td>11</td>
<td>278</td>
<td>4.3% (12/278)</td>
</tr>
<tr>
<td>Other deep samples&lt;sup&gt;c&lt;/sup&gt;</td>
<td>12</td>
<td>12</td>
<td>12</td>
<td>8.3% (1/12)</td>
</tr>
<tr>
<td>Ear</td>
<td>168</td>
<td>14</td>
<td>182</td>
<td>0.5% (1/182)</td>
</tr>
<tr>
<td>Cornea/Eye swab</td>
<td>6</td>
<td>6</td>
<td>Not detected</td>
<td></td>
</tr>
<tr>
<td>Tissues not specified/scar/puncture site</td>
<td>10</td>
<td>1</td>
<td>11</td>
<td>Not detected</td>
</tr>
</tbody>
</table>

<sup>a</sup> Includes samples marked as sputum/laryngeal aspirate, sinus, nose/nose vestibule biopsy/nasal aspirate/nose-throat.

<sup>b</sup> Includes samples marked as BAL/bronchial aspirate/pleura fluid and lung biopsy/lung/pleura.

<sup>c</sup> Cerebrospinal fluid, abscess/drain fluid/drain/abdominal swab, biopsy abdominal/biopsy organ not specified/pericardium/pericardial fluid and bone.

**FIGURE 1** MIC values for the included *Aspergillus fumigatus* isolates. Susceptible isolates (S) are shown green when susceptible at azole resistance screening. Susceptible isolates with an MIC are shown in blue, resistant isolates in red and isolates in the ATU for which the classification depends on the susceptibility of either itraconazole or voriconazole, respectively, are indicated in black. MIC values above 4 mg/L are shown as >4 mg/L. Isolates with no MICs for posaconazole (n = 1) and isavuconazole (n = 365) are not included in the diagrams.
The proportional distribution of resistance mechanisms at patient level are shown in Figure 2. cyp51A sequencing of azole-resistant A. fumigatus demonstrated environmental resistance (TR$_{34}$/L98H, TR$_{34}$/L98H, or TR$_{34}$/L98H/2977F/F495I) in isolates from 39 patients. This corresponds to an environmental resistance prevalence among the patients of 3.6% (39/1083 patients; 95% CI: 2.6%–4.9%) and accounted for 59.1% (39/66 patients; 95% CI: 47.0%–70.1%) of patients with resistant A. fumigatus isolates (Figure 2). Resistance with a tandem repeat was detected in samples from airways (Sputum [n = 44], BAL [n = 6] and tracheal aspirate [n = 4]) and one ear sample.

Resistance involving other cyp51A mutations accounted for 21.2% (14/66; 95% CI: 13.1%–32.5%). The corresponding alterations were G54R (n = 5), P216S (n = 2), F219L (n = 1), G54W (n = 1), M220I (n = 1), M220K (n = 1), M220R (n = 1), G432S (n = 1), G448S (n = 1) and Y121F (n = 1). One patient had sequential isolates with either M220R or G54R.

Non-cyp51A-mediated resistance (wild-type cyp51A) accounted for 19.7% (13/66; 95% CI: 11.9%–30.8%). Isolates from 12 of these patients were voriconazole resistant with MICs ≥4 mg/L or with MIC 2 mg/L and cross-resistance to the other azoles. One patient had an isolate that was classified as resistant solely due to a voriconazole MIC of 2 mg/L.

Among patients with a resistant isolate, both susceptible and resistant isolates were cultured intermittently during the surveillance from 38/66 (58%). Twenty-five patients had only one resistant isolate, and three patients had several consecutive resistant isolates.

Four unique resistant isolates did not undergo cyp51A-sequencing. Three isolates from a patient who had several resistant isolates with M220K, and another isolate from a patient who had isolates with P216S.

Isolates marked as 'Background samples' included 99 isolates from 94 patients (Table 1). A. fumigatus from three patients (3.2%; 95% CI: 0.9%–9.0%) were azole resistant and all harboured the TR$_{34}$/L98H resistance mechanism. One patient had consecutive isolates with TR$_{34}$/L98H-marked background and not marked as background.

Azole resistance was detected in samples from all five Danish regions (Figure 3). TR$_{34}$/L98H isolates were detected in four out of five regions and in samples from both the hospital and the primary health care sector, whereas isolates with single point mutations in cyp51A were found in three of five regions.

Comparing surveillances at national level, azole resistance prevalence was lower in Denmark than in the Netherlands (66/1083 [6.1%] vs. 508/4496 [11.3%]) (p < .0001).

### 4 | DISCUSSION

An azole resistance prevalence of 6.1% including a TR$_{34}$/L98H-related environmental resistance of 3.6% was documented at patient level during the first 2 years of the Danish nationwide surveillance programme. Whereas the first figure represents the current burden of azole resistance, the second provides information on what the chances are for facing azole resistance among azole-naïve patients. The resistance prevalence was higher in samples from the upper airways than in tracheal aspirates and lower airway specimens. We speculate, that this may reflect that out-patients with chronic lung disease including CF and aspergillosis are often provided with sputum containers for regular submission of sputum by mail and thus that the resistance frequencies across the different sample types are not directly comparable. Overall the resistance prevalence remains well below 10%, and azole therapy therefore remains the first choice for the initial treatment for aspergillosis in our country.

Several observations suggest that azole resistance in A. fumigatus and resistance due to TR$_{34}$/L98H specifically is increasing in Denmark. In 2007, 1.9% A. fumigatus isolates were azole resistant and none due to TR$_{34}$/L98H mechanisms in a 3 month multicentre survey. From 2010 to 2014, the azole resistance prevalence at patient level was 2.1% among referred (and thus selected) isolates with approximately half involving TR$_{34}$/L98H mechanisms. Moreover, in the Danish CF population, specifically, an azole resistance prevalence of 4.5% including 1.5% due to TR$_{34}$/L98H was observed in 2007 and 2009 compared to 7.3% including 3.7% TR$_{34}$/L98H in 2018. Although the studies are not directly comparable, we speculate that azole resistance is rising both overall and among CF patients and is driven by both medical and environmental azole use.
### TABLE 2  MIC values for *Aspergillus fumigatus* isolates resistant to at least one azole and which underwent cyp51A sequencing

<table>
<thead>
<tr>
<th>Resistance mechanism</th>
<th>Isolates (n)</th>
<th>MIC medians and ranges (mg/L)</th>
<th>ITR</th>
<th>POS</th>
<th>VOR</th>
<th>ISA</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Environmental</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TR$_{34}$/L98H</td>
<td>50</td>
<td>&gt;16 (2–&gt;16)</td>
<td>0.5/1 (0.5–4)</td>
<td>4 (2–16)</td>
<td>8 (4–16)</td>
<td></td>
</tr>
<tr>
<td>TR$_{34}^3$/L98H</td>
<td>4</td>
<td>&gt;16</td>
<td>1 (1–2)</td>
<td>8 (4–8)</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>TR$_{34}$/L98H/S297T/F495I</td>
<td>1</td>
<td>&gt;16</td>
<td>2</td>
<td>4</td>
<td>&gt;16</td>
<td></td>
</tr>
<tr>
<td><strong>Single point mutations</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y121F</td>
<td>2</td>
<td>&gt;4–16</td>
<td>1–4</td>
<td>&gt;4–16</td>
<td>&gt;8</td>
<td></td>
</tr>
<tr>
<td>G448S</td>
<td>2</td>
<td>&gt;4</td>
<td>(0.5–1)</td>
<td>&gt;4</td>
<td>&gt;8</td>
<td></td>
</tr>
<tr>
<td>G432S</td>
<td>3</td>
<td>&gt;4 (&gt;4–&gt;16)</td>
<td>2 (0.5–4)</td>
<td>4 (2–4)</td>
<td>8 (4–16)</td>
<td></td>
</tr>
<tr>
<td>M220R</td>
<td>4</td>
<td>&gt;4/16 (&gt;4–&gt;16)</td>
<td>1/4 (0.5–4)</td>
<td>2/4 (1–4)</td>
<td>4/8 (4–8)</td>
<td></td>
</tr>
<tr>
<td>M220K</td>
<td>9</td>
<td>&gt;16 (4–&gt;16)</td>
<td>2 (1–4)</td>
<td>2 (1–2)</td>
<td>2 (1–2)</td>
<td></td>
</tr>
<tr>
<td>M220I</td>
<td>1</td>
<td>&gt;16</td>
<td>0.5</td>
<td>0.5</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>G54R</td>
<td>6</td>
<td>&gt;16 (&gt;4–&gt;16)</td>
<td>4 (2–4)</td>
<td>1 (0.25–4)</td>
<td>2 (0.5–4)</td>
<td></td>
</tr>
<tr>
<td>G54W</td>
<td>1</td>
<td>&gt;16</td>
<td>&gt;4</td>
<td>0.5</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>F219L</td>
<td>1</td>
<td>&gt;4</td>
<td>0.5</td>
<td>1</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>P216S</td>
<td>3</td>
<td>2–16</td>
<td>0.25–0.5</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td><strong>Non-cyp51A mediated</strong></td>
<td>27</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Note:** Resistance mechanisms are shown according to environmental, single point mutations and non-cyp51A-mediated. Single point mutations are shown according to decreasing resistance.

**Abbreviations:** ISA, Isavuconazole; ITR, Itraconazole; POS, Posaconazole; VOR, Voriconazole.

One resistant isolate is not shown in this table since it was found with a F46Y/M172V/E427K, which is not associated with azole resistance, and the same patient had other resistant isolates with TR$_{34}$/L98H. Four isolates did not undergo cyp51A-sequencing and are not shown in the table.

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**FIGURE 3** Proportion of resistant *Aspergillus fumigatus* isolates and associated underlying resistance mechanism across the five Danish Regions. Each Region represented is the Region of the health care facility from which the isolate was referred. As some health care services are centralised this will not in all cases represent the patients’ place of residence or the place where the resistant fungus was acquired. Total numbers of isolates were for Capital $n = 910$, Zealand $n = 91$, Southern Denmark $n = 326$, Central Jutland $n = 419$ and Northern Jutland $n = 74$. The resistance mechanism remained uncharacterised in five isolates from five patients whom were known to harbour other cyp51A mutant isolates (blue bar). These included four resistant isolates that did not undergo cyp51A-sequencing of which three isolates derived from a patient who had other isolates with M220K, and one isolate from a patient who had other isolates with P216S, and one isolate with F46Y/M172/E427K from a patient who also had isolates with TR$_{34}$/L98H.
Of note, we did not observe any isolates harbouring TR\textsubscript{34}/L98H/T289A during the 2-year surveillance although this resistance genotype has been found once in DK in 2014.\textsuperscript{4}

Three single point amino acid alterations (G54A, G54R and G432S) have been associated with azole resistance in both azole-treated patients and the environment.\textsuperscript{27-30} Five patients in this surveillance programme harboured isolates with a G54R and one patient an isolate with a G432S alteration. Unfortunately, we did not have access to clinical information or prior medication data to enable a discussion of the origin of these resistance mechanisms.

The number of patients with azole-resistant \textit{A. fumigatus} was unevenly distributed across the five regions in Denmark. The reason for a higher occurrence in the capital region is likely that this is the largest region based on population size, and that one of the two CF centres is based in the capital region. TR\textsubscript{34}/L98H was not detected in northern Jutland during the study period, but was detected in a clinical isolate shortly before the surveillance programme was initiated.\textsuperscript{24} We therefore argue that TR\textsubscript{34}/L98H is found all over the country and pose a risk for any patient in Denmark susceptible to \textit{Aspergillus} infections.

In comparison to other surveillance studies, our azole resistance prevalence was higher than the 3.2% found in 2009 to 2011 in a multicentre study with 19 countries,\textsuperscript{31} but lower than the 11% in the more recent Dutch nationwide surveillance in 2017 and 2018.\textsuperscript{22} Larger studies and surveillances on azole resistance in \textit{A. fumigatus} are summarised in Table 3.\textsuperscript{22,25,32-39} These studies show that the

### Table 3: Studies and surveillances with azole resistance in \textit{Aspergillus fumigatus}

<table>
<thead>
<tr>
<th>Country and study period</th>
<th>Study type/Setting</th>
<th>Azole resistance prevalence</th>
<th>TR\textsubscript{34}/L98H and/or TR\textsubscript{46}/Y121F proportion of resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Europe</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Denmark</td>
<td>Nationwide surveillance 3-month</td>
<td>1.9% isolate level (2/107)</td>
<td>Not detected</td>
</tr>
<tr>
<td>(Mortensen et al.)\textsuperscript{25}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>The Netherlands</td>
<td>Nationwide surveillance</td>
<td>11.3% patient level (508/4496)</td>
<td>556/640 of resistant isolates\textsuperscript{a}</td>
</tr>
<tr>
<td>(Lestrado et al.)\textsuperscript{22}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>The Netherlands</td>
<td>Multicentre study</td>
<td>5.3% patient level (63/1192)</td>
<td>4.1% isolate level (74/1792)</td>
</tr>
<tr>
<td>(Van der Linden et al.)\textsuperscript{32}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Belgium</td>
<td>Multicentre study</td>
<td>5.5% patient level (9/164)</td>
<td>4.3% patient level (7/164)</td>
</tr>
<tr>
<td>(Vermeulen et al.)\textsuperscript{33}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spain</td>
<td>Multicentre study</td>
<td>4.7% patient level (34/715)</td>
<td>2.8% patient level (20/715)</td>
</tr>
<tr>
<td>(Escribano et al.)\textsuperscript{34}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Italy</td>
<td>Multicentre study</td>
<td>6.6% isolate level (19/286)</td>
<td>4.2% isolate level (12/286)</td>
</tr>
<tr>
<td>(Prigitano et al.)\textsuperscript{35}</td>
<td></td>
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<tr>
<td><strong>North America</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USA</td>
<td>Multicentre study/surveillance</td>
<td>1.5% isolate level (20/1356)</td>
<td>0.4% isolate level (5/1356)</td>
</tr>
<tr>
<td>(Berkow et al.)\textsuperscript{36}</td>
<td></td>
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<tr>
<td><strong>Asia</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Japan</td>
<td>Multicentre surveillance</td>
<td>12.7% isolate level (7/55)</td>
<td>3.6% isolate level (2/55)</td>
</tr>
<tr>
<td>(Tsuchido et al.)\textsuperscript{39}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Taiwan</td>
<td>Multicentre study</td>
<td>4% patient level (12/297)</td>
<td>3.4% patient level (10/297)</td>
</tr>
<tr>
<td>(Wu et al.)\textsuperscript{37}</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>China</td>
<td>Multicentre surveillance</td>
<td>2.5% isolate level (8/317)</td>
<td>2.5% isolate level (8/317)</td>
</tr>
<tr>
<td>(Chen et al.)\textsuperscript{38}</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

Note: Studies shown involve those who are either nationwide surveillances or multicentre studies in one country and not limited to a certain patient group or a referral hospital. Studies included are those that report azole resistance in \textit{A. fumigatus} specifically. Azole resistance in \textit{A. fumigatus} prevalence is shown in numbers of patients unless other specified.

\textsuperscript{a} Numbers of patients with either TR\textsubscript{34}/L98H or TR\textsubscript{46}/Y121F were not specified, and total number of isolates was not specified.
azole resistance prevalence in the present surveillance is in line with other European studies from 2011 to 2018 and the Netherlands from 2007 to 2009.

This study is associated with both strengths and limitations. The primary strength is that it is nationwide and thus population based. Results in studies limited to specific disease or centre will depend strongly on the case mix and use of azole therapy, which would favour selection of azole-resistant A. fumigatus. Limitations include a risk of ascertainment bias. We cannot exclude that centres managing many Aspergillus patients are more prone to prioritise referral of isolates – a so-called cluster sampling. Furthermore, the COVID-19 pandemic emerged during the surveillance, and we cannot be certain that routine sampling was performed as under regular circumstances. Indeed fewer BALs were performed, and out-patients with lung diseases were more often encouraged to send sputum samples by mail, than to visit the clinic in person.

Our classification of isolates as background samples is also associated with limitations. Not all laboratories adhered strictly to the inclusion criteria and not all laboratories referred background samples. Clinical data and information on prior antifungal treatment were not collected and therefore we cannot verify that the samples marked ‘Background’ actually represented a clinically insignificant background samples or that all such samples were indeed marked as Background samples. However, the fact that no isolates with medically driven point mutations were found among background samples suggests that Background samples at least are dominated by isolates from patients without prior azole therapy for clinically documented infection and thus representative for the background level of environmental resistance.

In conclusion, azole resistance is a significant problem for patients with clinical disease and in need of azole treatment. Few or no oral alternative drug options combined with long duration of treatment is a clinical challenge and results in a worsened prognosis. Initial treatment of invasive aspergillosis can remain unchanged in Denmark – but optimal treatment strategies do depend on the likelihood of azole resistance – highlighting the importance of continued surveillance, rapid susceptibility testing and a one-health approach to azole use.

ACKNOWLEDGEMENTS
The authors acknowledge the laboratory staff at the mycology unit at Statens Serum Institut.

CONFLICT OF INTEREST
MR: Has received research-and travel grants from Gilead. RKH: Has over the past 5 years received travel grants and speaker honoraria from Gilead. JBG: Has over the past 5 years received travel grants and speaker honoraria from Gilead. LC: No conflicts of interest. FSR: No conflicts of interest. SS: No conflicts of interest. NA: No conflicts of interest. JB: No conflicts of interest. BLR: No conflicts of interest. EM: No conflicts of interest. KA: Has received travel grant and speaker honoraria from Gilead. MP: No conflicts of interest. ED: No conflicts of interest. SLA: No conflicts of interest. MCA: has outside the current work, over the past 5 years, received research grants/contract work (paid to the SSI) from Amplyx, Basilea, Cidara, F2G, Gilead, Novabiotics and Scynexis, and speaker honoraria (personal fee) from Astellas, Chiesi, Gilead, MSD, and SEGES. She is the current chairman of the EUCAST-AFST.

AUTHOR CONTRIBUTIONS
Malene Risum: Data curation (lead); Formal analysis (equal); Investigation (equal); Project administration (equal); Writing – original draft (lead). Rasmus Krøger Hare: Data curation (equal); Investigation (equal); Methodology (lead); Writing – review & editing (equal). Jan Berg Gertsen: Data curation (equal); Investigation (equal); Methodology (equal); Project administration (equal); Writing – review & editing (supporting). Lise Kristensen: Investigation (equal); Methodology (equal); Writing – review & editing (supporting). Flemming Schanning Rosenvinge: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Sofia Sulim: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Karen Astavad: Investigation (equal); Methodology (supporting); Writing – review & editing (supporting). Nissrine Abou-Chakra: Investigation (supporting); Methodology (equal); Writing – review & editing (supporting). Jette Bangsbjerg: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Bent Røder: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Esad Dzajic: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Steen Lomborg Andersen: Investigation (supporting); Methodology (supporting); Writing – review & editing (supporting). Maiken Cavling Arendrup: Conceptualization (lead); Data curation (equal); Formal analysis (equal); Investigation (equal); Methodology (lead); Project administration (lead); Writing – original draft (lead).

DATA AVAILABILITY STATEMENT
Data are only available for research upon reasonable request to Statens Serum Institut and within the framework of the Danish data protection legislation.

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REFERENCES


