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Evaluating risk to people with epilepsy during the COVID-19 pandemic: Preliminary findings from the COV-E study


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ABSTRACT

The COVID-19 pandemic has caused global anguish unparalleled in recent times. As cases rise, increased pressure on health services, combined with severe disruption to people’s everyday lives, can adversely affect individuals living with chronic illnesses, including people with epilepsy. Stressors related to disruption to healthcare, finances, mental well-being, relationships, schooling, physical activity, and increased isolation could increase seizures and impair epilepsy self-management.

We aim to understand the impact that COVID-19 has had on the health and well-being of people with epilepsy focusing on exposure to increased risk of seizures, associated comorbidity, and mortality. We designed two online surveys with one addressing people with epilepsy directly and the second for carers to report on behalf of a person with epilepsy.

The survey is ongoing and has yielded 463 UK-based responses by the end of September 2020. Forty percent of respondents reported health changes during the pandemic (n=185). Respondents cited a change in seizures (19%, n=88), mental health difficulties (34%, n=161), and sleep disruption (26%, n=121) as the main reasons. Thirteen percent found it difficult to take medication on time. A third of respondents reported delay and difficulty in accessing health services (n=154), with 8% having had an appointment canceled (n=39). Only a small proportion reported having had discussions about epilepsy-related risks, such as safety precautions (16%, n=74); mental health (29%, n=134); sleep (30%, n=140); and Sudden Unexpected Death in Epilepsy (SUDEP; 15%, n=69) in the previous 12 months.

Keywords:
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Chronic illness
Mental health
Seizures
SUDEP

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1. Introduction

The COVID-19 pandemic has created particular challenges for people with chronic illness, who may have greater susceptibility to adverse consequences beyond direct risks from infection [1–3]. People with epilepsy are generally not considered to be more likely to contract COVID-19, or necessarily develop a more significant illness in case of infection [4,5]. The indirect consequences of the pandemic could, however, impair their health and well-being in numerous ways [2]. Stressors, including economic challenges, anxiety, isolation, decreased physical activity, as well as interruptions to schooling, work, and family life, may increase seizures [6,7]. Disruption to self-management practices, such as adherence to anti-seizure medications (ASMs), and a lack of access to healthcare may also play a role [8–10]. Together, these factors could increase seizures, worsen associated psychiatric comorbidities such as anxiety and depression, and contribute to an increased risk of premature mortality [11].

Medical services have reconfigured rapidly during the pandemic. In the United Kingdom (UK), many epilepsy nurses and neurologists with an interest in epilepsy were redeployed into general medical services, resulting in the temporary closure of epilepsy clinics [12]. Video-EEG monitoring and other elective admissions were suspended. Access to neuropsychology was significantly limited, and epilepsy surgery was postponed [13,14].

The pandemic has also sharply accentuated inequality on multiple levels [15,16]. For example, health service disruptions are likely to have a greater impact on vulnerable populations, including Black, Asian and Minority Ethnic (BAME) groups, the elderly, and people from lower socioeconomic strata [17–19]. All of these factors are predicted to adversely affect those with epilepsy, particularly those at high risk [17–21].

To examine this, the COVID-19 and Epilepsy (COV-E) Study Group, led by SUDEP Action & University of Oxford, launched online surveys to assess the experiences of people with epilepsy through the first wave of the pandemic. We aimed to understand the impact of COVID-19 on people’s health and well-being. Specifically, we evaluated whether people’s exposure to risk had changed, including an increased risk of seizures, worsening morbidity, and their corresponding risk of premature mortality. While the surveys achieved a global reach in multiple languages, here we present the UK data. These data are derived from people with epilepsy and their caregivers, submitted during the first four months of the COVID-19 pandemic when the rate of new COVID-19 infections had a considerable impact on epilepsy services [3,24]. We explore various themes associated with COVID-19 and epilepsy, including respondents’ demographics, health status, and their interactions with health services. We also assessed knowledge and exposure to risk, including factors associated with premature mortality in people with epilepsy [20–23].

2. Methods

2.1. Study design

In April 2020, we designed separate surveys for people with epilepsy and primary caregivers [https://sudep.org/epilepsy-risks-and-covid-19-survey-people-epilepsy; Supplementary material] hosted on Jisc online survey software (https://www.onlinesurveys.ac.uk/). Pilot testing was performed on volunteer people with epilepsy before the online launch. Initial surveys were in English and these have since been translated into 11 languages. The surveys focus on quantitative data collection while also presenting participants with the opportunity to provide qualitative data through free-text answers. The study was approved by the University of Oxford Ethics Committee (Reference: R69353/RE001).

Inclusion criteria stipulated that people completing the survey had to be over the age of 18 years and be a person with epilepsy or someone who cares for a child or adult with epilepsy. Participants were provided with study information and completed an online consent form. All data were collected online, and data entry was anonymous. Healthcare workers were also surveyed, data from which will be presented in future publications.

2.2. Measurements

2.2.1. Demographics

People with epilepsy provided background information including their age, gender, ethnic background, and postcode information [25,26].

2.2.2. Epilepsy type/health background

Respondents were asked about epilepsy type, seizure type(s), and frequency; nocturnal events; medications; primary epilepsy care provider; the number of specialist visits in the past year; unplanned/emergency hospital admissions due to epilepsy; associated injuries; and comorbidities. People were asked if they had contracted COVID-19 or had to self-isolate owing to possible exposure.

2.2.3. Risk factors for epilepsy morbidity and mortality

We enquired about changes in behavior, habits, and circumstances during the pandemic, potentially associated with increased risk. Specific questions were included that related to mental health status, alcohol and drug consumption, sleeping patterns, and changes to seizures [21–23,27]. We asked about individual living circumstances and whether respondents lived alone or with someone who could provide first aid [28].

We explored the discussion of risk between the participant and their clinician in the previous 12 months. We asked, where relevant, whether the following had been discussed: ASM side effects, rescue medication, alcohol, contraception, driving, life changes, employment, mental health, pregnancy, recreational drugs, safety aids, first aid, sleep, stigma, and Sudden Unexpected Death in Epilepsy (SUDEP) [21–23,27].

2.2.4. Access to healthcare

We investigated if there had been an impact on healthcare access (e.g. experience in obtaining prescriptions, changes to arranged appointments, communication with clinicians), and user satisfaction. Participants were encouraged to provide free-text answers to contextualize their responses.
2.2.5. Caregiver survey

The caregiver survey mirrored that of the person with epilepsy survey to ascertain the impact of COVID-19 on the individual with epilepsy, through a caregiver’s perspective. Caregivers were asked about their relationship to the person with epilepsy (e.g. parent, remunerated caregivers). Questions allowed caregivers to disclose gaps in their knowledge of the person’s background, health status, or treatment plan. In such instances, respondents could select “unsure” or “don’t know” options and elaborate where relevant.

2.3. Dissemination

Survey dissemination was led by SUDEP Action and was also shared on social media and promoted by multiple epilepsy support organizations, including, but not limited to, the BAND Foundation, Citizens United for Research in Epilepsy (CURE), Epilepsy Action, Epilepsy Foundation America, Epilepsy Research UK, Epilepsy Society, Epilepsy Sparks, the International Bureau for Epilepsy (IBE) and the International League Against Epilepsy (ILAE) (see acknowledgments).

2.4. Data analysis

Data collection is ongoing. We collated data in the UK from April to September 2020 and categorized these into: i) demographics; ii) reported health outcomes; iii) awareness of risk; and iv) access to epilepsy care.

Each survey was first analyzed in isolation, utilizing descriptive statistics and cross-tabulation of data using tools provided by Jisc. Data were then exported into Microsoft Excel, enabling cross-comparison of individual and caregiver surveys. We compared qualitative and quantitative data [29,30] to clarify responses, particularly for ethnicity, comorbidities, medication, and hospital admissions. Postcode data were processed using mapping tools provided by Datawrapper to produce choropleth maps of the geographic distribution of responders.

We further analyzed data from groups at potentially heightened vulnerability, including those who identify as members of BAME communities [17–19], people who are pregnant, people over the age of 60, and those who experience other comorbid conditions that make them directly vulnerable to COVID-19 [2,15]. Data analyses used descriptive statistics, case reports, and qualitative coding of free-text data [29,31]. Statistically significant conclusions were not drawn where the sample size was n ≤ 50.

3. Results

We received 463 responses, 68% from people with epilepsy (n = 316), and 32% from caregivers on behalf of a person with epilepsy (n = 147).

3.1. Population demographics

3.1.1. Geographical distribution

Data were collected from 106 of the 122 UK postcode areas with greater representation from the South East, the Midlands, and the East of England. Urban centers included Birmingham (n = 15), Liverpool (n = 11) and London (n = 23) (Fig. 1). Ninety percent of respondents live in England and Wales (n = 416), 6% in Scotland (n = 30), and 3% in Northern Ireland (n = 15). Two respondents did not provide valid postcodes.

3.1.2. Gender

The people with epilepsy survey yielded a greater response rate from females (78%, n = 246). Twenty percent identified as male (n = 64) and <1% as nonbinary (n = 2). Two percent did not disclose their gender (n = 4).

In the caregiver survey, 49% identified as female (n = 72) and 50% as male (n = 74). One respondent did not disclose their gender.

3.1.3. Age

People under 18 were highly represented in the caregiver survey, accounting for 61% of responses (Table 1). Across both surveys, response rates were higher in younger age groups and decreased with age (Fig. 2).

3.1.4. Ethnicity

In total, 2% of respondents identified as belonging to an ethnic minority group (n = 8). An additional 5% of respondents were unsure or did not disclose the relevant information (n = 24). Seventeen responses were omitted from analysis as they identified themselves as belonging to a group that is not regarded as a minority ethnic identity in the UK [26,32,33].

3.1.5. Comorbidities

Respondents reported a range of other health conditions, which could make them directly or indirectly vulnerable during the pandemic [2,34–37]. A small number of people reported conditions associated with COVID-19 complications, including diabetes (2%, n = 7), heart problems (5%, n = 23) or a respiratory condition (10%, n = 44). Other prevalent health conditions included intellectual disabilities (13%, n = 62), memory difficulties (31%, n = 143), and mental health difficulties (27%, n = 123).

3.2. Exposure to risk during the COVID-19 pandemic

3.2.1. Health and well-being

Forty percent of respondents reported a change in health or in the health of the person they care for (n = 185). Overall, 19% reported changes in frequency, type, and length of seizures (n = 88), although it was not possible to always determine if this was a positive or a negative change. Qualitative data analysis did, though, demonstrate a worsening in seizure profile for some respondents. 26% recorded disrupted sleep patterns (n = 121); and 34% increased mental strain, stress, worry, anxiety, or depression (n = 161). In responses from people over the age of 18 years, 4% reported increased alcohol consumption (n = 14), while 2% reported increased recreational drug consumption (n = 6).

Of those who take ASMs (total n = 454), 13% reported greater difficulties in taking them on time (n = 57). Changes to routine (8%, n = 38), difficulty in acquiring prescriptions (3%, n = 15), and stress (7%, n = 32) were cited as the predominant causes for impaired adherence (Fig. 3). An analysis of qualitative data provides additional insights into why these changes may have occurred. Six caregivers reported that changes to schooling and lifestyle contributed to declining mental and physical health in the person for whom they care. Notably, two caregivers reported improved health outcomes as the individual with epilepsy had fewer infections and benefited from spending more time at home. Twenty-five pieces of anecdotal data from the individual survey identified factors contributing to their declining health (Table 2). Four respondents in the people with epilepsy survey reported improved mental health owing to a reduced need to travel or commute and an overall healthier and more relaxed lifestyle.

3.2.2. Access to healthcare

3.2.2.1. Prescriptions. Of people taking ASMs (total n = 454), 4% reported delays in prescription deliveries (n = 16); 3% reported problems in ordering prescriptions (n = 14); and 3% difficulties in collecting their prescription (n = 14). Qualitative data highlight...
additional problems, including difficulty acquiring a repeat prescription and contacting primary healthcare physicians (General Practitioner, GP) or pharmacists. Five respondents, whose treatment plan had changed before or during the lockdown, reported that their new treatment plan had not been processed or adequately supported by their healthcare providers.

3.2.2.2. First aid. A quarter of people with epilepsy were living alone during lockdown (n = 112). In the sample of people who live alone, only 23% (n = 26/112) had discussed safety aids (seizure alarms, diaries) with an epilepsy specialist in the previous 12 months (n = 26/112), and 15% had discussed safety precautions and first aid (n = 17/112).

Additionally, of the 346 participants who reported living with another household member, 15% did not live with anyone who could provide first aid (n = 53/346).

3.2.2.3. Emergency care. Thirty-eight percent of respondents across both surveys reported acquiring injuries or needing emergency care due to epilepsy or related injuries during the previous

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**Fig. 1.** Geographical location of survey responses by postal code. Survey responses were received from almost all locations in the United Kingdom with a preponderance of responses from the south of England and the Midlands. Created using datawrapper.de.
12 months (total n = 176). Qualitative data show at least 11 people have had more than one emergency admission in the past 12 months, with an additional eight respondents requiring hospitalization of a day or more. Forty-one percent of the 176 people who have been injured or required emergency care in the past 12 months lived alone during the lockdown (n = 71).

3.2.2.4. Epilepsy services. A third of respondents reported difficulty accessing epilepsy services (n = 154), describing particular difficulty in reaching neurology, GP, and pharmacy services (Fig. 4). Qualitative data indicate that these difficulties stem from the cancellation or postponement of services, or changes to services. For example, people reported a change in the healthcare facility or clinician, or services being provided through virtual technologies.

Quantitative data from both surveys revealed that 48% of all respondents had a planned medical appointment changed (n = 221) and 8% had appointments canceled (n = 39). Twenty-two percent of respondents had not had any communication about prescheduled appointments (n = 100).

In analysis of the results from participants who had seen changes to their appointments (total n = 221), 25% were to take place later in the year (n = 56/221); 48% by telephone as originally scheduled (n = 107/221); and 17% by telephone at a later time (n = 38/221). Only 9% of participants were offered video consultations (n = 19/221). Six percent of video consultations were to take place at the originally scheduled time (n = 13/221), while 3% had video appointments moved to a different time (n = 6/221).

In total, across both surveys, 37% of individuals reported being satisfied with the current status of their next consultation (n = 173/463). Thirty-four percent reported being unsatisfied or unsure (n = 147/463) and 29% did not comment (n = 143/463). Qualitative data indicate that several respondents consider in-person appointments to be preferable, particularly for those with additional disabilities, including autism and hearing loss, who face additional challenges in adapting to health service delivery through telecommunication.

3.2.3. Risk awareness

Analysis of both surveys showed that few respondents reported communication on enduring risks, such as safety precautions (for example, carers being aware of first aid, not swimming in open water; 16%, n = 74); mental health (29%, n = 134); sleep (30%, n = 140); and SUDEP (15%, n = 69). 20% of respondents had not engaged in patient-clinician communication on any listed risk factors in the past 12 months (n = 92; Fig. 5). Of the 92 respondents who reported complete nonengagement on risk, 39% also reported seeing a healthcare professional less than once per year (n = 44/92). Risk awareness data varied between individual and caregivers surveys, likely owing to the high proportion of children in the caregiver survey, where risk factors and issues may differ from adults. Some risk factors included in the people with epilepsy survey may not be relevant for all individuals [27].

3.3. Special groups

3.3.1. Black, Asian and Minority Ethnic (BAME)

Four out of eight BAME participants reported changes to their health, citing changes in seizure activity (n = 1), sleep disruption (n = 2), and mental strain (n = 2) as reasons for this (Table 3). One respondent reported feeling healthier and more relaxed. Four respondents reported difficulty in accessing health services, including neurology (n = 3), general hospital services (n = 2), general practice (n = 1) and out-of-hours services (n = 1), pharmacy services (n = 1), NHS helpline services (n = 1), learning disability services (n = 1), occupational therapy (n = 1), physical therapy (n = 1) and speech therapy (n = 1). Multiple responses to these questions were permitted to cover all difficulties respondents had experienced.

Seven respondents demonstrated regular engagement with epilepsy specialists, with just one participant seeing an epilepsy specialist less than annually. No respondents had discussed alcohol consumption, drug use, or stigma. Two respondents reported that there had been no discussion on risk factors in the previous 12 months.
3.3.2. People with epilepsy aged over 60 years

People over the age of 60 years reported the highest prevalence of diabetes, heart conditions, hypertension, and memory difficulties. Frequency of mental health difficulties and respiratory conditions was lower than the overall average (Table 4). Fewer individuals in this older cohort reported changes to their overall health and well-being during the COVID-19 pandemic compared with the wider dataset. A smaller proportion reported difficulty in taking ASMs on time and accessing epilepsy care (Table 4). There were, however, multiple other areas in which the risks to older people with epilepsy could be modified. In this demographic, for example, 35% of individuals accessed a health specialist less than once a year ($n = 12$), 35% lived alone ($n = 12$), and 26% were living with someone who would not be able to provide first aid ($n = 9$).

### Table 2

<table>
<thead>
<tr>
<th>Perceived causes for changes in health</th>
<th>Number of responses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Difficulties in acquiring prescriptions</td>
<td>1</td>
</tr>
<tr>
<td>Difficulty in receiving healthcare</td>
<td>7</td>
</tr>
<tr>
<td>Adjusting medication during lockdown</td>
<td>1</td>
</tr>
<tr>
<td>Stress and anxiety related to COVID-19</td>
<td>7</td>
</tr>
<tr>
<td>Changes to weight or fitness</td>
<td>3</td>
</tr>
<tr>
<td>Job instability</td>
<td>1</td>
</tr>
</tbody>
</table>

3.3.3. Clinically vulnerable to COVID-19

When limiting analysis to individuals who have diabetes, hypertension, heart conditions, or respiratory conditions (total $n = 97$), a higher proportion of respondents reported epilepsy-related changes to their health ($56\%, n = 48/97$). Overall, 23%

![Fig. 3. Cited reasons for difficulties in ASM adherence and concordance as reported in the caregivers and people with epilepsy surveys, respectively. A larger proportion of respondents in the people with epilepsy surveys reported difficulties in adhering to ASM medication compared to responses from the caregiver survey (total $n = 57$). Changes in everyday routine was cited as the most common cause of nonadherence in both surveys, followed by stress or worry resulting in forgetfulness.](image1)

![Fig. 4. Number of people with epilepsy and caregivers who reported difficulties in accessing healthcare stratified by service type. A total of 154 respondents reported difficulties accessing healthcare during phase 1 of the COVID-19 pandemic. Respondents were able to answer stating that they had difficulty with more than one aspect of health care and so raw numbers of respondents are presented.](image2)
noticed an increase in seizures \((n = 22/97)\), 34\% experienced disrupted sleep \((n = 33/97)\), and 43\% reported increased mental strain \((n = 42/97)\). Thirty-seven percent reported difficulty in accessing epilepsy services \((n = 36/97)\).

### 3.3.4. Pregnancy

Five respondents, all aged between 30 and 39 years, reported being pregnant. All were living with other household members who could provide first aid. While the sample size is small, this cohort is possibly reflective of the broader data regarding health changes and accessing epilepsy services (Table 5). This group showed low engagement in epilepsy services and poorer risk communication [27], despite being a high-risk group for COVID-19 and epilepsy-related mortality.

### 4. Discussion

The COVID-19 pandemic is having an unparalleled global impact with far-reaching consequences on health and social care. While significant emphasis has been placed on the direct health impacts of COVID-19, our results suggest that people with epilepsy have also experienced a decline in health and well-being. Survey respondents reported increased seizure frequency and seizure intensity, decreased adherence to ASMs, impaired mental health, and, in some instances, increased substance or alcohol use. These factors can lead to increased seizure activity in people with epilepsy and heighten the potential for premature death from SUDEP, unintentional injuries, or suicide [11,27,28]. Crucially, these changes have occurred in the context of restricted healthcare provision, reducing capacity to address concerns [39]. The vulnerability appears to be compounded by the deficiency in risk awareness and insufficient engagement with healthcare services before and during the pandemic.

Rates of seizure-related deaths and SUDEP correlate with increased seizure frequency and insufficient engagement with clinicians [20–23,28,40–41]. Without appropriate medical attention, indicators that point to an increased likelihood of premature mortality may not be identified, restricting the ability to intervene [28].
The risk of mortality may increase owing to poor communication about risk. This can disempower people and inhibit their ability to make informed decisions on, for example, ASM adherence, alcohol and nonprescription drug consumption, family planning, driving, and sleep habits [28].

Epilepsy-associated risk, including premature mortality, is greater in lower socioeconomic groups, who make up the largest proportion of epilepsy cases in the UK [42]. These groups are already predisposed to higher levels of comorbidities and experience more years of life lost to disability [11,43–44]. This can hinder an individual’s stability and financial resilience, contributing to a decrease in earning potential and employment opportunities when unwell [11,42–45]. As services reconstitute, clinicians must engage in comprehensive communication to ascertain the specific needs of individuals, alongside an appraisal of their physical health, psychosocial well-being, and the interrelation between these components. To achieve this, a multimodal approach that includes social and economic interventions is essential.

Enabling good communication and accessibility are crucial in optimizing health care for people with chronic illness. Our study suggests that increased usage of remote consultations during the pandemic will benefit some, but may be a barrier for others (section 3.2.2.4). Ascertaining individual needs may streamline service provision, ensuring that those who benefit from face-to-face consultations can be prioritized in line with current capacities. Greater consensus on the role of technologies in medical care may facilitate better communication between healthcare providers. This could help avoid delays in referrals, treatments, or changes to medication regimes as well as alleviating confusion and uncertainty for people regarding their entitlements to healthcare.

To ensure that any potential gaps in care are mitigated, best-practice resources, such as the SUDEP & Seizure Safety Checklist, can be useful. These are able to provide personalized advice on reducing risks, including those that are of increasing relevance during the current circumstances [47]. Self-monitoring between appointments may also be beneficial for individuals in tracking their own experiences and behaviors, particularly when healthcare delivery is less straightforward. Resources such as EpSMon epilepsy self-monitoring app [28,46] and other technologies may empower individuals day-to-day and support them in monitoring and reflecting on their health while simultaneously providing resources to educate on risk reduction strategies [47–50].

Efforts should be made to ensure equitable health interventions that recognize the increased needs of special groups, a facet which has been highlighted previously in work examining the effect of COVID and lockdowns on children with epileptic encephalopathies [51,52]. COVID-19 disproportionately affects people of BAME heritage and the elderly [15–16,34–35], while the incidence of epilepsy is highest in older people and those from lower socioeconomic groups [17–18,42]. Despite extensive social media messaging, we only received small numbers of respondents from BAME and older communities. Indirectly, this highlights the need for a tailored approach to allow information to reach people from these groups [53–55]. Clinicians must be vigilant to the specific needs of increasingly diverse populations.

5. Limitations

Our study has limitations, predominantly relating to the need for the surveys to be online and owing to data anonymization.
Participant self-selection could introduce bias, meaning our findings cannot be extrapolated to the entire population with epilepsy. Variable reporting was partially identified through analysis, but some inconsistencies in reporting may remain. We cannot quantify health changes due to the absence of baseline data before the pandemic and the possibility of recall bias [56]. Also certain questions will need refining in future iterations as, for example, people could report a change in their seizures without necessarily specifying in which direction (worsening or improvement) that change was. Qualitative data enable us to conclude that some individuals have noticed deterioration in seizure control. With the second version of the surveys, more quantitative data on this aspect will be sought.

Socioeconomically vulnerable groups are under-represented (including those who are homeless, the elderly, ethnic minorities, LGBTQ+, and those with additional disabilities). These disparities can be partly attributed to the dependence on utilizing a text survey on an online platform which may not be accessible for all prospective participants [57]. Those marginalized by health and social care providers may be more reluctant to engage in research [53,54]. We aim to overcome these barriers as the study progresses by providing the surveys in multiple languages and increasing our efforts in engaging with outreach groups. Anonymous data collection using digital means may also facilitate greater inclusion of those who feel socially stigmatized and are more active in online spaces [55,58].

6. Conclusion

COVID-19 has caused a paradigm shift in healthcare provision. As the UK currently witnesses its second wave of cases, it is essential that those involved in healthcare delivery can ensure optimal care for people with epilepsy. A multifaceted, person-centered approach with appropriate use of currently available technologies can help bridge the emerging gap in healthcare provision, while mitigating the rising risks associated with epilepsy-related mortality. These interventions must also account for varying needs and levels of risk within populations. The results of the COV-E study carry relevance that extends beyond epilepsy care, and highlight themes that apply to many other long-term chronic illnesses [1,2,36]. Moving forward, we hope our research will facilitate further awareness and tailored targeting of strategies to help improve the care of people with epilepsy during and after the pandemic. We also hope to inspire a broader body of research on COVID-19 and its impact on people with chronic illnesses.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1101/jy.2020.107658.

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