Assessment and management of suicide risk in primary care

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Abstract

Background

Risk assessment and management of suicidal patients is emphasized as a key component of care in specialist mental health services, but these issues are relatively unexplored in primary care services.

Aim

To examine risk assessment and management in primary and secondary care in a clinical sample of individuals who were in contact with mental health services and died by suicide.

Method

Data collection from clinical pro formas, case records, and semi-structured face-to-face interviews with general practitioners.

Results

Primary and secondary care data was available for 198 of the 336 cases (59%). The overall agreement in the rating of risk between services was poor (overall kappa = 0.127; p = 0.10). Depression, care setting (post discharge), suicidal ideation at last contact and a history of self-harm were associated with a rating of higher risk. Suicide prevention policies were available in 25% of primary care practices and 33% of staff received training in suicide risk assessments.

Conclusion

Risk is difficult to predict, but the variation in risk assessment between professional groups may reflect poor communication. Further research is required to understand this. There appears to be a relative lack of suicide risk assessment training in primary care.
**Keywords:** suicide, risk assessment, general practitioners, primary care, secondary care.
**Background**

Suicide is a major public health problem internationally and in the UK.\(^1\) The majority of individuals are in contact with their general practitioner (GP) prior to suicide,\(^2\) particularly those with a mental illness who consult more frequently than other patients.\(^3\) People with a mental illness are at increased risk of suicide.\(^4\) As such, GPs are well placed to intervene and potentially improve outcome in patients at risk of suicidal behaviour.

To date, suicide prevention in primary care has largely focused on training GPs to identify, assess, and manage suicide risk in patients with a primary diagnosis of depression.\(^5,6,7\) Risk assessment and management of suicidal patients is emphasized as a key component of care in specialist mental health services, but these issues are relatively unexplored in primary care services. Knowing whether GPs are completing risk assessments, if they are identifying the ‘right’ factors to formulate risk assessments and whether practice is consistent between GPs and mental health specialists are important issues. Evidence suggests that different professional groups vary in their risk assessments.\(^8\) For example, one study found that non-specialists were more cautious in their risk assessment of patients who had self-harmed than mental health staff, but no less accurate in predicting repeat episodes.\(^9\) Understanding GPs procedures for assessing and managing risk in primary care is potentially important for future suicide prevention but the factors GPs take into account when assessing risk in primary care for patients who go on to die by suicide are relatively unexplored.
The overall aim of the study was to examine risk assessment and management prior to suicide in primary and secondary care in a sample of mental health patients who died by suicide. The specific objectives of this study were to:

1) compare the level of agreement of risk assessments in primary and secondary care;

2) investigate the factors associated with high versus low ratings of risk in primary and secondary care;

3) describe the primary care management including liaison between primary and secondary care for individuals assessed at different levels of risk;

4) describe key elements of suicide risk assessment in primary care, including the extent of GP training on risk assessment, and the policies and procedures in place in primary care.
Method

Setting and participants

This study was carried out in the English National Health Service (NHS). Most of the population is registered with a primary care physician – a general practitioner or ‘GP’. Specialist mental health services (generally referred to as ‘mental health trusts’) provide care to those seeking treatment. Primary care and specialist treatment, like other services provided by the National Health Service, is free at the point of use.

The sample for this study included individuals residing in the North West of England who had been in contact with mental health services before death and had died by suicide between 1st January 2003 and 30th June 2007.

Collection of suicide data from mental health services

Mental health service data were collected through the National Confidential Inquiry into Suicide and Homicide, a well-established national clinical study. Information on all general population suicides and deaths from undetermined external causes is obtained from the Office for National Statistics. Mental health services identify those people in contact with services in the 12 months before death. Clinical data on these patients is then obtained through questionnaires sent to consultant psychiatrists. The questionnaire comprises multiple sections including: demographic information (e.g., age, sex), clinical history (e.g., primary psychiatric diagnosis; history of self-harm), and clinical management (e.g., treatment and compliance with medication; last contact with mental health services). The Inquiry, therefore, provides comprehensive information on a
UK-wide case series of all suicides by people in mental health service contact in the year prior to death. The ascertainment procedures are robust and the response rates high (over 95%). Further details are available in previous publications.11

Collection of primary care and interview data

Contact details for GPs were obtained from coroner files or administrative departments of NHS Trusts. A retrospective review of primary care medical records (both written and electronic) and, where possible, semi-structured interviews were conducted with consenting general practices. A medical records proforma and interview schedules were used to collect data. These were adapted from tools used in previous research2 and included details of physical and mental health problems reported in all consultations and treatment offered in the year before death (specifically the final consultation), GP views on their concerns for the patient, antecedent factors contributing to death, and factors which could have prevented the death. Information on the mental health treatment a patient received in the 12 months prior to death was determined by direct inquiry from clinicians and also using evidence in the medical records. Additional interview data were collected on GP training on self-harm, suicidal ideation and suicide risk assessments and the policies GPs followed for patients at risk of suicide. The researcher and GP had access to the written or electronic medical notes in every interview providing a source of contemporaneous data.

All interviews and most case record reviews took place in the GPs’ surgeries, but some records were viewed at Primary Care Trust premises. Interviews were audio-recorded where participating GPs consented to this. Case reviews using the proforma took approximately one hour and interviews varied between 20-40 minutes.
**Ratings of risk.**

For the patients who died by suicide, clinicians in secondary care had estimated the risk at last contact on the Inquiry questionnaire as high, moderate and low or no risk, which we dichotomised for the purposes of the analysis (see below). Unlike in our previous paper, in this study we considered immediate and long-term risk identified by mental health teams not just immediate risk. Mental health services rate only a small minority of individuals (approximately 10%) who have died by suicide as at high immediate risk at last contact. Maximising the time horizon in this study meant that a greater number of potentially high risk individuals were included. We also thought that these more inclusive criteria were more comparable with the GPs ratings (which had no time cut off). In primary care we used the variable recording of whether GPs had been concerned for their patient’s safety prior to death as a proxy measure for high risk (with no concern being taken as a marker of low risk). The primary care risk data were obtained through the GP interviews.

**Ethical approval**

Ethical approval was granted by the North-west Research Ethics Committee (REC reference: 02/8/74) for the Inquiry as a whole and individual R&D approvals were obtained from all the relevant Mental Health Trusts included in the study.

**Analysis**
The secondary care assessments of risk were divided into two categories: 1) perceived moderate and high risk (hereafter referred to as high risk) and 2) perceived low or no risk (hereafter referred to as low risk). We took this approach because previous studies have suggested that even those rated as at moderate risk have a high incidence of adverse outcomes.\(^9\) Many of the independent variables in the questionnaire pro formas were in binary format (presence/absence of factors). For variables providing more than two possible responses, the main factor of interest was selected and the response recoded following a binary format. As some of the responses occurred in only a small number of cases; binary coding provided more reliable estimates of odds ratios.\(^{12}\) The independent variables were listed under the broad headings: demographic factors, clinical features and behavioural features (see Table 2 and 3).

Primary care and secondary care data were analysed separately. Descriptive analyses were used to compare the management and treatment of patients in primary care in the year prior to suicide. Both parametric and non-parametric statistical tests were carried out where appropriate. These included the chi-squared test of association, kappa statistic and analysis of variance. When percentages are quoted these refer to ‘valid cases’ (i.e. cases for which the relevant information was available). As a result the denominator varies between items. With respect to the interviews, much of the data were presented as simple frequencies.

We carried out forward stepwise logistic regression using SPSS version 20\(^{13}\) to compare the factors associated with primary care and secondary care doctors’ ratings of the perceived risk of suicide. We were looking at the best fitting data driven model for these variables. We have reported the frequencies, odds ratios, their confidence intervals and \(P\) values.
Results

Characteristics of the sample

Between 2003 and 2007, 336 patient suicides were recorded in the North West of England; approximately 6% of the entire national sample (n=5,552). GP records were reviewed in 291 (87%) cases and semi structured interviews were completed by 198 (59%) consenting GPs. Baseline characteristics of the patients are given in Table 1.

In terms of sex, age, civil status, living circumstances, employment status, and clinical characteristics no significant differences were noted between the patients for whom GP data were obtained and those for whom it was not (n = 45). Reasons for non-participation of GPs in interviews where medical records were available were: perceived lack of time; GP had retired or left the practice, no other GP knew the patient; or the GP had died.

Reported risk: comparison of primary care versus secondary care

Of the 198 cases, 162 (82%) patients had both primary and secondary data on risk assessment available. In 73/162 patients (45%) both GPs and mental health specialists rated risk at last contact as low, perhaps particularly surprising given that all of the sample had died by suicide.

There was overall agreement in the rating of risk between primary and secondary care for 60% of patients and disagreement in 40% (overall kappa = 0.127; p = 0.10; strength of agreement: poor). High risk was identified more often by secondary care than primary care (40% v 30%). In only 24 cases (15% of the suicide deaths) did both sets of clinicians rate the risk as high.
Factors associated with risk prior to suicide

Primary care

Data on GP concerns for safety were available in 189 (95% of 198) cases, of which 53 (27%) reported concerns for patient safety. Table 2 shows the factors associated with GPs ratings of risk (being concerned or not concerned about the safety of their patient). Four variables (depression; care setting- post discharge; suicidal ideation at last contact with primary care; and primary care consultation following self-harm) were individually associated with perceived risk. Stepwise regression was used, however the final logistic model included only one variable - suicidal ideation at last contact with primary care (OR: 21.61; 95% CI: 7.20-64.84; p-value<0.001) as an independent risk factor of perceived high risk.

Secondary Care

Data on secondary care patients’ reported risk were available for a total of 170 cases (86% of 198), of which 69 (41%) were rated at high or moderate risk. Table 3 shows the factors associated with an assessment of higher risk in secondary care. Three variables (care setting- post discharge; having suicidal ideas at last contact with secondary care; and a history of self-harm) were individually associated with perceived risk. The stepwise regression model included all three variables as independent risk factors of perceived high risk: care setting- post discharge (OR: 3.28; 95% CI: 1.29-8.38; p-value=0.013); having suicidal ideas at last contact with secondary care (OR: 3.43; 95% CI: 1.10-10.70; p-value=0.034); and a history of self-harm (OR: 3.86; 95% CI: 1.77-8.45; p-value=0.001).

Treatment
**Primary care**

Table 4 examines the association between consultation and treatment in primary care and GP’s ratings of risk. In terms of the consultation data, 186 (98%) patients consulted in the twelve months prior to their death. Reasons for the final consultation and the type of treatment offered at final consultation were significantly related to GP’s concern for their patient’s safety.

**Secondary care**

Table 5 examines the association between consultation and treatment in primary care and secondary care ratings of risk for a total of 170 (86%) patients where information on risk was available from secondary care. In terms of the consultation data, 165 (97%) patients consulted in the twelve months prior to their death. There were no significant differences between the groups.

**Policies, Procedures and Training in the primary care setting**

Only one in four practices had written policies to follow regarding suicide or self-harm (table 6) and one in five of those practices were unable to provide any specific information about what policies they followed. A third of the practices had training in place on suicide awareness and self-harm and on risk assessment for suicide. Training was not available to all staff in the practice and tended to be restricted to GPs. A quarter of the GPs had received training on only one occasion during their practising years, a third received ad hoc training, and a fifth could not report any information about their training.
Discussion

Main findings

This study is the first to our knowledge to compare assessments of risk and characteristics of patients treated by both primary and secondary care in the year prior to suicide. Many patients had been rated as at low risk of suicide prior to death. The level of agreement with respect to risk assessment in primary and secondary care was poor. In primary care, factors such as depression, care setting (post discharge), suicidal ideation at last contact with primary care and primary care consultation following self-harm were associated with GPs being concerned about their patient. Those who the GPs were concerned about were more likely to have consulted for psychological reasons and were more likely to have been referred to specialist services. In secondary care factors such as care setting (post discharge), having suicidal ideas at last contact with secondary care; and a history of self-harm were associated with a rating of high or moderate risk. There were few policies in primary care to guide practice and a discernible lack of training on suicidal behaviour and risk. We think that together these findings are of interest to those who plan and provide services and make a strong case for better more integrated assessment and management of suicide risk.

Methodological considerations

Our findings should be interpreted in the context of a number of methodological limitations. Both primary and secondary care clinicians were contacted after death. This may have resulted in recall bias. However, our main aim was to compare specialist and primary care risk assessments and understand the possible reasons for the rating of low risk in patients who went on to die by suicide. This could only be done with a clinical sample of people who were in contact with both...
mental health and primary care services. A prospective study would not have been feasible. In
addition, the researcher and GP had access to the written or electronic medical notes in every
interview and were therefore using data that was collected contemporaneously at the time the
patient consulted with their GP in order to inform their responses. Whilst the retrospective
assessment of risk has its potential weaknesses, it is a well-established methodology, used for
example in the UK’s National Confidential Inquiry into Suicide for a number of years. Previous
studies have identified variations in accuracy or the amount of detail provided in case records;
however, one study comparing GP records and patient self-report questionnaires found similar
figures for the mean number of consultations in both sources.\textsuperscript{14} A systematic review into the
quality of computerised medical records found that the recording of consultations on such systems
tended to be good.\textsuperscript{15} In addition our primary care data was supplemented by interviews with GPs
and our secondary care data collection was collected directly from the clinicians caring for the
patients by means of a dedicated proforma.

It is also possible that some clinicians may have been concerned about their own assessment of
suicidal risk and this may have introduced some bias (for example, with a tendency for some
clinicians to perhaps downgrade the estimated level of risk when the patient was last seen). The
emotional aspect of suicide and the personal role of the GP in (not) preventing the attempt could
have contributed to a possible ‘defensive reporting’ of the signals that were missed or wrongly
interpreted. In addition it should be borne in mind that risk is dynamic – it changes over time.
Some of the differences between primary and specialist services’ views of risk might be because
the risk assessments were carried out at different times. Equally some of the discrepancies
between primary and secondary care ratings of risk could have been due to the slightly different
nature of what clinicians were asked – in primary care it was about concerns about the patient’s safety.

The sample consisted of people in current, or recent, contact with mental health services from the North West of England and no comparison group of those who did not die by suicide, nor individuals who died by suicide and did not have contact with mental health services. Our risk recognition findings might in some senses represent a ‘best case’ scenario as ours was a clinical sample of patients in contact with services at the time of death. The recognition of risk is likely to have been lower in those not in contact with primary care or mental health services. In this context, we think our findings on the large number of individuals rated as at low risk prior to death are even more striking.

Our findings may not be representative of the rest of the UK although many of the issues we identified are likely to apply across services. It should also be noted that some of our data are now several years old. As a consequence some of the study findings might not necessarily reflect current clinical practice.

**Clinical and Research implications**

The assessment of suicide risk is clearly difficult. Part of the challenge is the poor predictive value of assessments and scales which means that many individuals rated as high risk will not go on to have adverse outcomes (i.e. ‘false positive’ on the basis of risk assessment).\(^{16,17}\) In this study we focussed on the complimentary issue of ‘false negatives’. Similar to previous studies we found
that many patients who died (nearly half the sample) had in fact been rated as at low risk when they were last seen by their clinicians.\textsuperscript{18} This ‘low risk’ paradox in patients who go on to die by suicide reflects the problems inherent in predicting low frequency events, but rapidly changing risk, desensitization to high risk situations (particularly in specialist care), and recall bias might also contribute.\textsuperscript{18} In primary care, presentation with physical complaints could mask psychological symptoms and lead to a downgrading of risk.

In this study we also found that there were potentially important differences in the risk assigned to patients in primary and secondary care prior to their death. Poor communication between care settings could account for these findings. Previous studies have highlighted poor communication and sharing of information between general and mental health services.\textsuperscript{19} Strengthening communication and liaison links between care services could lessen discrepancies and contribute to suicide prevention. The rapid improvement of information technology may facilitate both the collection and communication of risk information.\textsuperscript{20} Systems could flag up patients at risk, indicate who is responsible for follow up care and may be updated regularly, including for patients who do not attend appointments. Further research to investigate and understand the variation in risk assessment among health professionals may also help to improve practice.

Our data suggest that clinicians do take into account a number of important factors when assessing risk. Comprehensive risk assessments which take into account a wide range of demographic and clinical factors (for example, employment, living circumstances, age, gender, history of self-harm or substance misuse, physical health) are an important suicide prevention measure.\textsuperscript{2} Patients who express suicidal ideation at their last consultation but who are rated as low risk (6 patients in
primary care and 7 patients in secondary care in this study) may be an important group in whom to intervene.

We found a comparative lack of training for suicide risk assessment in primary care. This is consistent with previous studies.\textsuperscript{17,21} It is clear from guidelines\textsuperscript{22} and existing research that the evidence to guide the content and format of suicide risk assessment training is lacking. The use of screening or case finding instruments, for example the Columbia Suicide Severity Risk Scale (C-SSRS) which can be used by gatekeepers with minimal training may be a promising avenue to explore further.\textsuperscript{23} Of course, there is the caveat that it will always be extremely difficult to predict low incidence events like suicide.

Conclusion: 
Risk is difficult to predict, but the variation in risk assessment between professional groups may reflect poor communication. There appears to be a relative lack of suicide risk assessment training in primary care. Further research into the assessment and management of suicidal behaviour in primary care has the potential to contribute significantly to evidence-based suicide prevention.

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\textbf{Ethics:} North West Multi-centre Research Ethics Committee, 02/8/74.
Competing interests

NK was Chair of the Guideline Development Group for the National Institute for Clinical Excellence (NICE) guidelines for the longer term management of self-harm and sits on the Department of Health’s National Suicide Prevention Strategy Advisory Group.

Authors' contributions

The study was principally designed by PS, KW, NK but all authors had input into aspects of study design. Ethical approval was obtained by PS and KW. Data collection was carried out primarily by PS, supported by NS and KW. Initial data manipulation was carried out by PS supervised by DW. Data analysis was carried out by PS and supervised by DW and NK. Clinical input was provided by NK. The manuscript was prepared by PS with supervision from NK, KC and KW. All authors commented on drafts of the paper and contributed to the final version.

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