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Time trends in diagnostic testing for primary ciliary dyskinesia in Europe

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Time trends in diagnostic testing for PCD in

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- 3 To the Editor: Despite recent advances in diagnostic methods, diagnosis of primary ciliary dyskinesia (PCD) remains complex. We need a combination of different diagnostic tests, and all have their limitations [1]. In 2009, the first European Respiratory Society Task Force (ERS TF) on PCD in children published recommendations [2], suggesting that: 1) Nasal nitric oxide (nNO) should be measured to screen for PCD in patients aged \geq 5 years [3]; and 2) video microscopy (VM) analysis of ciliary beat pattern and frequency [4] plus electron microscopy (EM) [5] should be the key confirmatory diagnostic tests. Genetic testing was not recommended as part of the initial diagnostic testing, but as additional test for inconclusive cases. The recommended test combination was nNO, VM and EM for patients aged \geq 5 years and VM plus EM for younger patients. <u>18</u> In 2017, a second ERS TF on PCD diagnosis revised the accumulated literature and published 20 57 evidence-based guidelines [6]. Although evidence-based guidelines have become the norm in 23 24 research, their practical implementation can be challenging [7]. We wanted to assess whether the 2009 diagnostic recommendations had been implemented and how diagnosis of PCD changed in Europe over time. This knowledge will help to improve implementation of the new guidelines. 28 We analysed data from the international PCD cohort (iPCD) (details are published elsewhere [8]). 31 By May 2018, iPCD included data on 3733 patients from 26 centres in 21 countries. For this study, we included all datasets from European centres that tested patients with PCD, both before and after 2009, and had complete information on nNO, EM and VM testing. We excluded patients in 36 whom diagnosis was based only on clinical presentation, patients with unknown dates of testing. 38 We included 2108 patients from 16 centres (11 European countries) (Belgium, Cyprus, Czech Republic, France, Germany, Italy, Norway, Poland, Switzerland, Turkey and United Kingdom); 51% were male, 818 patients (39%) had been diagnosed before and 1290 after 2009. All three 42 70 recommended tests were available in all countries, with the exception of Norway where VM testing was not available neither before nor after 2009. 45 Based on the 2009 recommendations, we only considered nNO measurements in patients aged 5 years or older [2]. We considered the nNO test as positive when nNO was below 77 nL·min⁻¹ [9,10]. VM had been performed with different techniques over time, with high speed video analysis 49 74 50 75 being the most commonly used technique in recent years. We classified VM and EM results as 52 76 pathological based on information provided by the centres on the beat frequency, beat pattern and cilia ultrastructure. For each patient, we defined the calendar year of diagnosis based on the date of the earliest positive test result. We then assessed whether there was a change over time in the 5 5

- proportion of diagnosed patients who had received a) the recommended test combination; b) any 57 79
- 58 80 59 60 single test. We compared the proportion of patients with the recommended test combination (VM

2 3 81 and EM for patients aged <5 years and nNO, VM and EM for older patients) for the two time 4 5 6 82 periods, before and after 2009. We used R version 3.1.2 for all analyses. 7 83 Recommended test combination: Overall, we found no significant trend over time in the use of 8 the test combination. The three tests had been used in 54% of patients diagnosed before 2009 and 9 84 10 in 57% after 2009 (p=0.15) (Figure 1). In preschool children the proportion diagnosed with the 85 11 12 86 recommended combination was 72% before and 75% (p=0.47) after 2009; in older patients it increased from 46% to 52% (p=0.03). Results differed between countries. Few countries (e.g. 13 87 14 15 88 Belgium, Cyprus) combined all 3 tests already before 2009 for most patients and continued to do 89 so after 2009. In Germany, the UK and the Czech Republic, the combined use of all 3 tests was 16 17 18 90 common already before 2009 but increased even more after 2009, with almost ³/₄ of the patients 19 tested according to recommendations. The remaining countries (Turkey, Switzerland, Italy, France 20 91 21 22 and Poland) showed little or no change over time. In these countries, less than half of the patients, 92 were tested with all 3 approaches even in the later period. 23 93 24 25 Nasal NO testing increased overall from 63% before 2009 to 84% afterwards (p<0.001). This 94 26 increase was seen in most countries (Figure 1). After 2009, nNO was measured in over 34 of 27 95 28 29 patients in all countries, except in Czech Republic (65%), Italy (70%) and UK (77%). 96 Electron microscopy was frequently performed before 2009 (97%) but decreased to 80% 30 97 (p<0.001) in the later period. Its use became less common in Poland (79% to 69%), Switzerland 31 98 32 33 99 (88% to 62%) and Turkey (100% to 18%), in all other countries it remained stable or increased 34 100 after 2009. Video microscopy analysis increased overall from 76% to 87% (p<0.001). This was 36 101 mainly because the use of VM for PCD diagnosis increased considerably in Italy (36% to 69%) and 37 Turkey (25% to 88%). In most countries, its use remained stable, while in Switzerland (50% to 38 102 39 103 21%) it decreased substantially. 40 41 104 This is the first multi-national study that compared diagnostic testing in PCD patients between 4 43 ₁₀₅ countries and over time. Although a large number of countries contribute to iPCD, some had to be 44 45 106 excluded for this analysis as they only contributed patients diagnosed after 2009 to the iPCD 46 107 cohort. Thus, our study describes how the consensus recommendations were implemented in 11 47 countries. They are not representative for all European countries, but only for those with 48 108 49 109 established PCD diagnostic protocols. In this analysis we included both children and adults. 50 51 110 However when we limited the analysis to children only, for whom the 2009 recommendations were 52 53 ¹¹¹ intended, results remained similar. 55 112 Our results suggest that the implementation of the recommended diagnostic combination of nNO, 56 113 EM and VM testing after the 2009 consensus statement remained low. This reflects the complex 57 58 114 nature of PCD diagnostics and the regional resources. Many countries continued to perform only 59 one or two of the recommended tests. There are several explanations for this observation. First, 60 115

1 2 3 116 the availability of local resources could have led to the development of alternative diagnostic 4 5 pathways, which may have been most appropriate for the local situation at that time. All PCD 117 6 118 diagnostic tests need specialised expensive equipment and personnel experienced in analysis of 7 VM and EM results, which are not available in all settings. Limited resources or decentralised 119 8 healthcare might not have allowed to set up diagnostic centres with scientists experienced in all 9 120 11 121 methods. For countries with limited resources cost-effective alternatives for diagnostic testing have 12 been suggested, which might provide an acceptable diagnostic accuracy [11]. Second, since 2009 13 122 14 123 the use of other methods including genetic testing [12,13] and immunofluorescence microscopy 15 16 124 [14,15] became more widespread. These newer methods might have been used instead of the 17 125 recommended tests in some centres. Lastly, the lack of sufficient evidence supporting the use of 18 some diagnostic tests in 2009 might have prevented some countries to implement the full set of 19 126 20 21 127 recommended tests but let them to develop their own diagnostic algorithms. We found 22 128 23 considerable heterogeneity between countries in the use of the three tests. Overall, countries with low prior use of nNO showed improvement and nNO is now used in most patients aged \geq 5 years 24 129 suspected for PCD. For the proportion of patients who were still not tested after 2009, we 25 130 26 27 131 speculate that nNO was not performed as a screening test, and the primary investigators chose to do directly one or both of the other tests. In this case, if the diagnosis was already established 28 132 29 30 133 based on the results of the other tests, the patients might not have been invited posthoc to perform 31 also nNO measurement. This would be in line with the recommendations. We found that use of EM 32 134 33 135 analysis decreased, and VM increased, suggesting that there might be a shift from EM to VM 34 35 136 overall. Possible reasons are the realization that a significant proportion of patients have normal EM findings [16] and the high costs of EM analysis combined with an increased availability of VM, 36 137 37 38 138 39 so that only patients with inconclusive VM results were referred for EM testing. The overall changes in use of VM and EM analyses were strongly affected by the marked increase in VM and 40 139 decrease in EM analysis in Turkish patients. This shift is explained by the development of a new 41 140 42 PCD centre, which uses VM more and EM less frequently. 43 141 44 45 142 The 2009 PCD diagnostic consensus is a typical example of how difficult it is to implement 46 47 143 guidelines in clinical practice. Even though the recommendations were widely presented in 48 144 scientific conferences and meetings, improving knowledge is not sufficient to change daily 49 50 145 practices. A synthesis of systematic reviews on clinical guideline implementation strategies showed that passive dissemination was an ineffective measure and that implementation strategies should 51 146 52 be multifaceted, and actively engage clinicians throughout the process [7]. In the case of PCD 53 147 diagnosis, implementation is further hindered by fragmentation of national diagnostic services in 54 148 55 many centres and the cost of diagnostic equipment. In our study, countries with limited resources 56 149 57 (e.g. Poland, Turkey) or decentralised diagnosis (e.g. France, Italy, Switzerland) performed the 58 150 59 151 recommended test combination less frequently, than countries with more resources (e.g. Germany, 60 152 Belgium, UK) or established centralised PCD diagnosis (e.g. Cyprus, UK). National and multi-

1 2 3	153	national collaborations, such as the European Reference Network for respiratory diseases (ERN-
4 5 6	154 155	Lung; https://ern-lung.eu/) might play an important role, in the future to facilitate centralised diagnosis and standardised patient care. With the further development and improvement of
7 8	156	diagnostic tests for PCD and with new centres emerging, that might lack the necessary expertise,
9 1 11	157	there is an increased need for national and international collaboration in PCD diagnostic testing.
12	158	Overall, we found a low adherence to the 2009 consensus recommendations mainly due to the
14	159	test combination. To further improve DCD diagnosis, we must be more diligent and engaging in
15 16 17	160 161	implementing the new evidence-based guidelines published in 2017, putting more emphasis on
18 19	162	establishing specialised diagnostic centres and close international collaboration.
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Figure 1: Proportion of performed diagnostic tests in European countries before and after the 2009 consensus statement on PCD diagnostics.