Schizophrenia is not disappearing in south-west Scotland

J. ALLARDYCE, G. MORRISON, J. VAN OS, J. KELLY, R. M. MURRAY and R. G. McCREADIE

Background Recent work has reported a decline in the incidence of schizophrenia, but it is unclear if these findings reflect a true decrease in its incidence or are an artefact arising from methodological difficulties.

Aims To take account of these methodological difficulties and report service-based incidence rates for schizophrenia in Dumfries and Galloway in south-west Scotland for 1979–98.

Method Using both clinical diagnoses and diagnoses generated from the Operational Checklist for Psychotic Disorders (OPCRIT) computer algorithm for ICD–10 and DSM–IV schizophrenia, we measured change in the incidence rates over time. We used indirect standardisation techniques and Poisson models to measure the rate ratio linear trend.

Results There was a monotonic and statistically significant decline in clinically diagnosed schizophrenia. The summary rate ratio linear trend was 0.77. However, using OPCRIT-generated ICD–10 and DSM–IV diagnoses, there was no significant difference over time.

Conclusions OPCRIT-generated consistent diagnoses revealed no significant fall in the incidence of schizophrenia. Changes in diagnostic practice have caused the declining rates of clinically diagnosed schizophrenia in Dumfries and Galloway.

Declaration of interest This work was supported by the Stanley Foundation.

Dramatic declines (up to as much as 50%) in the first admission rates for schizophrenia have been demonstrated in different countries and continents over the past few decades (Munk-Jørgensen, 1986; Joyce, 1987; Der et al, 1990; Susvisaari et al, 1999). Several of these reports have come from Scotland (Eagles & Walley, 1985; Geddes et al, 1993). It is still not clear if this replicated finding reflects a true decrease in the incidence of schizophrenia or is simply an artefact (Jablensky, 1997). The uncertainty is due to the considerable methodological difficulties inherent in measuring trends over time in the incidence of schizophrenia. Because it is a rare disorder, its rates are sensitive to distortion by factors such as change in the underlying population structure, change in diagnostic habits, administrative inaccuracies when recording first admission or contact data and change in the organisation and delivery of psychiatric services (Kendell et al, 1993). In this study we set out to take account of these potential confounders and report service-based incidence rates for schizophrenia in Dumfries and Galloway in south-west Scotland over the 20-year period 1979–1998.

METHOD

Case identification

Dumfries and Galloway is a geographically well-defined area in south-west Scotland. It has a stable population of around 147000, of whom 99.5% are White. Psychiatric services are provided via one health trust and there is little or no private health care.

We identified all patients in contact with the psychiatric services between the years 1979 and 1998 who were given a clinical diagnosis of one of a broad range of psychotic disorders. During this period two different versions of the International Classification of Diseases (ICD) were used to code cases formally: ICD–9 (World Health Organization, 1978) from 1979 to March 1996, and ICD–10 (World Health Organization, 1992) thereafter. We reviewed all cases with a diagnosis of schizophrenia (295, F20), schizoaffective disorder (295.6, F25), delusional disorder (297, F22), mania (296.0, 296.2, 296.4, F30, F31.1, F31.2, F31.6), acute, transient or unspecified psychotic disorders (298, F28, F29) or drug-induced disorders (292.1, F12.5, F16.6, F19.5, F12.75, F16.75, F19.75).

This list of possible cases was derived from two sources: data for all in-patients held centrally by the Information and Statistical Division of the Scottish Office (ISD); and locally held registers of outpatients, domiciliary visits and ‘out of hours’ referrals. Cases were excluded if they had presented with a psychotic episode before 1979 or were not resident in Dumfries and Galloway at the time of their presentation. The case records (including medical, nursing, social work and occupational therapy notes and all correspondence) of the remaining patients with first-episode psychosis were examined and the Operational Checklist for Psychotic Disorders (OPCRIT) (McGuinness et al, 1991) was completed. The OPCRIT is a symptom checklist containing 90 items, which explores aspects of the mental state examination and the psychiatric history. It has a glossary of clear and explicit descriptions of each constituent item of psychopathology and instructions for coding them. It was designed with case note review in mind. Two of the authors (J.A., G.M.) completed the OPCRIT for the year following the initial presentation. This allowed us to compare each patient using similar information, irrespective of year of presentation. Case notes were examined in random order of date of presentation and raters were blind to the clinically coded diagnosis.

There was good interrater reliability for ICD diagnosis of schizophrenia: κ=0.79 (P=0.004). The OPCRIT checklist was then used to generate ICD–10 and DSM–IV (American Psychiatric Association, 1994) diagnoses via the associated computer algorithm OPCRIT 3.4.

General population data

(population at risk)

The General Register Office of Scotland provided detailed population data for the region. Census data for the years 1971, 1981 and 1991, stratified by age and gender, were provided, and population estimates for the intermediate years were
interpolated. The under-enumeration in Dunfries and Galloway was estimated at around 1%.

**Statistical analyses**

Analyses were carried out in four blocks of 5 years: 1979–83, 1984–88, 1989–93 and 1994–98. Incidence rates were calculated, stratified by age and gender, for the years 1979–83. These rates were used as our reference (standard) population rates. Using the indirect standardisation method, we applied this rate to the population structure in the subsequent year blocks. This allowed us to determine the number of expected cases. By dividing the number of cases actually observed by the number of cases expected, we calculated the standardised incidence ratio (SIR). Summary rate ratio linear trend, indicating the summary increase in risk with each consecutive time period, and gender modification effects were calculated using the Poisson Regression Procedure from the STATA program (Stata Corporation, 1999). These analyses were carried out for those patients receiving a clinical diagnosis of schizophrenia: OPCRIT-generated research diagnoses of ICD–10 and DSM–IV schizophrenia, as well as DSM–IV schizophrenia and schizoaffective disorder combined.

**RESULTS**

The number of patients with any one of the inclusion diagnoses in contact with the service during the 20-year period was 1460. After excluding cases with a previous psychotic episode prior to 1979, or whose home address was outside the catchment area, there were 464 cases for which the OPCRIT was completed. There was a monotonic decline (the incidence for each year was less than or equal to the previous one) over the 20-year period in the rates for clinically diagnosed schizophrenia. This was statistically significant. The summary rate ratio trend was 0.77 (95% confidence interval (CI) 0.68–0.88). However, using OPCRIT-generated diagnoses there was no significant difference in the rates over time, irrespective of the diagnostic system applied. The summary rate ratio linear trend for ICD–10 schizophrenia was 0.98 (95% CI 0.87–1.10) and for DSM–IV schizophrenia it was 1.10 (95% CI 0.85–1.19). As previous studies have analysed DSM–IV schizophrenia and schizoaffective disorder combined, we did the same; the summary rate trend was 0.89 (95% CI 0.86–1.11).

As recently published work has shown gender differences in the incidence of schizophrenia (De Alacron et al, 1990; Der et al, 1990), we examined the modification (interaction) effect of gender on the rates over consecutive time periods. There was no significant gender–time interaction in the models. For clinically diagnosed cases, gender–time interaction was 0.89 (CI 0.69–1.16) for OPCRIT ICD–10 schizophrenia it was 0.98 (CI 0.87–1.10) and for OPCRIT DSM–IV schizophrenia it was 0.87 (CI 0.62–1.22).

**DISCUSSION**

**Methodological issues**

**Strength of the study design**

(a) This study was carried out in an area with a stable population and no significant ethnic minority groups.

(b) By identifying not only all admissions to in-patient care, but also all day patients, out-patients, domiciliary visits and informal out-of-hours contacts, we avoided possible confounding effects due to the changes in service delivery.

(c) By reviewing all contacts with the service, we identified patients given a diagnosis of a psychotic disorder at first psychiatric presentation and also those psychotic patients who had previous periods of psychiatric care for non-psychotic episodes. This gives the most representative sample of first-ever psychosis (Driessen et al, 1998).

(d) Standardised rates were used to ensure that any change in the at-risk population has not led to spurious changes in the incidence rates reported.

(e) Reviewing the case notes of all contacts during the 20 years reduced distortion from inaccurate administrative coding of first contacts.

(f) Finally, any analyses of rates over a long time period will be susceptible to bias

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from gradually changing diagnostic practices, so that by using the OPCRIT checklist and computer algorithm we ensured a consistent diagnostic approach for all patients.

Limitations of the study design
There are a number of limitations to bear in mind, although we believe they are of minor importance. We report service-based rates for one geographical area. This could introduce bias if the referral patterns of general practitioners changed during the study. However, it is assumed in well-developed services in the UK that nearly all patients with schizophrenia come into contact with services sooner or later (Cooper et al, 1987; Kendall et al, 1993).

Moreover, case note reviews may not be the best way to make a diagnosis. However, OPCRIT has been used successfully in a number of clinical and epidemiological studies (Williams et al, 1996). The two raters in our study were specialist registrars in psychiatry with demonstrated good interrater reliability, and the case notes reviewed were comprehensive. There is, however, a small possibility that the contents of case notes have changed over the study period.

Interpretation of the findings
We have shown a decline in the first contact rates for clinically diagnosed schizophrenia. This is consistent with previous studies in Scotland (Eagles et al, 1988). There is however no such decline when ICD–10 and DSM–IV diagnostic systems are consistently applied to all cases of psychotic disorders.

The difference between rates for clinically diagnosed schizophrenia and the OPCRIT-generated rates suggests that changes in diagnostic habits have indeed operated to confound the reported rates. The OPCRIT procedure resulted in a number of cases who were clinically diagnosed as suffering from schizophrenia in the early years of the period studied being reallocated to a range of other diagnostic categories, whereas the reverse tended to take place in the later years. This must reflect an increasing hesitancy or caution in making a diagnosis of schizophrenia.

There are at least two possible reasons for this. First, clinicians may now consider reliability of diagnosis important and therefore be using more restrictive operational criteria (Crow, 1990). Second, there may be an increasing reluctance at the initial presentation to make a diagnosis with such profound prognostic implications without first reviewing all the possible differential diagnoses. That clinicians may now be using narrower criteria for diagnosing schizophrenia has been recognised in several studies (Parker et al, 1985; Munk-Jørgensen, 1986; Kendall et al, 1993); our findings strongly support this assertion.

The findings of this study are at odds with most of the recently published work investigating trends over time in the incidence of schizophrenia. However, other studies which have closely considered diagnostic difficulties in their methodology have failed to show the magnitude of decline reported in studies depending on routinely collected data (Castle et al, 1991; Brewin et al, 1997). Also, the Scottish age-period–cohort study of Takei et al (1996) showed that most of the decline in the incidence of schizophrenia could be accounted for by changes at or near the time of diagnosis rather than by aetiological factors operating early in life.

We conclude that changes in diagnostic practice have resulted in declining rates of clinically diagnosed schizophrenia in the area we have studied. Our results show no sustained fall in the incidence of consistently diagnosed cases of schizophrenia. In southwest Scotland we have found no evidence to support the view that schizophrenia is disappearing.

ACKNOWLEDGEMENTS

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