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Application of recent international epidemiological guidelines to a prospective study of the incidence of first seizures, newly-diagnosed epilepsy and seizure mimics in a defined geographic region in Ireland

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Short title: Methodology of a prospective incidence study on epilepsy, seizures and mimics

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1. Abstract

Studies adherent to international guidelines and epilepsy classification are needed to accurately record the incidence of isolated seizures, epilepsy and seizure-mimics within a population. Because the diagnosis of epilepsy is largely made through clinical assessment by experienced physicians, seizures and epilepsy are susceptible to misdiagnosis. Previous epidemiological studies in epilepsy have not captured ‘seizure mimics’. We therefore sought to quantify the incidence of isolated seizures, epilepsy and seizure-mimics using the International League Against Epilepsy (ILAE) classification system. In this study multiple overlapping methods of case ascertainment were applied to a defined geographic region from 1st January 2017 to 31st March 2017 to identify all patients presenting with first seizures (provoked and unprovoked), new diagnoses of epilepsy and seizure mimics. Over a three month period, from a population of 542,869 adults and children, 442 potential presentations were identified, and 283 met the inclusion criteria. Radiology databases were the source of the largest number of individual cases (n=153, 54%), while electroencephalogram (E.E.G.) databases were the source of the highest number of unique-to-source cases (those not identified elsewhere, n=60, 21%). No single case was picked up in every method of ascertainment. Among the 283 included presentations, 38 (13%) were classed as first provoked seizures, 27 (10%) as first unprovoked seizures, 95 (34%) as new diagnosis of epilepsy and 113 (40%) as seizure mimics. Ten (3%) presentations were indeterminate. We present and apply a rigorous study protocol for investigation of the incidence of first seizures, new diagnosis of epilepsy and seizure mimics in a geographically defined region which is adherent to recently published international guidelines for epidemiological studies and epilepsy classification. We highlight the challenges in making a diagnosis of new-onset epilepsy in patients presenting with a first seizure using the current ILAE definition of epilepsy, when epilepsy can be diagnosed in situations where the treating physician anticipates the risk of further seizures exceeds 60%.
2. Introduction

Epilepsy is a disorder of the brain characterized by an enduring predisposition to generate epileptic seizures and by the neurobiologic, cognitive, psychological, and social consequences of this condition [1]. Operationally, epilepsy can be diagnosed in 3 circumstances: (i) at least 2 unprovoked (or reflex) seizures occurring more than 24 hours apart, (ii) one unprovoked (or reflex) seizure and a probability of further seizures similar to the general recurrence risk (at least 60%) after 2 unprovoked seizures, occurring over the next 10 years, and (iii) diagnosis of an epilepsy syndrome [2]. The second situation is relatively common but difficult to apply in real life as judging that an individual patient has a 60% risk of further seizures is challenging. Epilepsy poses a substantial economic burden for health care systems, individuals and their families through direct costs of treatment and indirect costs such as loss of productivity and employment [3]. Epidemiological studies are necessary to define the full public health burden of epilepsy within a population, to provide information needed for early detection and treatment, and to set public health and health care priorities [4]. Assessment of patients presenting with first seizures is critical to differentiating epileptic seizures from other conditions that resemble epileptic seizures (‘seizure mimics’). Although first seizures and epilepsy are susceptible to misdiagnosis, to our knowledge there are no published epidemiological studies on seizure mimics.

There are significant disparities in reported prevalence and incidence of epilepsy worldwide. While some variation may be related to factors such as socioeconomic class, access to healthcare, stigmatization and exposure to environmental risk factors [5], a significant contributor to the variation in reported incidence and prevalence may be due to heterogeneous methodologies used across studies. A recent meta-analysis of 48 incidence studies noted significant heterogeneity between studies and estimated the pooled annual cumulative incidence of epilepsy as 67.77 per 100,000 persons (95% confidence interval [CI] 56.6 to 81.0) with one outlier report of 189.96 per 100,000 [6]. Methods of case ascertainment in published studies range from door-to-door population surveys, administrative database searches and surveys of neurology referral centers [7-9], thus providing variations in inclusion criteria and accuracy of case ascertainment. Many studies employ retrospective case ascertainment and lack of detailed diagnostic information often prohibits accurate classification of seizure and epilepsy type.

In 2011, in an effort to promote consistency in definitions and methods used in epidemiological studies and to facilitate comparisons between populations, the International League Against Epilepsy (ILAE) proposed standards for epidemiologic studies and surveillance of epilepsy [4]. These guidelines emphasize the importance of using multiple overlapping methods of case ascertainment to maximize the sensitivity of case ascertainment. Furthermore, Standards of Reporting of Neurological Disorders (STROND) guidelines for reporting of incidence and prevalence studies in neuroepidemiology have been developed to facilitate better reporting of published data and allow comparisons between studies [10]. When applied to future studies, these guidelines will enhance population-based
epidemiological studies and encourage the collection of data useful for the promotion of public health. Finally, the ILAE has recently commissioned updated classifications systems for both seizures and epilepsy types [11,12]. These classification systems are user-friendly and allow classification of seizure and epilepsy type by taking into account results of E.E.G. and imaging investigations. To our knowledge, this is the first epidemiological study to incorporate these classifications.

In Ireland, it has been estimated that 10 per 1,000 persons aged 18 years and older have a self-reported lifetime prevalence of epilepsy [13]. This study is the first to investigate the incidence of new onset seizures, epilepsy and seizure mimics in Ireland and, to our knowledge, the first to present epidemiological data on seizure mimics. We present our study protocol, adherent to ILAE and STROND guidelines, and findings for the first three months of data collection.

3. Materials and Methods

This study was carried out in the geographically defined area of Cork city and county over the course of the calendar year 2017, with an estimated total population of 542,868 persons based on a census in 2016. Herein we present our study protocol and, to demonstrate its application, the results of case ascertainment for the first three months of data collection, 1st January 2017 to 31st March 2017.

All acute medical hospitals were included with the exception of one solely private hospital that does not employ an on-site consultant neurologist. Acute seizures presenting to the medical assessment unit of that hospital are transferred to the tertiary university hospital in the city. The remaining 7 acute hospitals included in this study were as follows: 2 tertiary referral city center university hospitals, 2 regional secondary level hospitals, 1 solely private city center hospital which accepts acute referrals and employs 2 consultant neurologists, 2 city center hospitals with facilities for rehabilitation of geriatric patients who have recently been acutely admitted elsewhere e.g. ortho-geriatrics rehabilitation. This study also included community sources of case ascertainment as follows: all general practitioners (G.P.s) in Cork city and county who are registered with the Irish Medical Directory, all nursing homes and residential care centers registered with the Health Information and Quality Authority, and Epilepsy Ireland, the community based patient information and advocacy group.

3.1 Inclusion and exclusion criteria

We included all patients who had a suspected first seizure or new diagnosis of epilepsy from 1st January to 31st December 2017 whose registered address as per the hospital-based demographic information was within Cork city or county (defined area). The working diagnosis of ‘seizure’ had to be explicitly documented in medical correspondence during the patient’s assessment typically by a G.P. in the community, an emergency department triage nurse or physician, or senior hospital-based physician. Any patient, normally resident in the defined area, who had a
suspected seizure during 2017 but presented to a hospital outside the area was included. These patients were identified through survey of G.P.s and neurologists in Cork city and county, where their ongoing care and follow up was based. All patients with an address outside of the defined area were excluded.

We gathered information on all patients whose first presentation to medical services with a query of seizure occurred during 2017. In some cases, the date of the first clinical event was prior to the 2017 calendar year, but medical attention was first sought in 2017, for example, when a patient has a history of recurrent stereotyped events, but the possibility of these events being seizures had not yet been explicitly stated, or medical attention was not sought. Similarly, we continued to monitor a number of case ascertainment sources [Rapid Access Seizure Clinic (R.A.S.C.) referrals, electroencephalogram (E.E.G.) and radiology databases] up to 31st March 2018 in order to complete the classification of patients who first presented in late 2017. By restricting our count to first presentation during 2017, we will avoid over- or under-ascertainment in the 2017 calendar year, and the number of first presentations classified will estimate the true incidence of each clinical subcategory.

As per the ILAE epidemiologic guidelines [4], febrile seizures in children (3 months to 6 years old) and neonatal seizures (<28 days old) were excluded from this study as they are felt to be separate from epilepsy per se. With regard to acute provoked seizures, we noted the incidence of their occurrence as a specific research question, but they were separated from epilepsy in the final analysis.

### 3.2 ILAE Definitions

i) An epileptic seizure is a transient occurrence of signs and/or symptoms due to abnormal excessive or synchronous neuronal activity in the brain [1].

ii) Epilepsy is a disease of the brain defined by any of the following conditions: i) at least two unprovoked (or reflex) seizures occurring >24 hours apart ii) one unprovoked (or reflex) seizure and a probability of further seizures similar to the general recurrence risk (at least 60%) after two unprovoked seizures, occurring over the next 10 years, or iii) diagnosis of an epilepsy syndrome [2].

iii) Acute provoked seizures are epileptic seizures which occur in close temporal association with an acute systemic, metabolic or toxic insult or in association with an acute central nervous system insult [4]. The interval between the insult and the seizure may vary according to the underlying clinical condition (see Table 1). With regard to alcohol withdrawal seizures, the seizure must have occurred within 7-48 hours of the last drink to be classified as an alcohol withdrawal seizure [14]. Alternatively, any patient with a history of alcohol abuse and acute provoked seizures from alcohol, who developed their first seizure independent from alcohol within the study period [and was therefore commenced on anti-epileptic drug (A.E.D)] was counted as a ‘new diagnosis of epilepsy’. Seizures in the setting of acute alcohol intoxication due to extremely high quantities consumed and seizures in association with withdrawal of benzodiazepines were classified as acute provoked seizures [14].
iv) For all patients with a new diagnosis of epilepsy, the 2017 ILAE position papers on classification of seizures and epilepsy were applied. Seizures were therefore classified as focal onset, generalised onset or unknown onset and further descriptors were applied where possible [11]. Epilepsy was classified as focal, generalised, focal and generalised or unknown depending on available data [12].

3.3 Case ascertainment

We broadly divided our case ascertainment methods into ‘hot pursuits’, where information was gathered prospectively on a daily basis from active inpatient databases and services, and ‘cold pursuits’ where data were periodically reviewed retrospectively and cases were cross-referenced to the study database to prevent duplication. For an overview of the process of case ascertainment and case analysis, please see Figure 2.

3.3.1 ‘Hot pursuits’

i) Emergency departments. Triage recording systems vary in each of the hospitals and therefore it was necessary to screen emergency departments in different ways depending on the resources available. In the main university hospital of the city, electronic triage records allowed us to screen all patients presenting acutely to the emergency department and to identify all possible seizures. The electronic triage system was checked on a daily basis. Search terms including ‘seizure’, ‘epilepsy’, ‘collapse’, ‘seizure-like events’ and ‘fits’ were screened for among the initial post-triage electronic entries. The remaining hospitals operate paper based triage systems and a point of contact in each hospital was established to collect Medical Record Number (MRNs) by hand of all patients presenting with a possible seizure.

ii) Radiology. The clinical indication for all CT and MRI brains performed in each of the included hospitals was reviewed systematically to capture all brain imaging requested for an indication of possible seizure. We included direct queries of ‘first seizure’ as well as more indirect queries such as ‘collapse with jerking’.

iv) Inpatient services. Inpatient services with a high likelihood of assessing patients with first seizures were identified. These included neurology, geriatrics, oncology and neurosurgery inpatient services. These services were specifically targeted as it was felt that they might assess and treat patients for a possible seizure without necessarily ordering specific neurophysiology or neuroimaging investigations. These teams were contacted on a fortnightly basis in order to prompt recall of recent cases.

v) Clinical Nurse Specialists (CNSs). CNSs from services felt to have a high likelihood of encountering patients with a first seizure were contacted on a monthly basis and asked to inform the study team of any new case. CNS in paediatrics, neurology and oncology participated in this study.

3.3.2 ‘Cold pursuits’
i) E.E.G. databases. There are three neurophysiology departments in the capture region, two public and one private. All paediatric and adult E.E.G. are performed at one of these three sites. The clinical indication for all E.E.G.s performed during the study period was reviewed and all possible first seizures and new diagnosis of epilepsy were included.

ii) R.A.S.C. One consultant epileptologist works in the public service in the capture region (author D.C). This consultant runs a R.A.S.C. once per week with an aim of rapidly reviewing G.P. and emergency department referrals for ‘query seizure’. All new referrals to this clinic were reviewed on a weekly basis.

iii) Survey of hospital consultants. Consultant physicians in neurology and geriatrics departments in the seven hospitals outlined above were contacted via postal survey every 8-12 weeks and asked to alert us if they were aware of any new case.

iv) hospitals outside of the capture region. We acknowledge the possibility that a small proportion of patients whose residence is toward the outer edge of the capture region may present acutely to a regional hospital outside of the county. There are three acute hospitals in the neighboring counties with four consultant neurologists working between them. Each neurologist was contacted via a postal survey every 8-12 weeks and asked to alert us if they were aware of any new case which met our inclusion criteria.

v) General practice. All G.P.s in the defined region were contacted via postal survey every 12 weeks and asked to alert us if s/he was aware of any new case.

vi) Nursing Home and Residential Care survey. Clinical Nurse Manager of all registered nursing homes and residential care services were contacted via postal survey every six months and asked to alert us if s/he was aware of any new case. A six-month interval was chosen for nursing homes as case turnover is much lower than in general practice and we felt that recall would therefore be longer.

vii) Epilepsy Ireland. Epilepsy Ireland is a non-profit, nationwide patient advocacy and advice organisation for people with a diagnosis of epilepsy (www.epilepsyireland.ie). In order to rigorously identify any patient who may be resident in the defined area but who was diagnosed outside of one of the included hospitals, we contacted the local branch of Epilepsy Ireland. They reviewed their new patient database to identify patients who were ordinarily resident in the defined area but were diagnosed elsewhere. An Epilepsy Ireland staff member then consented the patient for inclusion in the study and forwarded their details to the study team.

Of note, we deliberately did not choose pharmacy data on A.E.D. prescribing or hospital coding databases for case ascertainment because in Ireland these databases are not linked to other medical data and typically do not discriminate between new cases and established cases of epilepsy.

3.3.3 Identifying under ascertainment

As a method of internal control, to determine if we were missing potential cases, we devised a method of assessment of case under ascertainment. We identified that patients with a known diagnosis of primary or secondary brain tumors are at increased risk of seizures [15]. We prospectively collected the MRNs of patients
discussed at the multidisciplinary neuro-oncology meeting of one tertiary referral university hospital from January to March 2017. Six months later, at the end of September 2017, we searched the medical records of these patients to identify if any had presented with a seizure. For any patient who had a seizure, we then crossed referenced them to our main database to see if they had already been captured through another method of ascertainment.

3.4 Case analysis

3.4.1 Demographic data and clinical data

The paper medical record of each patient identified through the above methods was obtained from the medical records department. The first author (E.M.) abstracted the demographic details, clinical admission details and results of all relevant investigations of each case through medical chart review. For any patient who was reviewed and diagnosed by a consultant neurologist, we adhered to the final diagnosis of the consultant neurologist. For all patients who were not reviewed by a consultant neurologist, the case was analysed by E.M and D.C. and a consensus decision was made. Any patient without a clear consensus was further analysed at panel review with a second consultant epileptologist (E.C.). D.C. and E.C. are trained neurologists who completed epilepsy fellowships and have more than 9 years clinical experience running epilepsy services.

3.4.2 Case classification

Following record review, patients were classified as one of the following five categories; first provoked seizure, first single unprovoked seizure, new diagnosis of epilepsy, seizure mimic or indeterminate event (see Figure 1). Individuals with a first single unprovoked seizure were classified as new diagnosis of epilepsy when it was estimated that there was a greater than 60% chance of a recurrent epileptic seizure in the next 10 years, as per the ILAE operational definition of epilepsy [2]. Otherwise, they remained classed as first unprovoked seizure. For definite new cases of epilepsy, the seizure and epilepsy type were subclassified as focal, generalized or unknown according to the ILAE 2017 classification system [3]. Cases were classified as indeterminate, following discussion panel review, if there was insufficient clinical information to accurately classify the case.

The 2011 ILAE Epidemiology Commission report on standards for epidemiologic studies and surveillance of epilepsy [4] provides a classification system for the level of evidence available to support a diagnosis of an epileptic seizure, or new diagnosis of epilepsy, based on the level of epidemiologic evidence available. Based on these standards, the following classification regarding level of evidence of both single epileptic seizures and new diagnosis of epilepsy were applied:

i) Definite: with primary documentation of (a) epileptic seizures, with evidence that these were unprovoked by any acute medical condition or transient brain disorder or (b) documentation of diagnosis by someone with appropriate specialised training in the recognition of epilepsy. Clear evidence of epileptic seizures was most often
based on documented collateral history or documented history by medical staff in the case on inpatient events. In the event of a single documented seizure, evidence of an approximate 60% risk of recurrence was determined based on the presence or absence of risk factors for recurrence such as epileptiform abnormality on E.E.G. or significant brain imaging abnormality [16-19].

ii) Probable: with other sources of information indicating the likelihood that criterion (a) or (b) above is met. For example, cases were defined as probable where there was a history strongly suggestive of an epileptic seizure, but a witnessed history was not recorded.

iii) Suspect: where primary or other sources of information suggest a possibility of epilepsy but neither (a) or (b) above is met. Possible epileptic seizures were defined as an event of which an epileptic seizure was one of a number of plausible differential diagnoses, but of which it was not possible to determine which was the most likely.

3.4.3 Accuracy of case information and classification

In order to ensure the data obtained from the medical records on each case was accurate and consistent, an internal audit of the data extraction was performed (by E.C.). Thirty-eight items were recorded from three randomly selected charts. Out of these 114 items (3x38), three items were discordant between E.C. and E.M. data extraction, resulting in a 97.4% concordance. None of the three discordant items altered the clinical diagnosis.

In order to ensure reliability of case classification, ten randomly selected cases were reviewed (by E.C). In the final classification, nine of the ten cases were in concordance. However, the internal audit highlighted the subjectivity of the ILAE practical diagnosis of epilepsy guidelines. In discussion, six of the 10 cases did not undoubtedly meet this criterion, and while all team neurologists agreed that these patients would be commenced on an A.E.D. due to risk of further seizures, whether this risk reached the 60% threshold was open to debate.

4. Results

The above protocol was applied to our study population from 1st January to 31st March 2018 in order to capture all patients who presented to medical services with a suspected first seizure or new diagnosis of epilepsy during the calendar year 2017. To demonstrate the protocol application, we present the results of the first three months of case ascertainment, 1st January 2017 to 31st March 2017.

A flow diagram of the potential cases of first seizure or newly diagnosed epilepsy from January 1st 2017 to March 31st 2017 is shown in Figure 2. Four hundred and forty two cases were initially identified. Following review of patient charts, 159 (40%) were excluded as they did not meet the inclusion criteria. The most common reason for exclusion was previously diagnosed epilepsy (n=69, 16%) or recurrent
provoked seizures (n=25, 6%). Two hundred and eighty-three patients were included and were distributed between 6 of the 7 hospitals in Cork city and county.

A pie chart illustrating the overlapping hospital-based sources of case ascertainment is shown in Figure 3. Of the 283 included cases, fifty-five percent (n=156) were identified by a single source of case ascertainment, 27% (n=76) were identified by two sources, 11% (n=31) by three sources and 7% (n=20) were identified by more than three sources of ascertainment. No single case was identified by all of the methods of case ascertainment. Regarding the contribution of individual sources of case ascertainment, almost 54% of cases were identified by reviewing radiology databases, see Table 2. The highest number of cases that was ‘unique to source’, meaning not identified elsewhere, came from review of E.E.G. databases.

Figure 4 illustrates the first round of postal surveys of G.P.s in the defined area. The response rate was 58%. Five G.P.s wrote back to indicate that they were retired and were therefore excluded from future surveys. Twenty-four potential cases were identified through this postal survey; however 8 were subsequently excluded as they did not meet the inclusion criteria. Sixteen cases were included in the study, none of which was unique to the G.P. survey in terms of source of case ascertainment. The community based patient advocacy group Epilepsy Ireland reviewed their records and did not yield any new cases diagnosed outside of a Cork hospital in the first three months of cases ascertainment.

Figure 5 illustrates the break-down of identified patients into each of the five subclassifications- first provoked seizure, first unprovoked seizure, new diagnosis of epilepsy, seizure mimics and indeterminate. Among the 95 cases of new diagnosis of epilepsy, 75 (79%) were patients who had a first seizure in 2017, and had an estimated greater than 60% chance of further seizures, therefore met the criteria for a practical definition of epilepsy [2]. Seventy-one of the new diagnosis of epilepsy were considered definite, and of these 71% (n=50) were focal, 18% (n=13) were generalized and in 11% (n=8) cases it was unknown whether epilepsy was focal or generalized according to the ILAE 2017 subclassification of seizure and epilepsy type. Seizure mimics represented the largest group (n=113, 40% of all reviewed case records) and the most commonly encountered seizure mimic was syncope (40%, n=45). There were 10 (3%) indeterminate cases in the first three months of data collection. Extrapolating the first three months of data, the crude incidence for new diagnosis of epilepsy in our population was 59 per 100,000 people per year, however it is necessary to highlight that this is an estimated rate which is likely to increase as it does not allow for ‘late entries’ due to lag in case ascertainment methods. The incidence is higher than previous studies outlined in Table 3 and therefore reinforces the importance of multiple methods of case ascertainment.

To estimate potential case under-ascertainment, 63 patients were identified by screening three months of multidisciplinary neuro-oncology meetings at a tertiary referral center. Of these, 5 had a previous diagnosis of epilepsy and 36 were not resident in the defined area therefore were not eligible for inclusion in the study.
The medical charts of the remaining 22 patients were reviewed and none had a documented seizure during follow-up.

5. Discussion

We present an epidemiological protocol that adheres to recently published international guidelines for investigation of the incidence of epilepsy within a population [4,10]. Furthermore, we have incorporated recently updated ILAE guidelines for the clinical classification of seizures and epilepsy [11,12] which have not yet been used in epidemiological studies but which will be of increasing importance in international epilepsy research. Our case ascertainment protocol is rigorous and, when data analysis is complete, will be the first to report the incidence of epilepsy, first unprovoked seizures and first provoked seizures in Ireland. Finally, to our knowledge, this will be the first study to additionally report the incidence and characteristics of ‘seizure mimics’ within the same population that seizures and epilepsy are under study. Inclusion of the related diagnosis of first seizures, epilepsy and seizure mimics in our study protocol gives a sense of the scale of this cohort and the significant impact on healthcare services investigating and treating these patients. Further details on age-adjusted incidence, seizure and epilepsy etiology, and classification will be reported separately when annual incidence data is complete.

Many previous epidemiologic studies have focused on a specific cohort within a population, for example children or the elderly. Relatively few have included all age-groups and, within those studies, all available case ascertainment methods range from use of a single source to multiple overlapping sources, see Table 3. With regard to case ascertainment in our study, we aimed for full capture of new events by combining as many methods of ascertainment as was practicable in our population. As shown in Table 3, the majority of other studies do not use such an extensive combination of methods. For example, Jallon et al., 1997 [20] used E.E.G. database as the sole method of case ascertainment. In our study, E.E.G. databases identified only 53% of cases. Furthermore, our interim three-month analysis demonstrates the importance of including all possible methods of ascertainment as no single case was picked up in every method. In addition, we specifically targeted inpatient teams and CNSs with a high likelihood of encountering a patient with a first seizure, for example, the neurosurgical and oncology teams, and contacted them on a regular basis to maximize case ascertainment. These methods identified 10 ‘unique to source’ cases, not identified by other methods in the first three months of case ascertainment. To our knowledge, this is the first study to apply such a method.

The seminal work by Hauser and Kurland, 1975 [21] in Rochester Minnesota demonstrates the usefulness of accurate medical record databases to perform epidemiologic studies. However, such medical record linkage systems are not available in many countries, including Ireland, and therefore in order to obtain accurate epidemiologic data overlapping methods of case ascertainment are required. Data obtained from cohort specific databases were used by Annegers et
al., 1999 [22] to describe the incidence of epilepsy in a multiethnic urban population (35.5 per 100,000). However, this database contained information only on those enrolled in a health maintenance organization served by a particular clinic and was therefore findings may not be generalizable to the whole population as the clinic served only people in employment and their dependents. Correspondingly, this study demonstrated low incidence among those over 65 years of age, in contrast to most other studies. Therefore, in populations where accurate, whole population, medical record databases are not in existence, multiple overlapping methods of ascertainment, in combination with medical chart review, as demonstrated by our study, provide detailed information for case identification, classification and determination of etiology.

An early study in Italy [25] was unique in its use of social workers and teachers in the community to maximize case ascertainment, see Table 3. However, due to ethical and practical considerations this would no longer be a viable source of case ascertainment. Finally, studies carried out in lower income countries, with less well developed health resources, have used door-to-door community surveys to ascertain cases, for example [26,27]. However, this method is vulnerable to under-ascertainment if stigma is associated with epilepsy and seizures in the population. Furthermore, accuracy of diagnosis and classification of cases is difficult when this method is used in isolation. Therefore, when adequate health resources permit, such as in our study, the use of hospital-based databases to identify and classify patients based on clinical information is preferable.

Other potential sources of case ascertainment were also considered. Previously published studies investigating adult populations have used A.E.D. prescription databases as a source of case ascertainment [23, 24]. However, it was not possible to use A.E.D. prescribing databases in our case ascertainment for a number of reasons. Firstly, drug prescribing data are not linked to other patient records in our population, therefore it would not have been possible to cross reference such information to our database. Secondly, patients presenting with a first seizure or seizure mimic may not routinely be prescribed an A.E.D. and therefore would not be captured by such a method. Thirdly, A.E.D.s are often used for indications other than epilepsy, such as neuropathic pain, and therefore may overestimate the diagnosis of epilepsy. Finally, use of a drug prescription database over the course of a year, without linkage to other patient records, would not give an accurate account of patients who were newly prescribed the medication. Therefore, in our population, such a method would be more appropriately applied to estimating prevalence of epilepsy rather than incidence, as was demonstrated by Linehan et al., in 2010 [13].

Very few studies address under-ascertainment. MacDonald et al., 2000 [28] cross-referenced general practice databases with local hospital patient administration system to try to ensure complete ascertainment. However, this method relies on accurate and complete case coding. For a number of patients, a seizure may not be the primary reason for attending hospital, for example, patients incorrectly initially diagnosed as a stroke, or patients who have their first seizure while medically unwell for another reason. To our knowledge, our study is the first to design a unique
method of following ‘at risk’ patients as a method of assessment of under-ascertainment. However, for the first three months of data collection, none of these at risk patients proceeded to have a seizure. We propose to follow this cohort for a longer time-frame in order to determine under-ascertainment and validate this unique method. On the other hand, it is reassuring that no community obtained case (from general practice or nursing homes) had not been identified already using the hospital-based methods, and therefore this serves as somewhat of a surrogate of complete ascertainment.

Our study highlighted the subjectivity involved in determining whether, following a single seizure, a person is at greater than 60% risk of further seizures, as found by our panel discussion. The Multicenter Trial for Early Epilepsy and Single Seizures (MESS) [29] provides some guidance on assessing this risk. However, the MESS study did not include neuroimaging as part of its risk stratification tool and there remains a lack of up-to-date, real world clinical data in certain populations, for example, older adults with established small vessel ischaemic disease with juxtacortical signal changes on brain imaging. In clinical practice, many patients commence an A.E.D. because they are considered at some risk of recurrent events, even though the exact risk is unquantifiable. The 2014 ILAE practical definition of epilepsy does not require the clinician to quantify the risk precisely. Therefore, we have included patients within the cohort of new diagnosis of epilepsy who were determined to be at an approximate 60% risk of recurrent seizures based on the evidence available to date. Further study of individuals at the margin of this risk threshold is required.

We are aware of potential gaps in case ascertainment, for example people who did not seek medical help after a seizure, people who do not recognize that they have had a seizure(s) and vulnerable patients unable to access medical care including homeless people, non-verbal individuals and people living alone. In addition, epilepsy and mimics largely are based on clinical history. In a small number of cases the treating physician may not recognize seizure as a potential diagnosis. However, through our study design, we have maximized case ascertainment in as much as possible.

6. Statements

6.1 Acknowledgement

The authors would like to thank all hospital-based medical, nursing, neurophysiology, radiology and medical records administrative staff as well as community-based G.P.s and nursing home staff who contributed and responded to the study. We would also like to thank Dr. Paul Corcoran (School of Public Health, University College Cork) and Dr. Carol Sinnott for advice during study design.

6.2 Statement of Ethics
Ethical approval was obtained from the Clinical Research Ethics Committee (CREC) of the Cork Teaching Hospitals. This was an observational study and following collection, all data were anonymized and stored on a password-protected computer in the Department of Neurology, Cork University Hospital.

6.3 Disclosure Statement

The authors have no conflicts of interest to declare.

6.4 Funding sources

No specific funding was received by any author with regard to this study.

6.5 Author Contributions

E. Maloney was involved in study design, data collection, data analysis and manuscript preparation. E. Chaila was involved in study design, data analysis and manuscript preparation. É. O’Reilly was involved in study design, data analysis and manuscript preparation. D. Costello was involved in study design, data collection, analysis and manuscript preparation.

7. References

19 Stosser S, Bockler S, Ludolph AC, Kassubek J, Neugebauer H: Juxtacortical lesions are associated with seizures in cerebral small vessel disease. J Neurol 2019
8. Figure Legends

Table 1. Examples of provoked seizures in association with disruption of the structural or functional integrity of the brain, adapted from the ILAE Standards for Epidemiologic Studies [4]

<table>
<thead>
<tr>
<th>Cause [14]</th>
<th>Period of occurrence</th>
<th>Notes/exceptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cerebrovascular disease [26-28]</td>
<td>First 7 days</td>
<td></td>
</tr>
<tr>
<td>Traumatic brain injury [28,29]</td>
<td>First 7 days</td>
<td>Included intracranial surgery. Long intervals are acceptable for subdural haematoma in the absence of known trauma or at first identification of haematoma. Subsequent seizures are unprovoked.</td>
</tr>
<tr>
<td>CNS infection [28]</td>
<td>First 7 days</td>
<td>Included seizures occuring after 7 days in patients with persistent clinical and/or laboratory signs of infection.</td>
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<tr>
<td>Cerebral tuberculoma [14]</td>
<td>During treatment</td>
<td>Seizures occuring after successful treatment are unprovoked</td>
</tr>
<tr>
<td>Brain abscess [14]</td>
<td>During treatment</td>
<td>Seizures occuring after successful treatment are unprovoked</td>
</tr>
<tr>
<td>HIV infection [14]</td>
<td>Acute infection or severe metabolic disturbance</td>
<td>Seizures occuring in the absence of oppotunistic CNS infection or severe metabolic disturbance are unprovoked</td>
</tr>
<tr>
<td>Arterovenous malformation [14]</td>
<td>In the presence of acute haemorrhage</td>
<td>All other seizures are unprovoked</td>
</tr>
<tr>
<td>Multiple sclerosis [14]</td>
<td>First presenting symptom within 7 days of relapse</td>
<td></td>
</tr>
<tr>
<td>Autoimmune disease [14]</td>
<td>Signs or symptoms of activation</td>
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</table>
Table 2. Cases identified at each source expressed to the nearest percentage. R.A.S.C. = Rapid access seizure clinic; C.N.S. = Clinical nurse specialist

<table>
<thead>
<tr>
<th>Source</th>
<th>Number of cases (%)</th>
<th>Unique to source (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiology</td>
<td>153 (54%)</td>
<td>45 (16%)</td>
</tr>
<tr>
<td>E.E.G.</td>
<td>151 (53%)</td>
<td>60 (21%)</td>
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<tr>
<td>Emergency department</td>
<td>70 (25%)</td>
<td>27 (9%)</td>
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<tr>
<td>Hospital Doctors</td>
<td>53 (19%)</td>
<td>9 (3%)</td>
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<tr>
<td>R.A.S.C.</td>
<td>45 (16%)</td>
<td>14 (5%)</td>
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<tr>
<td>Community survey</td>
<td>16 (6%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>C.N.S.</td>
<td>11 (4%)</td>
<td>1 (&lt;1%)</td>
</tr>
<tr>
<td>Epilepsy Ireland</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
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</table>
Table 3. Comparison of case ascertainment methods in previous studies investigating the incidence of epilepsy and/or first seizures within a population. Studies included all age groups within the population studied. E.E.G.= electroencephalogram. R.A.S.C= Rapid Access Seizure Clinic. G.P.= General practitioner.

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<tbody>
<tr>
<td>Geographic area (population)</td>
<td>Cork, Ireland, (542, 868)</td>
<td>Iceland, (882, 151)</td>
<td>London, United Kingdom, (100,250)</td>
<td>Martinique Island (383,596)</td>
<td>Genevea, Switzerland (384,657)</td>
<td>Southwest France (1,128,164)</td>
<td>Faroe Island (41,144)</td>
<td>Copparo, Italy (45,153)</td>
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<td>Northern Norway (215,000)</td>
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**Case ascertainment**

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</table>

**Incidence of first unprovoked seizures (per 100,000)**

| 56.8 | 11  | 64.1 | 45.6 | 71.3 (all seizures) | 23.6 (all seizures) |

**Incidence of first provoked seizures (per 100,000)**

| 16.4 | 25.2 |

**Incidence of epilepsy (per 100,000)**

| 33.3 | 46  | 42.8 | 33.1 | 48.7 | 32.8 |
Figure 1. Overview of case ascertainment, case analysis and case classification protocol. E.E.G.= electroencephalogram. R.A.S.C= Rapid Access Seizure Clinic. G.P.= General practitioner.
Figure 2. Flow diagram of identification of potential cases of first seizure and newly diagnosed epilepsy in the capture region over the study period 1st January to 31st March 2017.
Figure 3. Pie chart demonstrating overlapping sources of hospital based case ascertainment during study period 1st January to 31st March 2017 (n=283). The chart shows the proportion of included cases identified uniquely by each of the six hospital based methods as well as the proportion of cases identified by 2, 3 or more than 3 individual sources. C.N.S.= Clinical Nurse Specialist, E.D.= Emergency Department, E.E.G.= electroencephalogram, R.A.S.C.= Rapid Access Seizure Clinic.
Figure 4. Flow diagram illustrating case ascertainment by postal survey of all G.P.s in the capture region during study period 1st January to 31st March 2017. A total of 16 cases were included from this method of ascertainment, all of whom had been previously identified through hospital based ascertainment. G.P.= General Practitioner
Figure 5. Number of cases in each subgroup following panel review of case information. Of the 95 cases of new diagnosis of epilepsy, 75 also presented with their first seizure and were determined to be at greater than 60% risk of recurrent seizures. IQR= interquartile range, m=months, w=weeks, y=years.