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1 Attitudes towards epilepsy in the UK population: results from a 2018 national survey

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20 **Abstract**

21 Purpose: To measure stigma resulting from negative attitudes toward epilepsy in the United Kingdom
22 (UK) population.

23 Methods: An online survey of a stratified quota sample of UK adults in July 2018. The primary
24 outcome measure was the 46-item Attitudes and Beliefs about Living with Epilepsy (ABLE) scale,
25 scored on a five-point Likert scale. Items on sociodemographic characteristics, experience of epilepsy,
26 and knowledge of epilepsy were also included. Mean scores were calculated for the ABLE and
27 subscales: risk and safety concerns, personal fear and social avoidance, work and role expectations, and
28 negative stereotypes. Hierarchical regressions tested the association between mean ABLE and subscale
29 scores with sociodemographic and epilepsy related factors.

30 Results: 4,000 responded, 3875 responses were included in the analysis. Mean ABLE score was 2.28
31 (95% CI: 2.26-2.29) (1=no stigma, 5=high stigma). Subscales: risk and safety concerns 3.22 (95% CI:
32 3.20-3.25), personal fear and social avoidance 2.13 (95% CI: 2.11-2.16), work and role expectations
33 2.07 (95% CI: 2.05-2.09), and negative stereotypes 1.67 (95% CI: 1.65-1.69). Mean knowledge score
34 was 78% (95% CI: 76.15-77.02).

35 Conclusion: Findings of the first UK national survey of attitudes and beliefs about living with epilepsy
36 suggest relatively low stigma among the sampled population. The subscale with the least stigma was
37 negative stereotypes. Risk and safety concerns were associated with highest stigma. Improving public
38 knowledge about epilepsy has potential to reduce stigma, however this may also raise risk and safety
39 concerns. The results from this project could inform future work to improve awareness and
40 understanding of epilepsy.

41 **Keywords:**

42 Epilepsy, attitude, belief, stigma, survey, UK

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44 **Graphical Abstract**

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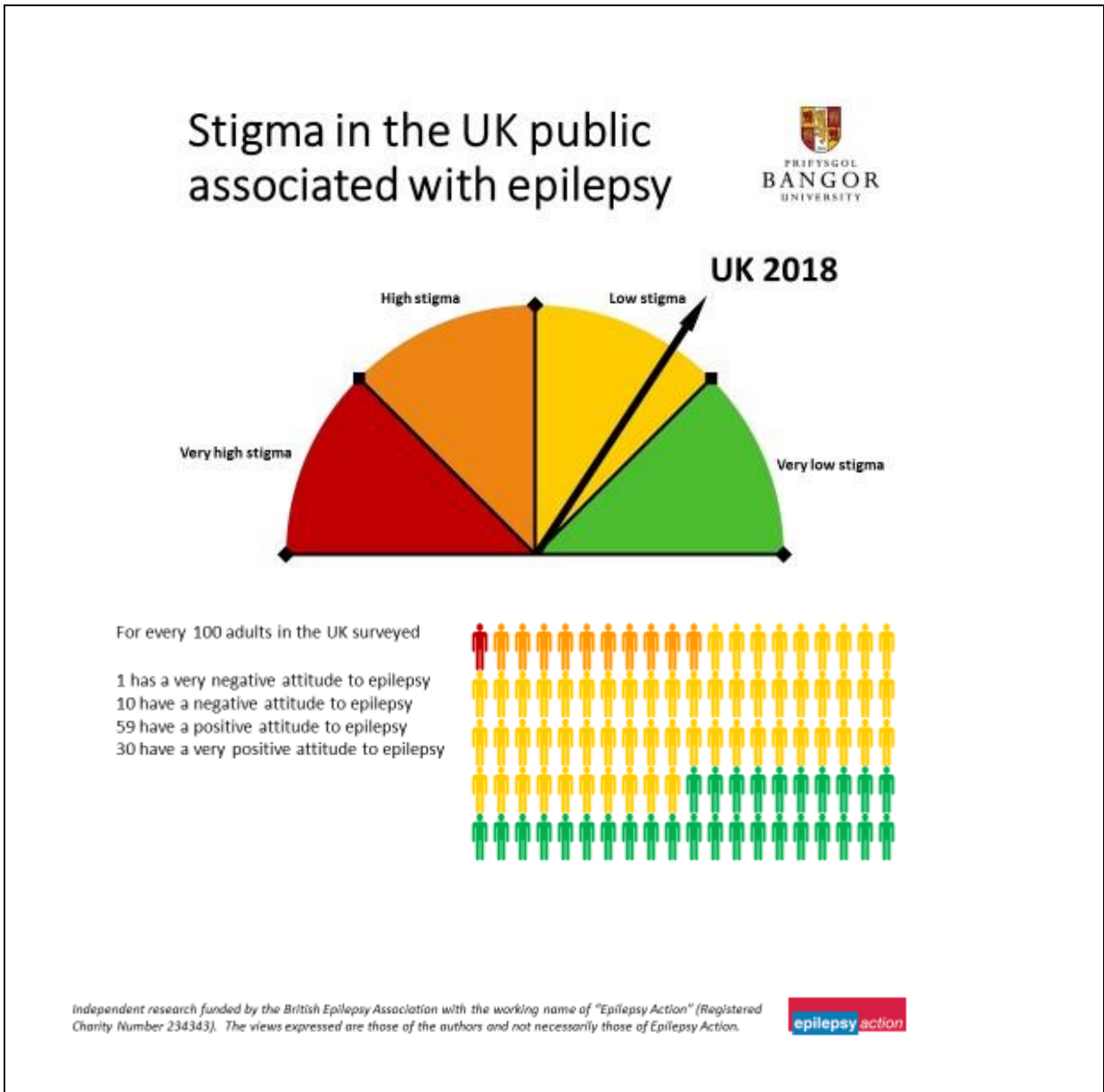
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63 **Introduction**

64 Epilepsy is a common chronic neurological condition affecting at least 50 million people globally ^[1].
65 People with epilepsy often experience stigma and discrimination because of the condition and as such,
66 their social and personal expectations are often restricted^[2]. The experience of stigma may even affect
67 their quality of life even more than the medical condition itself^[3]. Where people with epilepsy feel
68 prevented from living an ordinary life due to stigma resulting from negative attitudes, this may be an
69 internal perception rather than widely enacted discrimination^[4]. It is important to understand any stigma
70 that people with epilepsy may face. This may help to improve quality of life for people with epilepsy.

71 One in 103 people in the United Kingdom (UK) are living with epilepsy^[5] yet little is known about the
72 UK public attitude towards epilepsy and people with epilepsy. Research to date, has predominantly
73 focused on experience of stigma by people with epilepsy; and, evidence of the associated negative
74 attitude in society is more limited. Researchers in the United States (US) developed the Attitudes and
75 Beliefs about Living with Epilepsy (ABLE) scale to measure public attitudes to epilepsy and people
76 with epilepsy^[6,7] and have used this longitudinally to measure changes in public perceptions^[8]. Tracking
77 public attitudes over time has potential to assist patient organisations, such as Epilepsy Action in the
78 UK, plan future work to improve awareness and understanding of epilepsy and to evaluate the success
79 of such campaigns. Evidence on public attitudes may also contribute to the debate on internal
80 perceptions of stigma. Consequently, creating, strengthening and implementing policies to promote
81 access to information and knowledge may help reduce the stigma surrounding people living with
82 epilepsy.

83 The aim of our current study was to quantitatively assess public attitudes to epilepsy and people with
84 epilepsy in the UK in 2018. Specific objectives were to (i) measure public attitudes to epilepsy and
85 people with epilepsy in the UK using the ABLE scale^[6,7,8] by assessing levels of stigma related to: risk
86 and safety concerns, personal fear and social avoidance, work and role expectations, and negative
87 stereotypes. (ii) measure associations between stigma and sociodemographic characteristics, experience
88 of epilepsy, and knowledge of epilepsy. We hypothesised that attitude would vary by age, gender,
89 ethnicity, and education, consistent with previous findings^[6]. We also hypothesised that people with
90 experience of epilepsy (self or other) and/or higher knowledge of epilepsy would demonstrate less
91 stigma.

92 **Methods**

93 *Study design*

94 A cross-sectional web-based survey was conducted in the UK on a nationally representative sample of
95 adults aged 18 or over. . All data were collected in July 2018. The study was approved by Bangor
96 University Healthcare and Medical Sciences Academic Ethics Committee. The study was conducted
97 and reported according the STROBE checklist for cross-sectional studies^[9]. All data were analysed in
98 STATA 13 (StataCorp LLC, College Station, TX).

99 *Participants*

100 Members of the public were recruited using a consumer marketing panel. The panel is made up of pre-
101 recruited members of the public who have agreed to participate in online surveys. The company used
102 quota sampling to achieve a sample that was representative of the UK public, stratified by age, gender,
103 region and socioeconomic group. Respondents were required to check a series of boxes to confirm that
104 they had read and understood the participant information before proceeding to the questionnaire. The

105 participant information contained details relating to the survey (e.g. length of time to complete) and
106 participation (e.g. that participants may withdraw at any stage, confidentiality etc.). Respondents were
107 incentivised to participate by collecting points, tradable for goods (equivalent to £0.17).

108 *Outcomes*

109 The primary outcome measure was the ABLE scale^[6]. The ABLE scale contains 46-items to measure
110 stigma associated with epilepsy, in the general public. Twenty-three items assess cognitive beliefs about
111 people with epilepsy, which include items describing characteristics of a person with epilepsy, and the
112 abilities and limitations of people with epilepsy. Thirteen items assess affective reactions towards
113 people with epilepsy (e.g. discomfort, shame, fear, pity). Ten items assess respondent intentions toward
114 social distancing behaviours (e.g., would be nervous around a person with epilepsy because she or he
115 might have a seizure). All 46-items were included in the survey, assessed on a five point Likert scale
116 ranging from strongly disagree to strongly agree^[8], the underlying construct of each item varied in terms
117 of whether they were positively or negatively phrased. Secondary outcomes were four subscales,
118 constructed from items within the ABLE, that measure difference stigma domains: risk and safety
119 concerns (6-items), personal fear and social avoidance (8-items), work and role expectations (8 items),
120 and negative stereotypes (7 items).

121 The research team assessed the US version of the ABLE for UK language specificity. Five items were
122 adapted by replacing terms that had a direct equivalent in the UK (i.e. elementary school was replaced
123 to primary school), otherwise, by providing further explanation in parentheses alongside the original
124 US wording.

125 Additionally, questions were included to measure personal experience of epilepsy, attitude towards
126 epilepsy, knowledge about epilepsy, and sociodemographic characteristics. In addition to the
127 stratification variables of age, gender, region and socioeconomic class, sociodemographic factors
128 included ethnicity, employment status, education level, experience of marriage/civil partnership, having
129 children, and annual household income. A purposive review of epilepsy knowledge and experience
130 measures was conducted to determine items for inclusion in the questionnaire. The final questionnaire
131 contained 4-items on experience of epilepsy (self or other) and witness of seizures. As the purpose was
132 to assess whether the respondent had witnessed a seizure in real life, rather than on TV, this question
133 was phrased “*Have you ever seen someone have a seizure in person?*”. Respondent’s knowledge about
134 epilepsy, was tested using 15 true or false items and 1 multiple choice question adapted from the
135 Epilepsy Knowledge Profile-General^[10,11]. This section was specifically designed to assess knowledge
136 of the condition and its symptoms, characteristics, prevalence and treatment. Responses to each
137 question were mandatory to progress to the next item.

138 A pilot study was conducted in a convenience sample of 22 members of Bangor University and Epilepsy
139 Action employees. Twelve people completed the questionnaire and provided feedback on completion
140 time and comprehension of items. The mean completion time of the pilot questionnaire was 12 minutes.
141 In response to the pilot feedback, one ABLE item was rephrased, and knowledge questions were refined.
142 Overall, participants were asked to complete a maximum of 77 questions.

143 *Study sample*

144 The minimum sample size was defined as 2,401 responses, based on a 95% confidence level, an assumed
145 standard deviation in ABLE score of 1.25, and a confidence interval width of 0.1^[12].

146 To allow for analysis of multiple subgroups and to allow for potential irrational responders to be
147 removed, 4,000 responses were collected

148 Representativeness of the sample was assessed for the stratification variables, and socio demographic
149 characteristics by comparing participant responses with data for the general UK population^[13].
150 Response bias was assessed by identifying non-differentiation (straight lining) and inconsistency in
151 responses^[14]. Nondifferentiation occurs when respondents give identical answers to all items. Given
152 the nature of the design of the ABLE, with the direction of the underlying construct being reversed in
153 over 40% of questions, we considered a participant to be non-differentiating when they provided the
154 same response to over 41 (90%) of the questions within the ABLE. It was not possible to assess sample
155 bias or self-selection bias, as the required data was not available from the consumer marketing panel.

156 *Analysis of primary outcome*

157 Each item from the ABLE questionnaire was scored on a five-point Likert scale ranging from strongly
158 disagree to strongly agree, consistent with the most recent US application^[8]. Twenty- seven items were
159 reversed so that positively and negatively worded items were scored in the same direction as the
160 underlying scale construct. Thereafter, higher scores indicated more negative attitudes towards a person
161 with epilepsy^[6]. On review of the responses and in consultation with Epilepsy Action, scoring of a
162 further item, “*I believe people with epilepsy cannot have as good quality of life as people without*
163 *epilepsy*” was also reversed.

164 Mean score and 95% confidence intervals were generated for the ABLE scale (46-items) and four
165 subscales. Chronbach Alpha was reported as a measure of internal consistency for each scale.

166 *Analysis of sociodemographic, experience and knowledge items*

167 Frequencies and descriptive statistics were reported for demographic items and experience of epilepsy.
168 Subgroup analyses, using t-tests and ANOVAs were conducted to describe differences in responses
169 according to respondent characteristics. Items on knowledge were scored and summed such that higher
170 scores indicate a higher level of knowledge. Knowledge score was reported as a percentage of items
171 answered correctly. Association between actual knowledge (knowledge test score) and the ABLE item
172 “*I believe I know a lot about epilepsy*” was compared descriptively. Due to multiple comparisons
173 resulting in an increased risk of a false positive result, significant results were conservatively reported
174 at $p < 0.001$, following Bonferroni correction.

175 *Regression analysis*

176 Hierarchical regression was used to adjust for any confounders, and assess how sociodemographic
177 characteristics, experience and knowledge were able to explain stigma, multivariately. Responses of
178 “*unknown*” or “*prefer not to say*” were treated as missing in the regression. Regression analyses were
179 performed for the ABLE scale, and each subscale.

180 **Results**

181 Four thousand responses were received and examined for eligibility. All responses were complete, with
182 no missing data. One-hundred and twenty-five respondents were excluded from analysis due to
183 nondifferentiation, the remaining 3875 were included in the analysis.

184 *Descriptive data: Demographics*

185 The mean age of the sample was 49 years (range 18 to 92). Sample demographics (Table 1) were
186 representative of the UK general public with our sample falling within 3% of UK proportions for each
187 of the stratification variables. One-hundred and eighty-nine people with epilepsy or a seizure disorder
188 completed the survey; which is higher than estimates based on UK prevalence of epilepsy alone. Less
189 than half the sample knew someone with epilepsy, most commonly a friend or family member. Just
190 over half had witnessed a seizure in their lifetime; one in four had witnessed a seizure within the last 5-
191 years.

192 Outcome data: ABLE scale

193 The population mean for the 46-item ABLE scale questions was 2.28 (95% CI:2.26-2.29) (Table 2).
194 When considering the mean scores of each of the 3875 respondents, 5 (<1%) have a very negative
195 attitude to epilepsy (ABLE score 4+), 396 (10%) have a negative attitude to epilepsy (ABLE score
196 between 3&3.9), 2317 (59%) have a positive attitude to epilepsy (ABLE score between 2&2.9), and
197 1157 (30%) have a very positive attitude to epilepsy (ABLE score less than 2). Our findings therefore
198 suggest that the average person in the UK has a somewhat positive attitude toward epilepsy, in that they
199 agree (but do not strongly agree) with positive statements. Items from the ABLE scale showed very
200 good internal consistency, (Chronbach alpha 0.94), consistent with DiIorio and colleagues^[6].

201 Mean scores for individual items suggest that seven items were associated with a negative attitude
202 (supplementary material 1; score >3 in Table 2). Items associated with the most negative attitude were
203 “*I believe I know a lot about epilepsy*” (57% disagreed with this statement) and “*I would let my child*
204 *ride in a car with a driver who has epilepsy*” (50% disagreed with this statement). In general,
205 respondents expressed a level of agreement or disagreement to each item, there were only 9 items where
206 the modal response category was “*uncertain*”. “*I believe people with epilepsy can safely operate heavy*
207 *machinery*” had a mean score of 3 (95%CI:2.97-3.03), with 42% of respondents selecting the
208 “*uncertain*” option, suggesting the average person in the UK is uncertain of their opinion on this.

209 *Subscales*

210 Internal consistency was “*good*” or better for all four subscales (Table 2). The highest score was on the
211 risk and safety concerns domain (mean=3.22, 95%CI: 3.20-3.25). Among the items describing risk and
212 safety concerns, four had average scores that indicated negative responses (mean item score >3). All
213 four of these items were associated with driving and/or children. Among the items in the personal fear
214 and social avoidance subscale (mean 2.13, 95%CI: 2.11-2.16) the majority of respondents indicated
215 they would not be embarrassed if someone in their family had epilepsy. A lower score on the work and
216 role expectations subscale (mean 2.07, 95% CI: 2.05-2.09), indicated a perceived work normality for
217 people with epilepsy. The negative stereotypes subscale had the least negativity of all subscales (mean
218 1.67, 95% CI: 1.65-1.69), indicating that negative stereotyping is relatively low in the UK. All seven
219 items in this subscale had mean scores that indicated agreement with positive statements, or
220 disagreement with negative statements.

221 *Epilepsy knowledge*

222 Percentages of people responding correctly to each of the knowledge items are listed in table 3. The
223 mean overall knowledge score was 77% (95% CI: 76.15-77.02). Less than half of respondents
224 responded correctly to questions on prevalence, seizure first-aid, and epilepsy manifestations. Most
225 people were aware that epilepsy is not contagious (92%). A quarter of respondents did not know that

226 people with epilepsy are protected by the Equality Act. We explored association between actual
227 knowledge and the ABLE item “*I believe I know a lot about epilepsy*”. Perceived knowledge (ABLE
228 item “*I believe I know a lot about epilepsy*”) negatively predicted actual knowledge (knowledge test
229 score). People who strongly agreed with the ABLE statement had a lower mean knowledge test score
230 (68%).

231 *Subgroup analysis*

232 Univariate, statistically significant differences in mean attitude and subscale scores were identified for
233 gender, employment, marriage, children, people with epilepsy, people who knew someone with
234 epilepsy, and witnesses of seizure (Table 4).

235 Using ANOVA, statistically significant differences in attitude between age categories were observed
236 for all scales ($p \leq 0.001$) (Figure 1). Men had attitudes that are more negative on all scales, with the
237 exception of risk and safety concerns. People in employment had significantly lower risk and safety
238 concerns but more fear and social avoidance, and negative stereotyping, compared to people not in paid
239 employment. Risk and safety concerns were also lower for respondents who had never been married or
240 in a civil partnership and respondents with children.

241 People with epilepsy had significantly lower risk and safety concerns, but significantly higher scores
242 (were more negative) on all other subscales, compared to people who do not have epilepsy. Whereas,
243 people who knew someone with epilepsy had significantly higher risk and safety concerns, more
244 personal fear and social avoidance, and more negative work and role expectations, than people who did
245 not know anyone with epilepsy. Those who had witnessed a seizure had significantly less negativity
246 on all domains other than negative stereotypes, but both groups had low stigma on this domain.

247 Despite reaching statistical significance, the actual difference in mean score between groups was quite
248 small and explained a limited amount of the variance in stigma (less than 5%).

249

250 *Hierarchical regression*

251 The results of the regression between attitude towards epilepsy and other factors are summarised in
252 table 5, for the full ABLE scale and subscales. Multivariate results were largely consistent with
253 univariate results (subgroup analyses). Overall, more negative attitudes were significantly associated
254 with sociodemographic factors (age <35 or >65 , male), experience of epilepsy (not knowing a person
255 with epilepsy, never having witnessed a seizure) and having lower epilepsy knowledge. Experience of
256 epilepsy was the greatest contributor to risk and safety concerns, interestingly, people with epilepsy had
257 significantly lower risk and safety concerns.

258 These models explained a limited amount of the variance in attitude (between 8% and 48%). The ABLE
259 scale model explains 35% of the variance in attitude. The sociodemographic factors entered into the
260 regression model explained 9%, experience factors explain a further 2%, and adding knowledge about
261 epilepsy explained a further 25%. Across subscales sociodemographic explained no more than 14% of
262 the variance around the score, and experience added no more than 4%. Knowledge was the largest
263 contributor, in all models except risk and safety concerns. Ranking of individual knowledge factors
264 within the regression analyses are presented in Table 3. Lower knowledge of the most commonly known
265 items appears to predict more negative attitude, with the exception of the risk and safety concerns

266 domain, where higher knowledge of three items was associated with higher concerns (epilepsy is not
267 contagious; seizure types vary; and you cannot tell by looking at someone with they have epilepsy).

268 **Discussion**

269 *Key results*

270 Stigma toward epilepsy and people with epilepsy was present among the sampled UK population,
271 however the findings suggests the attitude of the average person is associated with relatively low stigma.
272 Attitudes differed by subscales. The least negative subscale was negative stereotypes, suggesting that
273 the average respondent disagreed or strongly disagreed with statements on negative stereotypes. Risk
274 and safety concerns were associated with the most stigma.

275 Sociodemographic and experience factors explain a limited proportion of the variance in stigma score
276 and across domains. Knowledge factors contributed the most explanation of negative attitude, however,
277 the direction of effect varied. Overall, lower knowledge was associated with more negative attitude,
278 with the exception of risk and safety concerns, where higher knowledge in some areas was associated
279 with higher concerns.

280 *Interpretation*

281 Reducing stigma is a major activity of patient support groups globally^[15]. Epilepsy Action, in the UK,
282 has a goal to improve awareness and understanding of epilepsy and our findings suggest that this has
283 the potential to reduce stigma. In the current study, 95% of the population scored between 44-100% on
284 the knowledge test. Less than half the people surveyed were aware of how many people are living with
285 epilepsy in the UK, knew how to respond when someone was having a seizure, or, understood that
286 seizures can have different triggers. Only a quarter of respondents were aware that people with epilepsy
287 are protected by the Equality Act, which may also allude to a lack of understanding about the
288 classification of disability under this act. Whilst 20% of people believed they knew a lot about epilepsy,
289 of interest was that respondents who believed they knew a lot about epilepsy had a significantly lower
290 knowledge score. Researchers in other fields have suggested that individuals who express a belief that
291 they are more knowledgeable and may have a perception that they are better informed, despite
292 displaying higher evidence gaps between their perceived and actual knowledge^[16]. This finding
293 warrants further research in the context of epilepsy.

294 When assessing the relationship between stigma and knowledge, an incorrect response to the most
295 commonly known items was associated with greatest stigma; this suggests that increasing the
296 knowledge of this minority may have the greatest impact. Association between attitude and knowledge
297 reversed on some items in the risk and safety domain, suggesting that higher knowledge was associated
298 with being more risk averse. Furthermore, the most negative finding on work and role expectations,
299 related to safety – which concurs with the findings of the risk and safety are of highest concern to the
300 UK public. Future agendas should therefore focus on both improving knowledge, addressing risk and
301 safety concerns, and dispelling misconceptions.

302 Stigma in the UK was found to be relatively low, with ~90% of the UK public having a neutral to
303 positive attitude. This evidence may contribute towards the debate on internal perceptions and enacted
304 stigma. Whilst the concepts of perceived and enacted stigma represent different constructs, targets for
305 intervention may share some common ground. In the UK, Taylor and colleagues reported that 54% of
306 people with newly diagnosed epilepsy (n=1566) indicated feeling stigmatised; and, that reduced
307 mastery, younger age (<50), and seizures frequency, were amongst factors significantly associated with

308 this feeling^[17]. A recent systematic review of 25 quantitative studies of correlates of stigma in epilepsy,
309 also found that stigma was associated with demographic, illness-related, and psychological factors; and,
310 that these associations were highly culturally specific^[18]. This suggests that campaigns targeting factors
311 associated with public attitudes and knowledge, may also have a positive impact on people with
312 epilepsy. Furthermore, previous research has demonstrated the benefits of people with epilepsy
313 themselves being involved target audience-directed content and mode of delivery^[19]. This warrants
314 further exploration in the interpretation of the current findings and development of future campaigns.

315 *Comparison with other studies*

316 To our knowledge our study is the first application of the ABLE scale in the UK. A review of the
317 literature identified four studies reporting the results of the ABLE scale^[6,7,8,20]. Direct comparison across
318 all studies is not possible, due to heterogeneity in Likert scales^[6,7], subscale content^[20], and population.
319 The findings of this study, however, can be compared with US public attitudes on fear and social
320 avoidance, where the US public were found to be slightly more negative (2.18 versus 2.13)^[8]. Our
321 results also follows the general trend of previous studies, in that risk and safety concerns are associated
322 with the most stigma, and negative stereotypes are associated with the least stigma. Health-related
323 stigma has been measured for several other conditions, for example HIV/AIDS, leprosy, tuberculosis,
324 mental health, obesity^[21,22,23,24]. The literature suggests that the consequences of stigma being
325 remarkably similar between conditions and cultures, however, instruments to measure stigma tend to
326 be condition specific^[24]. The Time to Change anti-stigma campaign in mental-illness-related public
327 stigma among the English population^[25] is based on the “Attitudes to Mental Illness” national survey.
328 This includes 26 attitude items derived from the Community attitudes toward the Mentally Ill scales
329 and an additional item on employment-related attitudes. There are two subscales: prejudice and
330 exclusion, and tolerance and support; in this case higher scores indicate more positive attitudes.
331 Attitudes to mental illness in the English population in 2013 were in the region of 3.9 to 4.0, where 5.0
332 is positive^[25]. When compared to our findings of epilepsy stigma in the UK, the levels appear to be in
333 the same region (2.1 to 2.0 when scaled in the direction of 1.0 being the most positive).

334 *Strengths*

335 Our study has several key strengths. Firstly, we surveyed a large sample of the UK general population,
336 representative of age, gender, region and socioeconomic group. Secondly, our analysis considered the
337 distinction between what could predict negative attitude across all of the stigma domains (subscales).
338 Thirdly, we considered the contribution of distal and more proximal factors associated with attitude
339 (e.g. sociodemographic, experience and knowledge). Finally, we conducted further exploratory analysis
340 on more modifiable factors such as knowledge. Our consideration of perceived and actual knowledge
341 also gave insight to both the need to improve knowledge and address potential misconceptions.

342 *Limitations*

343 There are several limitations to our analysis. Firstly, our study used quota sample, we must therefore
344 acknowledge the risk of selection bias. The survey was administered online, via a consumer marketing
345 panel which may influence the results insofar as only people who were actively interested in completing
346 web-based surveys participated, which may reduce the external validity of our findings. Review of our
347 stratification variables confirmed that our sample was representative of the UK general population in
348 terms of age, gender, region, and socioeconomic class; and our approach to sampling was balanced
349 against efficient data collection. Secondly, a pragmatic approach was taken to assessing rational
350 responses, as with all surveys there is potential for false responses. Thirdly, sociodemographic
351 groupings used in the regression models were often broad (e.g. ethnic group), therefore, we cannot make

352 any inferences on any specific categories within groups, which may be associated with attitudes that are
353 more negative. Finally, the ABLE instrument was developed using attitudes of the US population in
354 2002^[6]. We identified six ABLE-items that required changes to reflect UK English language; we also
355 recoded a single item to represent a UK perspective on what constitutes stigma. This reduces
356 comparability of our results with US applications but increases reliability in our UK context.

357 *Conclusion*

358 Findings of the first UK national survey of attitudes and beliefs about living with epilepsy suggest
359 relatively low stigma among the sampled population. The subscale with the least stigma was negative
360 stereotypes, suggesting that the average respondent disagreed or strongly disagreed with statements on
361 negative stereotypes. Risk and safety concerns were associated with highest stigma. Attitude and
362 knowledge of epilepsy significantly correlate, suggesting that improving public knowledge about
363 epilepsy has potential to reduce stigma. The results from this project have potential to inform future
364 work to improve awareness and understanding of epilepsy.

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367 interpretation of this survey; and, thank the members of the consumer panel who participated in this
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Table 1: Sociodemographic and experience characteristics of study sample

Sociodemographic characteristics	
Age: Mean (SD)	49 (17)
Gender: N (%)	
Male	1,821 (47)
Female	2,040 (53)
Unknown (Other; Prefer not to say) *	14 (<1)
Ethnicity: N (%)**	
Asian or Asian British	189 (5)
Black or Black British	98 (3)
Chinese or Chinese British	33 (1)
British Mixed heritage	74 (2)
White	3,400 (88)
Unknown (Any other ethnic background; Prefer not to say)*	81 (2)
Employment: N (%)	
Employed	2107 (54)
Not in paid employment	1768 (46)
Education: N (%)	
Compulsory education	1240 (32)
Further education (i.e.. A-level, technical or professional qualification)	1332 (34)
Higher education (i.e. University degree)	1303 (34)
Have you ever been married or in a civil partnership: N (%)	
Yes	2,584 (67)
No	1,291 (33)
Do you have children: N (%)	
Yes	2,393 (62)
No	1,482 (38)
Household income: N (%)	
Low (under £19,999)	1,063 (27)
Medium (£20,000 - £39,999)	1,314 (34)
High (over £40,000)	1056 (27)
Unknown (Don't know; Prefer not to say)*	442 (11)
Do you consider yourself to have a disability or a long-term health condition: N (%)	
Yes	1,199 (31)
No	2,601 (67)
Prefer not to say *	75 (2)
Do you have epilepsy or a seizure condition: N (%)	
Yes	189 (5)
No	3686 (95)
Do you know someone with epilepsy or a seizure condition: N (%)	
Yes	1730 (45)
No	2,145 (55)
Have you ever witnessed a seizure?	
Yes	2,117 (55)
No	1,758 (45)

437 *Treated as missing in the regression analyses. ** The ethnic groupings used here are broad; there is
438 no breakdown of data for the more specific ethnic groups each contains. Some of the specific ethnic
439 groups have very different experiences to one another ([https://guide.ethnicity-
440 factsfigures.service.gov.uk/a-z](https://guide.ethnicity-factsfigures.service.gov.uk/a-z)).

Table

Table 2: ABLE mean scores. Higher scores represent more negative attitudes (range 1 to 5)

ABLE subscale and items	Mean	Confidence interval (95%)	
ABLE Score (Chronbach alpha 0.94)	2.28	2.26	2.29
Risk and Safety (Chronbach alpha 0.85)	3.22	3.20	3.25
I would let my child ride in a car with a driver who has epilepsy	3.50	3.46	3.53
I would feel comfortable if my child rode in a car with a driver who has epilepsy	3.44	3.40	3.47
I would ride in a car if the driver has epilepsy	3.33	3.30	3.36
I would hire someone with epilepsy to babysit my infant child	3.23	3.20	3.27
I believe people with epilepsy can safely operate heavy machinery.	3.00	2.97	3.03
*I believe people with epilepsy should not drive, even if the DVLA allows them to get a driving license	2.84	2.80	2.87
Personal Fear and Social Avoidance (Chronbach alpha 0.90)	2.13	2.11	2.16
*I would be afraid to be alone with a person with epilepsy	2.57	2.54	2.61
*I would be nervous to be around a person with epilepsy because he or she might have a seizure	2.51	2.48	2.55
*I would NOT want my child to date someone with epilepsy	2.24	2.21	2.27
*I would avoid a person with epilepsy who has frequent seizures	2.16	2.13	2.20
*I believe being around a person with epilepsy would make me uncomfortable	2.09	2.06	2.12
*I would NOT go out again with a person I just started dating if I found out he or she has epilepsy	1.97	1.94	2.00
*I would NOT want to work with someone with epilepsy	1.85	1.82	1.88
*I would be embarrassed if someone in my family had epilepsy	1.67	1.63	1.70
Work and role expectations (Chronbach alpha 0.79)	2.07	2.05	2.09
*I believe there are many work activities people with epilepsy cannot do safely that I can do safely	2.84	2.80	2.87
I would expect just as much from people with epilepsy as from others	2.16	2.13	2.19
*I believe people with epilepsy cannot have as good quality of life as people without epilepsy	2.11	2.08	2.15
I believe people with epilepsy can do anything I can do	2.03	2.00	2.06
I believe people with epilepsy can work 40 hours per week	1.99	1.97	2.02
I believe people with epilepsy are able to cope with everyday life as well as other people	1.94	1.92	1.97
I believe people with epilepsy can lead normal lives	1.77	1.74	1.79
I believe people with epilepsy can be as successful at work as others	1.70	1.68	1.72
Negative stereotypes (Chronbach alpha 0.92)	1.67	1.65	1.69
*I believe people with epilepsy should NOT have biological children.	1.85	1.82	1.88
*I believe people with epilepsy have a mental health problem	1.85	1.82	1.88
*I believe people with epilepsy are unreliable.	1.81	1.78	1.84
*I believe people with epilepsy are NOT as smart as other people who do not have epilepsy.	1.61	1.58	1.64
*I believe people with epilepsy should NOT marry	1.58	1.55	1.61
*I would consider getting a divorce if my spouse were diagnosed with epilepsy.	1.54	1.51	1.57
*I believe people with epilepsy are possessed by a supernatural spirit.	1.45	1.43	1.48

*Item coded in reverse to match direction of underlying construct

Table 3: Results of knowledge questionnaire, and rank order of association between attitude score and lower knowledge of item (1= largest impact on stigma).

Knowledge item (<i>correct answer</i>)	Correct responses: n of 3875 (%)	ABLE score	Risk and safety	Fear and social	Work and role	Negative stereotypes
K13. You can catch epilepsy from someone who has it (<i>False</i>)	3,564 (92)	3		1	1	2
K15. All people with epilepsy have the same type of seizure (<i>False</i>)	3,507 (91)	1		3	3	1
K14. You can tell from looking at someone if they have epilepsy (<i>False</i>)	3,515 (91)	2		2	4	3
K10. With treatment, most people with epilepsy can go a year or more without a seizure (<i>True</i>)	3,431 (89)	4		4	7	4
K2. All people with epilepsy lose consciousness during seizures (<i>False</i>)	2,846, (73)	5	2	5	5	5
K4. For most people with epilepsy, seizures are well controlled with drug treatment (<i>True</i>)	3,492 (90)	6			2	
K11. There is a high likelihood of death every time a person with epilepsy has a seizure (<i>False</i>)	2,791 (72)	7	15		6	8
K12. Some seizures may last for a matter of seconds and may not be noticed by others (<i>True</i>)	3,477 (90)	9			8	6
K3. Some people get a warning or feeling shortly before a seizure(<i>True</i>)	3,337 (86)	10		6		7
K6. Anyone can develop epilepsy at any time (<i>True</i>)	2,629 (68)	11		8	9	
K7. When you see someone having a seizure, you should put some put something in the person's mouth to prevent the person from biting or swallowing their tongue (<i>False</i>)	1,778 (46)	8		7		10
K1. A seizure can be described as an abnormality in the function of the nerve cells in the brain (<i>True</i>)	3,543 (91)					
K8. Most people with epilepsy have seizures when looking at flashing lights (<i>False</i>)	1,811 (47)	12				
K9. People with epilepsy are protected by the Equality Act (<i>True</i>)	2,915 (75)					9
K5. For people with epilepsy, stress may cause some seizures (<i>True</i>)	3,444 (89)					
K16. Approximately how many people do you think are affected by epilepsy in UK (<i>1 in 100</i>)*	1,404 (36)					

*Options were 1 in 10, 1 in 100, 1 in 1,000, 1 in 10,000. **Note:** In regression N=3368. Rank score = incorrect response associated with more negative attitude (1=greatest association). i.e. correct response associated with increased concern or more negative stereotypes. Blank cells = did not reach statistical significance i.e. there no associated with stigma in specified domain.

Table 4: Mean able scores of subgroups

	ABLE	Risk and safety concerns	Fear and social avoidance	Work and role expectations	Negative stereotypes
	Mean score (95% CI)	Mean score (95% CI)	Mean score (95% CI)	Mean score (95% CI)	Mean score (95% CI)
Male	2.32* (2.30-2.34)	3.18 (3.14-3.22)	2.22* (2.18-2.26)	2.1* (2.07-2.12)	1.75* (1.71-1.79)
Female	2.24* (2.22-2.26)	3.26 (3.22-3.29)	2.05* (2.02-2.09)	2.03* (2.01-2.07)	1.60* (1.57 -1.62)
Employed	2.30(2.27-2.32)	3.14* (3.10-3.17)	2.17* (2.14-2.21)	2.07 (2.05-2.10)	1.76* (1.72-1.79)
Not in paid employment	2.25 (2.23-2.28)	3.33* (3.29-3.36)	2.08* (2.05-2.12)	2.06 (2.04-2.09)	1.57* (1.54-1.60)
Compulsory education	2.29 (2.26-2.31)	3.30 (3.26-3.35)	2.12 (2.08-2.17)	2.08 (2.04-2.11)	1.65 (1.16-1.69)
Further/higher education	2.27 (2.25-2.29)	3.19 (3.15-3.22)	2.14 (2.11-2.17)	2.06 (2.04-2.09)	1.68 (1.65-1.71)
Been in marriage or civil partnership	2.28 (2.27-2.30)	3.25* (3.223.29)	2.16* (2.132.19)	2.07 (2.05-2.10)	1.68 (1.65-1.71)
Never been in marriage or civil partnership	2.26 (2.23-2.29)	3.16* (3.11-3.20)	2.07* (2.03-2.11)	2.06 (2.02-2.09)	1.66 (1.61-1.70)
Have children – yes	2.27 (2.25-2.29)	3.27* (3.233.30)	2.14 (2.11-2.17)	2.06 (2.04 -2.08)	1.66 (1.63-1.69)
Have children – no	2.28 (2.26-2.30)	3.15* (3.11-3.20)	2.12 (2.08-2.16)	2.08 (2.05-2.11)	1.69 (1.65-1.73)
Annual household income					
Low	2.29 (2.26-2.32)	3.03 (3.26-3.35)	2.11 (2.06-2.16)	2.10 (2.07-2.14)	1.64 (1.60-1.68)
Medium	2.26 (2.24-2.29)	3.22 (3.18- 3.26)	2.12 (2.07-2.16)	2.06 (2.03-2.09)	1.65 (1.61-1.69)
High	2.30 (2.27-2.34)	3.12 (3.07-3.17)	2.21 (2.16- 2.27)	2.06 (2.02-2.09)	1.78 (1.72-1.84)
Person with epilepsy – yes	2.51* (2.42-2.60)	2.86 * (2.74-2.98)	2.53* (2.09-2.14)	2.18 (2.09-2.27)	2.43* (2.24-2.62)
Person with epilepsy – no	2.7 * (2.25-2.28)	3.24 * (3.22-3.27)	2.11* (2.36-2.70)	2.06 (2.04-2.08)	1.63* (1.61-1.65)
Know someone with epilepsy* - yes	2.14* (2.19-2.24)	3.29 * (3.25-3.32)	2.23* (2.20-2.26)	2.10 * (2.07-2.12)	1.68 1.64-1.71)
Know someone with epilepsy* - no	2.33* (2.31-2.35)	3.14* (3.10-3.18)	2.01* (1.98-2.05)	2.03* (2.00-2.06)	1.66 (1.62-1.70)
Has witnessed a seizure - yes	2.25* (2.23-2.27)	3.19 (3.16-3.23)	2.08* (2.04-2.11)	2.05 (2.03 -2.07)	1.70 (1.66-1.74)
Never witnessed a seizure	2.31* (2.29-2.33)	3.26 (3.22-3.30)	2.20* (2.16-2.23)	2.09 (2.063-2.12)	1.63 (1.60-1.67)

*Significant result at $p < 0.001$

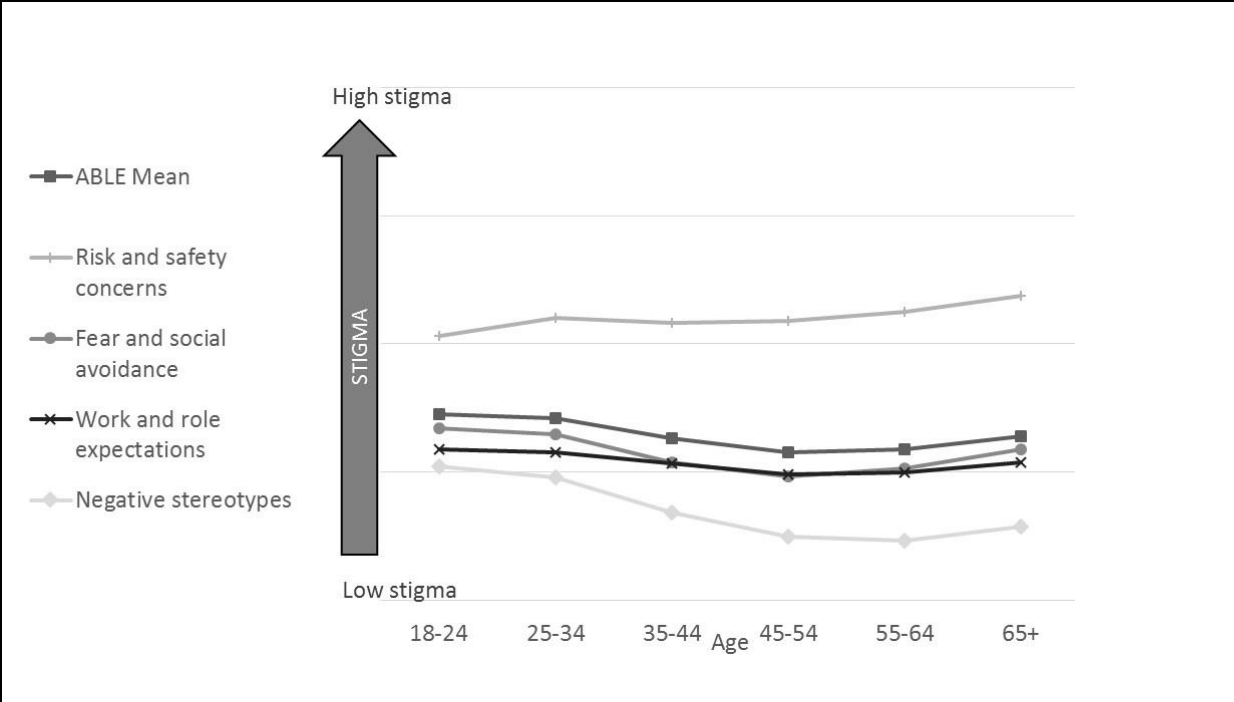
**Other than self.

Table 5: Regression models

	ABLE score	Risk and safety concerns	Fear and social avoidance	Work and role expectations	Negative stereotypes
Sociodemographic factors					
Age: base category = 45-54 years					
18-24 years	0.0897	-0.0407	0.102	0.00956	0.151*
25-34 years	0.0810	0.0629	0.068	0.0141	0.119
35-44 years	0.0391	0.00549	0.0196	0.0354	0.04
55-64 years	0.0475	0.0517	0.102	0.0438	0.014
65 years +	0.134*	0.117	0.263*	0.0893	0.132*
Gender: male	0.0636*	-0.0794	0.0996*	0.0572	0.136*
Ethnicity: base category = white					
Asian	0.0700	0.0107	0.0984	0.0037	0.093
Black	0.0281	0.0979	-0.0467	0.102	-0.0365
Chinese	0.0951	0.177	0.142	0.0143	0.0696
Mixed	-0.00269	-0.122	-0.0313	-0.0134	0.0345
Employment: in employment	0.00851	-0.0955	0.0653	-0.0154	0.0559
Education: further/higher	0.00453	-0.0269	0.0337	0.0198	0.0151
Marriage/civil partnership	0.0323	0.0243	0.0542	0.0351	0.0605
Children	-0.002	0.0795	0.0109	-0.0203	-0.0237
Household income: lower	-0.0216	-0.0537	0.012	-0.0593	-0.0063
R² (stage 1)	0.088	0.032	0.073	0.034	0.143
Experience factors					
Has epilepsy	-0.00882	-0.345*	0.0845	-0.112	0.330**
Knows someone with epilepsy	-0.105*	-0.113*	-0.199*	-0.0517	-0.0674*
Has witnessed a seizure	-0.0542*	0.00825	-0.109*	-0.0366	0.00497
R² (stage 2)	0.112	0.046	0.108	0.039	0.180
Knowledge factors (see table 6 for full wording)					
k1: abnormality of the function of the brain	0.0486	0.0487	0.0860	0.0644	-0.019
k2: people lose consciousness during a seizure	-0.167*	-0.155*	-0.204*	-0.139*	-0.214*
k3: some people get a warning before a seizure	-0.0868*	0.00166	-0.147*	-0.0584	-0.123*
k4: for most seizures are well controlled ...	-0.118*	-0.101	-0.108	-0.164*	-0.117
k5: stress may cause some seizures	-0.00717	0.0747	-0.0386	0.0191	-0.0787
k6: anyone can develop epilepsy...	-0.0742*	-0.0919	-0.0883*	-0.0920*	-0.0419
k7: put something in the person's mouth	-0.0921*	-0.0448	-0.137*	-0.0620	-0.0770*
k8: have seizures when looking at flashing lights	-0.0497*	-0.0801	-0.0412	-0.031	-0.0358
k9: protected by the Equality Act	-0.0517	-0.000693	-0.0547	-0.0660	-0.0810*
k10: with treatment most people can ...	-0.173*	-0.0785	-0.269*	-0.130*	-0.233*
k11: high likelihood of death every time a person ...	-0.112*	-0.149*	-0.0890	-0.137*	-0.116*
k12: seizures may last for a matter of seconds ...	-0.0886*	0.0366	-0.0783	-0.109*	-0.153*
k13: you can catch epilepsy ...	-0.216*	0.216	-0.404*	-0.180*	-0.510*
k14: you can tell by looking at someone ...	-0.217*	0.169	-0.383*	-0.151*	-0.488*
k15: all people have same type of seizure ...	-0.235*	0.160	-0.378*	-0.161*	-0.523*
k16: Prevalence of epilepsy in the UK	0.025	0.00013	0.0404	0.00568	0.0623
R² (stage 3)	0.354	0.082	0.309	0.179	0.481
Other unobserved factors					
_cons	3.564*	3.158*	3.955*	3.187*	3.865*
Model statistics					
N	3368	3368	3368	3368	3368
R-sq	0.354	0.082	0.309	0.179	0.481
adj. R-sq	0.348	0.073	0.302	0.171	0.476
rmse	0.413	0.797	0.683	0.526	0.58

*Significant result at p≤0.01

Figure 1: Attitude by age groups



Supplementary material 1: ABLÉ item responses

Percentage of respondents who agreed with each statement, in order of worst to best: n of 3785 (%)

	Strongly disagree					Strongly Agree
	1	2	3	4	5	
I believe I know a lot about epilepsy	796 (21%)	1,408 (36%)	915 (24%)	586 (15%)	170 (4%)	
I would let my child ride in a car with a driver who has epilepsy	893 (23%)	1,054 (27%)	1,199 (31%)	552 (14%)	177 (5%)	
I would feel comfortable if my child rode in a car with a driver who has epilepsy	785 (20%)	1,046 (27%)	1,291 (33%)	588 (15%)	165 (4%)	
I believe people being treated for epilepsy can have a seizure at any time*	83 (2%)	489 (13%)	1,380 (36%)	1,530 (39%)	393 (10%)	
I would ride in a car if the driver has epilepsy	664 (17%)	1,008 (26%)	1,308 (34%)	731 (19%)	164 (4%)	
I would hire someone with epilepsy to baby-sit my infant child	503 (13%)	908 (23%)	1,620 (42%)	683 (18%)	161 (4%)	
I believe seizures in people with epilepsy require emergency medical assistance*	145 (4%)	1,045 (27%)	1,334 (34%)	1,033 (27%)	318 (8%)	
I believe people with epilepsy can safely operate heavy machinery	296 (8%)	801 (21%)	1,644 (42%)	885 (23%)	249 (6%)	
I believe I feel sorry for people who have epilepsy*	601 (16%)	1,024 (26%)	865 (22%)	1,122 (29%)	263 (7%)	
I believe there are many work activities people with epilepsy cannot do safely that I can do safely*	444 (11%)	965 (25%)	1,446 (37%)	816 (21%)	204 (5%)	
I believe people with epilepsy should not drive, even if the DVLA allows them to get a driving license*	498 (13%)	1,019 (26%)	1,287 (33%)	760 (20%)	311 (8%)	
I would be worried that a seizure could happen at any time if I were around a person with epilepsy*	467 (12%)	1,267 (33%)	823 (21%)	1,075 (28%)	243 (6%)	
I would feel comfortable if my child were in primary school class in which the teacher has epilepsy	204 (5%)	599 (15%)	1,377 (36%)	1,327 (34%)	358 (10%)	
I would be afraid to be alone with a person with epilepsy*	749 (19%)	1,358 (35%)	772 (20%)	792 (20%)	204 (5%)	
I would be nervous to be around a person with epilepsy because he or she might have a seizure*	730 (19%)	1,543 (40%)	673 (17%)	750 (19%)	179 (5%)	
I believe I am prepared to help a person with epilepsy who is having a seizure	181 (5%)	407 (11%)	774 (20%)	1,712 (44%)	801 (21%)	
I believe if people with epilepsy have seizures, it is because they are not doing what their doctor tells them to do*	1,011 (26%)	1,469 (38%)	918 (24%)	346 (9%)	131 (3%)	

I would NOT want my child to date someone with epilepsy*	1,024 (26%)	1,460 (38%)	955 (25%)	302 (8%)	134 (3%)
I believe people with epilepsy should NOT be primary school teachers*	1,061 (27%)	1,472 (38%)	855 (22%)	369 (10%)	118 (3%)
I would avoid a person with epilepsy who has frequent seizures*	1,225 (32%)	1,423 (37%)	719 (19%)	392 (10%)	116 (3%)
I would expect just as much from people with epilepsy as from others	55 (1%)	269 (7%)	702 (18%)	2,065 (53%)	784 (20%)
I believe people with epilepsy have trouble managing their day to day activities*	1,209 (31%)	1,505 (39%)	753 (19%)	321 (8%)	87 (2%)
I believe people with epilepsy cannot have as good quality of life as people without epilepsy*	1,286 (33%)	1,461 (38%)	654 (17%)	348 (9%)	126 (3%)
I believe being around a person with epilepsy would make me uncomfortable*	1,254 (32%)	1,566 (40%)	605 (16%)	345 (9%)	105 (3%)
I believe people with epilepsy should NOT do many recreational activities that I am able to do*	1,338 (35%)	1,594 (41%)	543 (14%)	278 (7%)	122 (3%)
I believe people with epilepsy can do anything I can do	65 (2%)	254 (7%)	468 (12%)	2,031 (52%)	1,057 (27%)
I believe people with epilepsy can work 40 hours per week	78 (2%)	140 (4%)	597 (15%)	1,925 (50%)	1,135 (29%)
I would NOT go out again with a person I just started dating if I found out he or she has epilepsy*	1,528 (39%)	1,318 (34%)	736 (19%)	197 (5%)	96 (2%)
I believe people with epilepsy are able to cope with everyday life as well as other people	60 (2%)	157 (4%)	492 (13%)	1,963 (51%)	1,203 (31%)
I believe parents should expect less of their child if the child has epilepsy*	1,492 (39%)	1,560 (40%)	466 (12%)	268 (7%)	89 (2%)
I believe people with epilepsy have a mental health problem*	1,820 (47%)	1,239 (32%)	483 (12%)	235 (6%)	98 (3%)
I would NOT want to work with someone with epilepsy*	1,675 (43%)	1,462 (38%)	477 (12%)	173 (4%)	88 (2%)
I believe people with epilepsy should NOT have biological children*	1,716 (44%)	1,395 (36%)	500 (13%)	174 (4%)	90 (2%)
I believe children with epilepsy can perform well in mainstream schools (not a special needs school)	30 (1%)	103 (3%)	432 (11%)	1,855 (48%)	1,455 (38%)
I believe people with epilepsy are unreliable*	1,722 (44%)	1,507 (39%)	389 (10%)	169 (4%)	88 (2%)
I believe people with epilepsy should tell their employers that they have epilepsy	49 (1%)	96 (2%)	403 (10%)	1,814 (47%)	1,513 (39%)
I believe people with epilepsy can lead normal lives	30 (1%)	99 (3%)	358 (9%)	1,836 (47%)	1,552 (40%)
I believe people with epilepsy should hide their condition from others except for close family and friends*	1,982 (51%)	1,233 (32%)	394 (10%)	195 (5%)	71 (2%)
I believe having epilepsy is nothing to be embarrassed about	92 (2%)	93 (2%)	255 (7%)	1,641 (42%)	1,794 (46%)

I believe people with epilepsy can be as successful at work as others	37 (1%)	62 (2%)	248 (6%)	1,884 (49%)	1,644 (42%)
I would be embarrassed if someone in my family had epilepsy*	2,172 (56%)	1,192 (31%)	244 (6%)	171 (4%)	96 (2%)
I believe having epilepsy is nothing to be ashamed about	54 (1%)	98 (3%)	222 (6%)	1,533 (40%)	1,968 (51%)
I believe people with epilepsy are NOT as smart as other people who do not have epilepsy*	2,342 (60%)	1,030 (27%)	264 (7%)	150 (4%)	89 (2%)
I believe people with epilepsy should NOT marry*	2,405 (62%)	993 (26%)	253 (7%)	152 (4%)	72 (2%)
I would consider getting a divorce if my spouse were diagnosed with epilepsy*	2,518 (65%)	898 (23%)	262 (7%)	127 (3%)	70 (2%)
I believe people with epilepsy are possessed by a supernatural spirit*	2,872 (74%)	542 (14%)	225 (6%)	174 (4%)	62 (2%)

*During analysis. item coded in reverse to match direction of underlying construct