Histological precursors of oesophageal squamous cell carcinoma: results from a 13 year prospective follow up study in a high risk population

G-Q Wang, C C Abnet, Q Shen, K J Lewin, X-D Sun, M J Roth, Y-L Qiao, S D Mark, Z-W Dong, P R Taylor, S M Dawsey

Oesophageal squamous cell carcinoma is a common malignancy with a very poor prognosis. It is the sixth most common fatal cancer in the world, causing over 300,000 deaths each year. Between 1992 and 1999, the five year relative survival rate for oesophageal cancer in the USA was 14.0%, among the lowest for all cancers. The main reason for this poor prognosis is that most cases are asymptomatic and go undetected until they have spread beyond the oesophagus and are unresectable. In this setting, successful strategies for primary prevention and early detection of curable lesions are critically needed to control this disease.

Worldwide, the great majority of oesophageal cancers are oesophageal squamous cell carcinomas (OSCC). One important step in designing effective prevention and early detection strategies for OSCC is the identification of valid histological precursors that can be used as intermediate end points in prevention research and as targets for early detection screening and treatment. Identification and validation of such precursors requires classification of lesions by well defined histological categories and subsequent prospective follow up of biopsied patients to document which precursor lesions are valid and can be used as targets for screening and treatment.

In this study, squamous dysplasia and carcinoma in situ were the only histological lesions associated with a significantly increased risk of developing OSCC within 13.5 years after endoscopy. There was no evidence that oesophagitis predisposed to this tumour. Increasing grades of dysplasia were strongly associated with increasing risk, indicating that the histological grading was clinically meaningful. The follow up experience of severe dysplasia and carcinoma in situ was equivalent, suggesting that this distinction is not clinically relevant. Documenting these precursor lesions of OSCC should assist in the development of effective prevention, early detection, and treatment strategies for this disease.

Materials and Methods

Endoscopic survey

Endoscopic examinations were conducted among subjects enrolled in the Linxian Dysplasia Trial, a six year randomised prospective nutrition intervention trial limited to individuals with a previous Chinese cytological diagnosis of oesophageal dysplasia. Active intervention, consisting of daily tablets containing 26 vitamins and minerals at 2–3 times US Recommended Daily Allowances or matched placebos, began on 1 May 1985 and ended on 30 April 1991.

Abbreviations: BCH, basal cell hyperplasia; CICAMS, Cancer Institute of the Chinese Academy of Medical Sciences; CIS, squamous carcinoma in situ; dysplasia NOS, dysplasia not otherwise specified; OSCC, oesophageal squamous cell carcinoma; IEN, intraepithelial neoplasia; NCI, US National Cancer Institute
Normal
A stratified squamous epithelium was present which showed no features diagnostic of the other histological categories listed below. Mature squamous cells with abundant clear cytoplasm, scattered lymphocytes, and compressed nuclear fragments (“squiggly cells”) were occasionally seen in the epithelium. The lamina propria, if present, commonly contained a few scattered mononuclear inflammatory cells.

Acanthosis
An otherwise normal epithelium was \( \geq 0.5 \) mm thick.

Oesophagitis
One or more of the following three criteria were present: elongation of lamina propria papillae into the upper third of the epithelium together with basal cell hyperplasia (BCH) \( >15\% \) of total epithelial thickness; epithelial infiltration by neutrophils or eosinophils; or a dense non-follicular infiltrate of mononuclear inflammatory cells or neutrophils in the lamina propria.

Basal cell hyperplasia (BCH)
An otherwise normal epithelium had a basal zone thickness \( >15\% \) of total epithelial thickness, without elongation of lamina propria papillae or other abnormality.

Squamous dysplasia
Nuclear atypia (enlargement, pleomorphism, and hyperchromasia), loss of normal cell polarity, and abnormal tissue maturation were present in the lower third (mild), in the lower two thirds (moderate), or in all thirds (severe) of the epithelium, without full thickness involvement or invasion. Dysplastic biopsies which could not be graded because of biopsy size or orientation were categorised as squamous dysplasia, not otherwise specified (NOS).

Squamous carcinoma in situ (CIS)
Dysplastic squamous cells were present throughout the full thickness of the epithelium, without invasion.

Squamous cell carcinoma
Neoplastic squamous cells were present which had invaded through the basement membrane.

Follow up procedures
Incident cancers and deaths were identified through several methods which assured essentially complete ascertainment of these events.5 During the intervention period, all trial participants were visited each month by their village doctors to deliver the intervention pills, and all those with cancer symptoms and those who died from any cause were identified and reported. In addition, all medical facilities, including commune hospitals, the Linxian County Cancer Hospital, and the Cancer Hospital in the prefecture capital of Anyang, notified investigators of all cancer diagnoses among residents of the communes in the Dysplasia Trial. After the active intervention, information was obtained every month from village doctors on the vital status, occurrence of any incident cancer, and cause of death for each trial participant. In addition, all living participants were individually surveyed by questionnaire at the end of intervention in 1991, in 1996, and again in 2000. Throughout the trial and the post trial follow up period, symptomatic individuals were referred to the CICAMS field station in Yaocun Commune or to their commune hospital for further clinical evaluation. Case records and diagnostic materials (histology slides, cytology slides, and/or x ray films) of all subjects developing cancer were reviewed and the diagnosis of cancer confirmed by members of an International Endpoints Review Committee.
Histological precursors of oesophageal SCC

composed of expert pathologists, cytologists, and radiologists from both the US and CICAMS.

Analysis
In the 1987 endoscopic survey, 754 of 833 endoscoped subjects had one or more squamous biopsies which were satisfactory for histological diagnosis. Of these, 72 had received a diagnosis of cancer (of any site) before or during the 1987 endoscopy examinations. The analytic cohort for this study consisted of the remaining 682 subjects who had at least one satisfactory squamous biopsy and were free of invasive cancer at the beginning of the follow up period.

The age, sex, smoking status (ever smoked regularly ≥6 months), and alcohol use (any drinking of alcohol during the last 12 months) were recorded by questionnaire. The pretrial cytology diagnoses (dysplasia 1 [low grade]; dysplasia 2 [high grade]) came from balloon cytology examinations performed in 1983. For each subject, a worst oesophageal diagnosis was determined using the hierarchy of invasive cancer > carcinoma in situ > dysplasia (graded) > oesophagitis > acanthosis > BCH > normal.

Descriptive statistics of the follow up results were generated based on all subjects in the analytic cohort, with deletions as necessary for missing data. Age adjusted oesophageal cancer incidence rates for each of the histological categories were calculated by determining the number of cases which had occurred in each age and histological category and dividing that number by the number of person years of observation in that category. Rates were age adjusted using the 1987 age distribution of all subjects in the analytic cohort as weights (≤50 years = 22.4%; 50–59 years = 46.0%; ≥60 years = 31.5%). A cumulative incidence plot was generated using the Kaplan-Meier method by plotting (1 − survival function) versus follow up time. Relative risks and 95% confidence intervals (CI) for the histological categories (modelled as indicator variables with normal as the reference) were estimated by Cox proportional hazards regression models using SAS PHREG (SAS Institute, Cary, North Carolina, USA) with adjustment for age (continuous variable), sex, smoking, alcohol use, 1983 cytology diagnosis, and treatment group (dichotomous variables).

RESULTS
Baseline characteristics of the analytic cohort have been described previously. There was little variation in age across the diagnostic categories but subjects with dysplasia or carcinoma in situ were more likely to be male, to smoke, to have a 1983 cytology diagnosis of dysplasia 2, and to be in the placebo treatment group than were subjects with a normal mucosa.

There were 114 new cases of OSCC diagnosed in the analytic cohort during the follow up period, including 52 cases during the intervention period (1987–1991), 48 cases during the next five years (1991–1996), and 14 cases during the last five years (1996–2001). There was a total cumulative incidence of 16.7% during the follow up period, and a crude incidence rate of 1510 cases per 100 000 person years of follow up.

Table 1 and fig 2 show the relationship between the 1987 oesophageal biopsy diagnoses and OSCC incidence during the 13.5 year follow up period. Eight per cent of patients with a normal mucosa or acanthosis and 6% of those with a worst biopsy diagnosis of oesophagitis developed OSCC in the follow up period. Fifteen per cent of patients with BCH developed OSCC. In contrast, 24% of those with mild dysplasia, 50% of those with moderate dysplasia, and 74% of those with severe dysplasia developed OSCC during this time. Seventy five per cent of patients whose initial biopsies showed CIS developed OSCC, the same proportion as those who started the follow up period with severe dysplasia. The cumulative incidence for patients originally classified as dysplasia NOS was 58%, consistent with this category being a

Table 1 Incidence and relative risk of oesophageal squamous cell carcinoma during 1987–2001, by initial histological diagnosis

<table>
<thead>
<tr>
<th>1987 diagnosis</th>
<th>No of subjects</th>
<th>No of OSCC cases</th>
<th>Cumulative OSCC incidence (%)</th>
<th>OSCC incidence rate*</th>
<th>Relative risk (95% CI) of OSCC incidence†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>375</td>
<td>31</td>
<td>8.3</td>
<td>605</td>
<td>1.0 (ref)</td>
</tr>
<tr>
<td>Acanthosis</td>
<td>77</td>
<td>6</td>
<td>7.8</td>
<td>597</td>
<td>0.9 (0.4–2.2)</td>
</tr>
<tr>
<td>BCH</td>
<td>40</td>
<td>6</td>
<td>15.0</td>
<td>1637</td>
<td>1.9 (0.8–4.5)</td>
</tr>
<tr>
<td>Oesophagitis</td>
<td>33</td>
<td>2</td>
<td>6.1</td>
<td>493</td>
<td>0.8 (0.2–3.2)</td>
</tr>
<tr>
<td>mD</td>
<td>76</td>
<td>18</td>
<td>23.7</td>
<td>1518</td>
<td>2.9 (1.6–5.2)</td>
</tr>
<tr>
<td>MD</td>
<td>30</td>
<td>15</td>
<td>50.0</td>
<td>10507</td>
<td>9.6 (5.3–18.3)</td>
</tr>
<tr>
<td>SD</td>
<td>23</td>
<td>17</td>
<td>73.9</td>
<td>21364</td>
<td>28.3 (15.3–52.3)</td>
</tr>
<tr>
<td>NOS</td>
<td>12</td>
<td>7</td>
<td>58.3</td>
<td>7689</td>
<td>12.7 (5.5–29.6)</td>
</tr>
<tr>
<td>CIS</td>
<td>16</td>
<td>12</td>
<td>75.0</td>
<td>21194</td>
<td>34.4 (16.4–71.4)</td>
</tr>
<tr>
<td>Total</td>
<td>682</td>
<td>114</td>
<td>16.7</td>
<td>1510</td>
<td>na</td>
</tr>
</tbody>
</table>

BCH, basal cell hyperplasia; mD, mild dysplasia; MD, moderate dysplasia; SD, severe dysplasia; NOS, dysplasia not otherwise specified; CIS, carcinoma in situ; OSCC, oesophageal squamous cell carcinoma.

*Oesophageal cancer incidence rates (per 100 000 person years) adjusted to the age distribution of the analytic cohort at baseline (<50 years = 22.4%; 50–59 years = 46.0%; ≥60 years = 31.5%).

†Relative risk (95% confidence interval (CI)) for OSCC adjusted for age, sex, tobacco use, alcohol use, 1983 cytology diagnosis, and treatment group.

Figure 2 Cumulative incidence of oesophageal squamous cell carcinoma by initial histological diagnosis. BCH, basal cell hyperplasia; Mild dys, mild dysplasia; Moderate dys, moderate dysplasia; Severe dys, severe dysplasia; Dysplasia NOS, dysplasia not otherwise specified; CIS, carcinoma in situ.
mix of all grades of dysplasia. The age adjusted incidence rates and multivariate adjusted relative risks for developing OSCC paralleled the cumulative incidence values. Relative to those with a normal mucosa, only subjects with initial biopsies showing squamous dysplasia or carcinoma in situ had a significantly elevated risk. Exclusion of cases diagnosed in the first year after endoscopy did not significantly change the relative risk estimates (BCH = 2.0, mild dysplasia = 2.9, moderate dysplasia = 9.6, severe dysplasia = 25.7, dysplasia NOS = 11.5, and carcinoma in situ = 34.5).

In addition to the 114 OSCC, there were 60 gastric adenocarcinomas (including 53 from the gastric cardia) and 13 other cancers (six liver, four lung, one breast, one colon, and one rectum) diagnosed in the analytic cohort during the follow up period. There were 266 deaths in the analytic cohort during this time, including 85 from OSCC, 37 from gastric cardia cancer, 15 from other cancers, 36 from heart disease, 60 from stroke, and 33 from other causes. Of the 30 patients with an initial diagnosis of moderate dysplasia, 15 developed OSCC, 10 died of other causes, and five were alive without evidence of cancer at the end of the follow up period. Of the 59 patients with initial severe dysplasia or CIS, 29 developed OSCC, four died of other causes, one was alive with gastric cardia cancer, and five were alive without evidence of cancer at the end of the follow up period.

DISCUSSION
Identification of clinically relevant precursor lesions is a prerequisite for designing successful prevention and early detection strategies for OSCC because such lesions are the targets for prevention and early detection efforts. Clinically relevant precursor lesions, which may be histological or molecular lesions, are abnormalities that precede invasive OSCC by months to years and are reliably associated with an increased risk of tumour development. Once documented, such lesions can be used for risk stratification of individuals, they can be gold standards for evaluating primary screening tests, and they can be targets for endoscopic therapy. They can also potentially be used as intermediate end points in prevention studies such as chemoprevention trials.

Relatively little has been written about the histological precursors of OSCC in low risk Western populations. Squamous dysplasia (including carcinoma in situ) is thought to be the relevant lesion because it is the accepted precursor in other organs with squamous epithelia, such as the cervix, and is commonly found adjacent to foci of invasive OSCC in oesophageal biopsy specimens.11 12

Most studies of OSCC precursor lesions have been performed in high risk Asian populations. Crespi et al and Munoz and colleagues13 15 described a high prevalence of histological oesophagitis, atrophy, and dysplasia in endoscopic surveys of Iranian and Chinese populations which have high OSCC rates and little or no evidence of oesophageal reflux. Based on differences in the prevalence of these lesions between these high risk populations and a low risk Chinese population and a one year follow up study of 20 patients, they proposed that oesophagitis, atrophy, and dysplasia may be precursor lesions of OSCC in these groups.14 16 Using similar methods, Yang and Qiu17 reported no difference in the prevalence of oesophagitis or the prevalence of atrophy in high versus low risk Chinese populations but found a higher prevalence of BCH and dysplasia in their high risk group. They also re-endoscoped 186 patients 30–78 months after their initial surveys and found progression to cancer in 34% of subjects whose initial biopsies showed both oesophagitis and dysplasia but similar progression to cancer in only 4% of those whose original biopsies showed oesophagitis alone, leading them to suggest that dysplasia is the primary precancerous lesion in the high risk areas of China.16 17 Others have since referred to Yang and Qiu’s comparative population findings to propose that BCH is also a precursor lesion of OSCC, preceding dysplasia in the carcinogenic sequence.18

The current study is the largest and longest prospective follow up of an endoscoped cohort looking for the development of OSCC. Our findings show no evidence that oesophagitis is associated with an increased risk of developing OSCC within 13.5 years after the initial diagnosis (relative risk 0.8 (95% CI 0.2–3.2)) and so we believe it is unlikely to be a significant precursor lesion in this population. Our initial endoscopic survey found no examples of atrophy16 so we cannot comment on this lesion. It is noteworthy however that Yang and Qiu also found this lesion to be rare in their surveys,16 17 making it unlikely to be an important precursor lesion.

We found a nearly twofold increased risk of OSCC associated with an initial diagnosis of BCH (relative risk 1.9 (95% CI 0.8–4.5)). Although this increased risk was not statistically significant, the cumulative incidence curve (fig 1) was very similar to that of mild dysplasia, and it is our experience that the morphological distinction between BCH and mild dysplasia can be difficult. Thus we think that patients with a diagnosis of BCH should be viewed clinically with some increased concern, even if only because of the risk of histological misclassification between this diagnosis and mild dysplasia.

In our analysis, all grades of squamous dysplasia and carcinoma in situ were associated with a significantly increased risk of developing OSCC, identifying these lesions as histological precursors of this tumour. In addition, we found that increasing grades of dysplasia were associated with dramatically increasing risk, implying real biological differences in these histological categories. Thus we believe that stratifying treatment based on dysplasia grade is appropriate, and that a shift in dysplasia grade over time (during passive follow up or during an intervention) may confer a real change in risk. The follow up experience of severe dysplasia and carcinoma in situ was equivalent in this study, suggesting that this diagnostic distinction is not clinically relevant and should be abandoned. This is not surprising, considering that common artefacts such as retention of surface epithelium in the biopsy forceps and imperfect biopsy orientation during tissue processing can significantly influence whether dysplastic cells appear to reach the upper third or the full thickness of the epithelium.4

Previous reports have questioned the utility of histological grading of oesophageal squamous intraepithelial neoplasia (IEN).19 In the current study, higher dysplasia grades identified distinctly higher cancer risks, showing that clinically useful histological grading of oesophageal squamous IEN is possible, using the extent of epithelial involvement by dysplastic cells as the basis for grading. Although occasional cases of invasive OSCC can be seen in which only the lower layers of the overlying epithelium contain dysplastic cells, the current findings demonstrate that over a series of cases, grading by extent of epithelial involvement correctly predicts different levels of OSCC risk. Accurate grading by this system requires good biopsy orientation, so the distribution of dysplastic cells from the base to the surface of the epithelium can be evaluated, but this can be reproducibly achieved by spreading the biopsies on supporting material in the endoscopy suite before fixation.20

Recent publications have advocated a unified histological classification for epithelial neoplasia throughout the gastrointestinal tract, including a two grade (low grade, high grade) classification of intraepithelial neoplasia.21 While the major findings of this study (significant OSCC risk associated with
Interventions aimed at preventing the development of OSCC during this time. Increasing grades of dysplasia were strongly associated with increasing risk, indicating that this precursor lesion can be divided morphologically into clinically meaningful grades. The follow up experience of severe dysplasia and CIS was equivalent, suggesting that this distinction is not clinically relevant. Documenting these precursor lesions of OSCC should assist in the development of effective prevention, early detection, and treatment strategies for this disease.

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**Authors’ affiliations**

G-Q Wang, X-D Sun, Y-L Qiao, Z-W Dong, Cancer Institute, Chinese Academy of Medical Sciences, Beijing, China.

C C Abnet, M J Roth, P R Taylor, S M Dawsey, Cancer Prevention Studies Branch, Center for Cancer Research, National Cancer Institute, Bethesda, Maryland, USA.

S D Mark, Biostatistics Branch, Division of Cancer Epidemiology and Genetics, National Cancer Institute, Bethesda, Maryland, USA.

Q Shen, Henan Medical University, Zhengzhou, China.

K J Lewin, Department of Pathology, UCLA Center for the Health Sciences, Los Angeles, California, USA.

Correspondence may also be addressed to Dr Y-L Qiao, Cancer Institute, Chinese Academy of Medical Sciences, 17 S Panjiayuan Lane, Chaoyang District, PO Box 2258, Beijing 100021, Peoples’ Republic of China; qiaoy@public.bta.net.cn

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**REFERENCES**


**EDITOR’S QUIZ: GI SNAPSHOT**

Black oesophagus—cause?

**Clinical presentation**
A 64 year old woman was admitted with coffee ground emesis. She had been bedridden for the previous two years due to osteoarthritis but had no past history of upper gastrointestinal ulcer, gastro-oesophageal reflux disease, or corrosive intake. Initial upper endoscopy revealed a patchy black appearance of the oesophageal mucosa extending from the proximal third of the oesophagus, with adherent yellow exudates, which became circumferential in the distal third of the oesophagus (fig 1). The black mucosa showed a sharp boundary at the gastro-oesophageal mucosal junction (fig 1), and the gastric and duodenal mucosa showed a normal appearance. The patient was treated with intravenous ranitidine for seven days and oral ranitidine thereafter. Endoscopy one week later revealed that the oesophageal mucosa was diffusely covered by thin whitish exudates with no stenosis, and biopsies revealed ulcerated mucosa with massive necrosis.

**Question**
What is the diagnosis?

*See page 227 for answer*

This case is submitted by:

A Sako, J Kitayama, S Kaizaki, H Nagawa
Department of Surgical Oncology, University of Tokyo, Tokyo, Japan

H Suzuki
Suzuki Hospital, Tokyo, Japan

**Correspondence to:** Dr A Sako, Department of Surgical Oncology, University of Tokyo, Hongo7-3-1, Bunkyo-ku, Tokyo 113-8655, Japan;
sakou-dis@h.u-tokyo.ac.jp
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Figure 1  Upper gastrointestinal endoscopy of the oesophagus revealing a diffusely black mucosa that ends sharply at the gastro-oesophageal mucosal junction.