The top 10 research priorities in cystic fibrosis developed by a partnership between people with CF and healthcare providers

ABSTRACT
There remain many treatment uncertainties in cystic fibrosis (CF). With limited resources, research should focus on questions which are most important to the CF community. We conducted a James Lind Alliance Priority Setting Partnership in CF. Research questions were elicited and then prioritised in successive surveys. A workshop agreed the final top 10. Online methods avoided cross infection and widened participation. The elicitation survey had 482 respondents (1080 questions) and prioritisation survey 677 respondents. Participants were drawn equally from the patient and clinical communities globally. We have achieved a consensus on 10 research priorities which will be attractive to funders.

INTRODUCTION
Therapies targeting the basic defect of cystic fibrosis (CF) are now available in the clinic, with more in development.1 However, conventional therapies for persistent airway infection and pancreatic malabsorption are still required and there remain many treatment uncertainties, which have yet to be clarified through well designed and adequately powered clinical trials.2

Both the Patient-Centered Outcomes Research Institute, in the USA, and the National Institute for Health Research (NIHR), in the UK, support patient involvement and ‘co-production’ of research.3 However, there has been no systematic and inclusive dialogue between clinicians, patients and parents about priorities for clinical research in CF. Face-to-face contact between patients is precluded due to risk of transmission of organisms such as Pseudomonas aeruginosa, providing a challenge to consultative exercises involving patients with CF.4

The NIHR James Lind Alliance (JLA) supports Priority Setting Partnerships which bring patients, carers and clinicians together to prioritise research questions. Outcomes from JLA Priority Setting Partnerships have a track record of attracting significant research funding.5 We conducted a JLA Priority Setting Partnership in CF to identify the key treatment uncertainties for the CF community and produce the top 10 research questions for treatment and management of CF.

METHODS
Our Priority Setting Partnership took place in March 2016–January 2017. We published our protocol before commencing (http://eprints.nottingham.ac.uk/35223/). We followed JLA methodology,6 adapting the process using online methods to avoid risk of cross infection. Our steering group was selected to include lay members (two people with CF and two parents) and professionals from each discipline: two respiratory paediatricians, respiratory physician, physiotherapist, dietitian, nurse, pharmacist, clinical psychologist and social worker. We also included a specialist commissioner, UK Cystic Fibrosis Trust representative and four academics (systematic reviewer, qualitative researcher, project manager and academic clinical fellow).

From March to June 2016, we conducted the first of two online surveys (Survey Monkey). The elicitation survey requested respondent demographics and asked a single question ‘What question on Cystic Fibrosis would you like to see answered by research?’ Respondents could submit up to five questions. The survey was advertised through professional groups, clinic posters and our bespoke Twitter account (@questionCF).

Two researchers (NR and SS) independently reviewed all questions submitted with adjudication by the steering group. Non-questions and questions unrelated to treatment were removed. We conducted a systematic review of evidence gaps in CF7 and excluded any questions already answered. Where several questions addressed the same issue, they were consolidated into a single ‘standardised question’.

The steering group undertook a Delphi8 assessment of the standardised questions to produce a shortlist for the prioritisation

Table 1  Demographic characteristics of respondents to elicitation survey and prioritisation survey. Of the 513 responses to elicitation survey, 31 were from respondents who completed the survey more than once (482 individual respondents). Data are presented as number (%) or median (range)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Elicitation survey (n=482)</th>
<th>Prioritisation survey (n=677)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>111 (23.0)</td>
<td>153 (22.6)</td>
</tr>
<tr>
<td>Female</td>
<td>315 (63.4)</td>
<td>454 (67.1)</td>
</tr>
<tr>
<td>Did not answer</td>
<td>56 (11.6)</td>
<td>70 (10.3)</td>
</tr>
<tr>
<td>Median age (range)</td>
<td>42 (6–82)</td>
<td>41 (6–82)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Category of respondent</th>
<th>Elicitation survey (n=482)</th>
<th>Prioritisation survey (n=677)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lay persons</td>
<td>224 (46.5)</td>
<td>314 (46.4)</td>
</tr>
<tr>
<td>Person with cystic fibrosis</td>
<td>95 (19.7)</td>
<td>121 (17.9)</td>
</tr>
<tr>
<td>Parent of person with cystic fibrosis</td>
<td>105 (21.8)</td>
<td>160 (23.6)</td>
</tr>
<tr>
<td>Family and friends</td>
<td>24 (5.0)</td>
<td>33 (4.9)</td>
</tr>
<tr>
<td>Professionals</td>
<td>211 (43.8)</td>
<td>303 (44.8)</td>
</tr>
<tr>
<td>Medical doctor</td>
<td>61 (12.7)</td>
<td>85 (12.6)</td>
</tr>
<tr>
<td>Nurse</td>
<td>22 (4.6)</td>
<td>32 (4.7)</td>
</tr>
<tr>
<td>Physiotherapist</td>
<td>49 (10.2)</td>
<td>65 (9.6)</td>
</tr>
<tr>
<td>Dietitian</td>
<td>31 (6.4)</td>
<td>41 (6.1)</td>
</tr>
<tr>
<td>Pharmacist</td>
<td>6 (1.2)</td>
<td>18 (2.7)</td>
</tr>
<tr>
<td>Social worker</td>
<td>7 (1.5)</td>
<td>16 (2.4)</td>
</tr>
<tr>
<td>Non-clinical researcher</td>
<td>10 (2.1)</td>
<td>7 (1.0)</td>
</tr>
<tr>
<td>Psychologist</td>
<td>21 (4.4)</td>
<td>38 (5.6)</td>
</tr>
<tr>
<td>Other</td>
<td>4 (0.8)</td>
<td>1 (0.1)</td>
</tr>
<tr>
<td>Did not answer</td>
<td>47 (9.8)</td>
<td>60 (8.9)</td>
</tr>
<tr>
<td>Country</td>
<td></td>
<td></td>
</tr>
<tr>
<td>UK</td>
<td>249 (51.7)</td>
<td>474 (70.0)</td>
</tr>
<tr>
<td>USA and Canada</td>
<td>91 (18.9)</td>
<td>70 (10.3)</td>
</tr>
<tr>
<td>Rest of Europe</td>
<td>45 (9.3)</td>
<td>42 (6.2)</td>
</tr>
<tr>
<td>Australia and New Zealand</td>
<td>35 (7.3)</td>
<td>22 (3.2)</td>
</tr>
<tr>
<td>Other</td>
<td>5 (1.0)</td>
<td>5 (0.7)</td>
</tr>
<tr>
<td>Did not answer</td>
<td>57 (11.8)</td>
<td>64 (9.5)</td>
</tr>
</tbody>
</table>
survey. This allowed participants to rank their top 10 questions (September–November 2016). The prioritisation survey was advertised through Twitter and by emailing participants who completed the elicitation survey.

The top 20 questions chosen through the prioritisation survey were then discussed at a workshop in January 2017. Lay and professional participants (recruited through the prioritisation survey), all with equal voice, led by three independent JLA facilitators, selected the final top 10 research priorities, through a process of small group discussion and plenary voting. Patient representatives joined remotely from home and hospital using video conferencing.

RESULTS
We had 513 replies to the elicitation survey, of which 31 completed the survey more than once, leaving 482 respondents who submitted 1080 questions. Table 1 shows the demographics of elicitation survey respondents. There were 224 (46.5%) lay respondents, 211 (43.8%) professionals and 47 (9.8%) were unknown. Just over half were from the UK, with submissions from 23 countries in total.

Figure 1A describes how the questions were sorted and refined. There were 704 treatment and management questions which were combined into 127 standardised questions. Following the Delphi process, 71 questions were taken forward for prioritisation. The prioritisation survey was completed by 677 respondents (see table 1 for demographics). The final top 10 questions are shown in figure 1B.

DISCUSSION
We have undertaken the first JLA Priority Setting Partnership in CF, with equal numbers of participants from the patient and clinical communities. We have produced a top 10 list of clinical research questions in CF, which will be attractive to both researchers and funders.

There has been little previous work of this kind. The Italian Patient-Centered Outcomes CF Research Group canvassed opinion from 12 clinical researchers and eight ‘expert stakeholders’.9 This exercise prioritised five topics: transplantation, Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) modulators, P. aeruginosa, Burkholderia cepacia and allergic bronchopulmonary aspergillosis. The management of infection in CF also featured in our top 10—non-tuberculous mycobacteria (Q3) and P. aeruginosa (Q10). Infection with P. aeruginosa has previously been shown to be a great concern to parents and patients with CF.5 Surprisingly, questions on CFTR modulators were absent from our top 10. Respondents might assume this research will progress whether or not prioritised.

The strength of our study is our global reach and the large numbers of respondents, representing the whole CF community. Online surveys give less granular data than approaches such as focus groups. An
alternative approach to ours—focus groups, using video conferencing—would also avoid cross infection. Key aspects of our robust methodology were a representative steering committee, a systematic review of research gaps and having two researchers independently processing questions. Our experience of using online surveys, promotion through social media and video conferencing may be useful to other patient engagement exercises in respiratory medicine and beyond where face-to-face meetings are restricted by infection control, geography or frailty of participants. The US CF Foundation ‘Insight CF’ engagement programme has adopted a similar approach to registry-based research.

The items on the top 10 list are not prescriptive, rather they allow research groups to formulate a testable hypothesis with the appropriate study design. We are in discussion with the NIHR Evaluation, Trials and Studies Coordinating Centre about topics from our top 10, suitable for NIHR-commissioned research. We hope that this exercise will invigorate research in areas of shared importance to both the patient and the clinical community and demonstrate the value of involving the whole CF community in all steps of the research pathway.

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Contributors All authors contributed to the design, recruitment, data analysis and in the preparation of the manuscript.

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Provenance and peer review Not commissioned; externally peer reviewed.

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