The effect of metabolic risk factors on the natural course of gastro-oesophageal reflux disease

Y-C Lee,1,2 A M-F Yen,1 J J Tai,1 S-H Chang,1 J-T Lin,2,3 H-M Chiu,2 H-P Wang,2 M-S Wu,2 T H-H Chen1

ABSTRACT
Background and aims: The effect of metabolic risk factors on the natural course of gastro-oesophageal reflux disease (GORD), which remains elusive, was quantified.

Methods: The population included 3669 subjects undergoing repeated upper endoscopy. Data were analysed using a three-state Markov model to estimate transition rates (according to the Los Angeles classification) regarding the natural course of the disease. Individual risk score together with the kinetic curve was derived by identifying significant factors responsible for the net force between progression and regression.

Results: During three consecutive study periods, 12.2, 14.9 and 17.9% of subjects, respectively, progressed from non-erosive to erosive disease, whereas 42.5, 37.3 and 34.6%, respectively, regressed to the non-erosive stage. The annual transition rate from non-erosive to class A–B disease was 0.151 per person year (95% CI 0.136 to 0.165) and from class A–B to C–D was 0.079 per person year (95% CI 0.063 to 0.094). The regression rate from class A–B to non-erosive disease was 0.481 per person year (95% CI 0.425 to 0.536). Class C–D, however, appeared to be an absorbing state when not properly treated. Being male (relative risk (RR) 4.31; 95% CI 3.22 to 5.75), smoking (RR 1.20; 95% CI 1.03 to 1.39) or having metabolic syndrome (RR 1.75; 95% CI 1.29 to 2.38) independently increased the likelihood of progressing from a non-erosive to an erosive stage of disease and/or lowered the likelihood of disease regression. The short-term use of acid suppressants (RR 0.54; 95% CI 0.39 to 0.75) raised the likelihood of regression from erosive to non-erosive disease.

Conclusions: Intrathoracic reflux damage is a dynamic and migratory process in which the metabolic syndrome is associated with accelerated progression to or attenuated regression from erosive stages. These findings have important implications for the design of effective prevention and screening strategies.

Gastro-oesophageal reflux disease (GORD) is becoming increasing prevalent in the population, paralleling similar rises in the frequency of metabolic disorders, and resulting in the concomitant growth of an already considerable economic burden. Although GORD substantially affects public health, its natural history remains elusive. Two opposing theories have been proposed to explain the GORD heterogeneity. The category theory holds that GORD can be treated as three distinct entities (non-erosive (NE) reflux disease, erosive reflux disease and Barrett’s oesophagus11) and arises from the fact that therapeutic responses differ substantially between erosive and NE disease stages. The continuum theory suggests that GORD is a spectrum of diseases with differing severities. Support derives from the fact that transitions from NE to erosive disease are observed during endoscopic follow-up, and disease severity might account for observed variations in therapeutic responses.

Several studies have reported that the adverse effect of obesity on GORD is through mechanical alterations at the oesophago-gastric junction. However, since not every obese patient develops GORD, the pathogenesis must be multifactorial and cannot be explained by a single physiological parameter. Therefore, knowledge of GORD’s natural history and its relationship with metabolic risk factors is very informative not only to identify which individuals should undergo endoscopic screening but also to develop individually tailored prevention strategies. Unfortunately, it is difficult to assess GORD’s natural history because of the paucity of data from large, long-term endoscopic follow-up studies and the fact that symptoms of the disease cannot be treated as a surrogate measure for endoscopy. Even when data are available, serial observations with irregular inter-examination intervals render the quantification of transition between states and the derivation of kinetic curves that shows how each state evolves with time hard to assess without using complex multistate models. Our primary aim was to quantify the effect of putative factors, particularly the effect of metabolic risk factors, on the rates of onset, progression and regression between NE and erosive disease stages.

METHODS
Participants and evaluation
Our study was based on a voluntary health promotion programme at National Taiwan University Hospital (NTUH) that used a standard protocol including a physical examination, blood chemistries, plain radiography, abdominal ultrasound and endoscopy. Most subjects were invited to undergo an upper gastrointestinal endoscopy annually. Such a scheme is confirmed to be effective for cancer prevention in areas where the upper gastrointestinal cancers are prevalent. The ethics committee at our hospital approved the study protocol.

We enrolled patients who underwent at least two endoscopic examinations. Excluded were those who received proton pump inhibitors (PPIs) or histamine-2 receptor antagonists (H2RAs) in the 4 months preceding the first endoscopy, those who underwent gastrectomy and those with malignancy. National Health Insurance in Taiwan...
covers a 4-month course of treatment with a PPI or an H2RA for those who show signs of erosive oesophagitis or peptic ulcer disease after endoscopy. These medications are not available over-the-counter in Taiwan. Subjects who received less than 4 months of treatment were defined as short-term users. They were included in the analysis since a short-course PPI/H2RA treatment for erosive oesophagitis is the common clinical practice. Those who required two or more successive courses of PPI or H2RA treatment were defined as long-term users. They were excluded because such treatment may strongly affect GORD’s natural history.

Prior to the examination, a self-administered questionnaire was used to collect information on demographics, social habits and medical/medication histories. We defined symptoms of GORD as the presence of troublesome heartburn, acid regurgitation or both. Heartburn was defined as a burning sensation in the retrosternal area and acid regurgitation as the perception of flow of refluxed gastric contents into the mouth or hypopharynx. The frequency was once a week or more over the past 3 months. Self-reported data were confirmed in a face-to-face interview with an internist. This GORD-specific approach has been validated in population-scale research. Participants were evaluated for metabolic risk factors, including measurements of body mass index (BMI), waist circumference, blood pressure, plasma glucose, total cholesterol, high-density lipoprotein (HDL) cholesterol, triglycerides and uric acid levels. According to the modified criteria for Asians, metabolic syndrome is the presence of three or more of the following: waist circumference >90 cm for men or >80 cm for women, serum triglyceride levels ≥150 mg/dl, HDL cholesterol levels <40 mg/dl, blood pressure ≥150/85 mm Hg and serum glucose levels ≥110 mg/dl. Those receiving antidiabetic or antihypertensive therapy were assumed to have high fasting glucose levels or high blood pressure (details have been described elsewhere).

After the evaluation of metabolic risk factors, subjects underwent endoscopy. Erosive oesophagitis was scored using the Los Angeles (LA) classification system with standard comparator photos. The original LA classification for erosive oesophagitis consists of four classes but we combined these into two categories (classes A–B and C–D) to reduce interobserver variation and increase statistical power. The overall κ changed from 0.45 to 0.65, suggesting good consistency.

Three-state Markov model
We modelled GORD’s natural history as a three-state Markov process by defining state 1 as NE disease, state 2 as LA class A–B oesophagitis and state 3 as class C–D oesophagitis (fig 1). Transitions between serial states were quantified by two instantaneous progression rates from state 1 to 2 (λ1) and from state 2 to 3 (λ3), and two instantaneous regression rates from state 2 to 1 (λ2) and state 3 to 2 (λ4). Cumulative risk for each transition was computed by transition probabilities that were a function of transition rates λ1–λ4 and follow-up time by using the method of Chen et al.

The evolution of these cumulative risk curves corresponding to state-to-state transitions of GORD’s natural course with follow-up time are called kinetic curves. For subjects free of erosive disease at baseline, kinetic curves showed the evolution of cumulative risk for developing class A–B and class C–D oesophagitis and probability of remaining in the NE disease state. For subjects with class A–B disease at baseline, kinetic curves showed the evolution of cumulative risk for developing class C–D, remaining in class A–B and regressing to the NE disease state.

Statistical analysis
Time intervals between endoscopic examinations were recorded to build up a continuous-time Markov process for the three-state model. We estimated the transition rates labelled in fig 1 and their 95% CIs based on the total likelihood, a product of transition probabilities from a series of endoscopic examinations in all subjects. Statistical analyses for this model have been described in several papers. As time intervals between assessments are irregular and vary from individual to individual, a multistate model is adopted to tackle this technical problem with the incorporation of different time intervals into transition probabilities, which is elaborated in Appendix 1.

We presented the model parameters derived from the complete data set of subjects. To test the predictive validity of the current model, we also performed cross-validation by splitting data into 2/3 for deriving the model and 1/3 for validation of the model. The observed transition histories in the validated data set were compared with the predicted ones that were computed by the application of parameters trained from the derived data set.

As we are interested in the effect of metabolic risk factors on transition rates, a univariate regression analysis was therefore done to assess the effect of each component on transition rates. The exponential Markov regression form was adapted to estimate relative risk (RR), which is done by taking exponentials of the regression coefficients of the Markov regression. Besides the metabolic profile, factors considered in the regression included basic demographic information, lifestyle factors, symptoms of reflux and the use of acid suppressants following screening. It should be noted that each predictor in the same individual may vary from time to time and was repeatedly evaluated along with each endoscopic screening. They are treated as time-varying covariates, which means that their contributions to progression and regression of GORD may depend on the status they had at the time preceding the next transition during a given epoch. Thus, the net force of dynamic change of each covariate contributing to progression and regression is worthy of investigation.

Take smoking status (smoking, X = 1; no smoking, X = 0), for example. An individual at time t0 was a current smoker (X = 1), he developed A–B during time interval (t1–t0), quit smoking, was treated as non-smoking (X = 0), at time t1, regressed to NE during the time interval (t2–t1), and stayed as NE without smoking until time t3. Thus, as smoking status changed with time, its effect in each epoch makes different contributions to disease progression and regression in the same individual. The net force of smoking on the state-to-state transitions can be considered in this manner to aggregate each individual change into a population-average net effect expressed by the difference of regression coefficients between progression and regression.

Figure 1 Three-state Markov model of the natural history of gastro-oesophageal reflux disease. The transition rates, λ1–λ4, are parameters in the model and will be estimated.

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The phenomenon may be applied to PPI/H2RA use that is associated with progression on the grounds of indication and regression probably causally related through treatment.

Hypothesis testing for such net force for each risk factor mentioned above is performed as follows. The transition rate function is first developed:

$$
\lambda_j(t) = \lambda_{0j} \exp(\beta_j X(t)), \quad j = 1, 2
$$

where $$\lambda_{0j}$$ and $$\lambda_{02}$$ are baseline progression and regression rates, $$\beta_1$$ and $$\beta_2$$ are the corresponding regression coefficients, and $$X(t)$$ is the indicator variable concerned with the presence of risk factors at time $$t$$ prior to the next transition. The null hypothesis for an attributed lack of effect due to a risk factor is defined as $$\beta_1 - \beta_2 = 0$$. The alternative hypothesis was set as $$\beta_1 - \beta_2 > 0$$, where $$\beta_1 - \beta_2 > 0$$ indicates a detrimental effect and $$\beta_1 - \beta_2 < 0$$ indicates a protection. By using the estimated variance–covariance matrix, the significance of a risk factor was determined using the Wald test statistic.

To build a multivariate model, we used forward selection to evaluate the additive effects of risk factors. The presence of metabolic syndrome, and its individual components, were added one by one into the model. The final model was selected based on the log-likelihood ratio test. Using the set of risk factors that were significant in the final model, we created a predicted risk score based on regression coefficients estimated from the model and the assigned time-invariant and time-varying factors. Following the above notation of $$\beta_1$$ and $$\beta_2$$, the predicted risk score at time $$t$$ ($$S(t)$$) for each set of risk factors is expressed by:

$$
S(t) = \sum_{j=1}^{q} (\beta_{1j} - \beta_{2j}) X_j(t)
$$

where $$X_j(t)$$ is a set of significant factors and $$q$$ is the number of significant factors. Risk stratification according to this predicted risk score yielded a series of different kinetic curves with different shapes, showing heterogeneous natural courses of disease. Analysis was done using an SAS/IML procedure (version 9.1; SAS Institute Inc., Cary, North Carolina, USA). All p values were two sided; $$p<0.05$$ indicated significance.

**RESULTS**

**Descriptive findings**

Between June 2003 and December 2006, 19 812 subjects underwent screening upper endoscopy at NTUH. Of these, 3669 had at least two examinations and comprised our study group (table 1). Mean age (SD) was 56.3 (10.5) years; mean BMI was 24.1 (5) kg/m². GORD symptoms were reported in 11.3%. The prevalence rate (16.4%) of erosive oesophagitis among these 3669 subjects was similar to that (15.7%) obtained from the whole population (n = 19 812) in the previous study. This may suggest that we did not have findings on the first endoscopy that warranted more frequent surveillance for these subjects with repeated endoscopy. Stratified by their endoscopic findings, 10.5% of NE, 15.5% of A–B and 35.5% of C–D groups had GORD symptoms. The average duration of short-term PPI or H2RA treatment was 2.1 (0.8) months. Thirty patients (0.15%) were excluded for being long-term PPI or H2RA users. Barrett’s oesophagus was too rare (0.06%) to be included. As only 0.55% had chronic renal insufficiency (serum creatinine concentration $$\geq$$ 2 mg/dl), this was also excluded.

**Transition rates between states in GORD’s natural history**

Subjects underwent up to four endoscopies creating three epochs (baseline endoscopy to endoscopy 2, endoscopy 2 to 3, and endoscopy 3 to 4). The mean duration of epochs 1, 2 and 3 (in days) was 528 (210), 392 (108) and 352 (60). Table 2 shows aggregate numbers of observed transitions between states.

There were 5145 transitions, including 3669 in epoch 1, 1401 in epoch 2, and 536 in epoch 3. Observed rates of transition from NE to erosive oesophagitis were 12.2% (95% CI 8.9% to 15.5%), 14.9% (95% CI 9% to 20.8%) and 17.9% (95% CI 13.2% to 22.6%). The risk of progressing from class A–B to class C–D

**Table 1 Baseline characteristics of the 3669 study subjects**

| Characteristic | No. of subjects (%) |
|---------------|---------------------|
| Male          | 2483 (67.7)         |
| Smoker        | 416 (11.3)          |
| Drinks alcohol (at least once per week) | 2219 (60.5) |
| Chronic illnesses |               |
| Cardiac       | 47 (1.3)            |
| Pulmonary     | 92 (2.6)            |
| Hepatic       | 501 (13.7)          |
| Peptic ulcer  | 648 (17.7)          |
| Cholesterol  $$>$$ 200 mg/dl | 314 (8.6) |
| Hyperuricaemia* and/or history of gout | 283 (7.7) |
| Metabolic syndrome | 498 (13.6) |
| Enlarged waist circumference | 1239 (33.8) |
| Hypertension or blood pressure $$>$$130/85 mm Hg | 559 (15.2) |
| Diabetes or fasting glucose $$>$$110 mg/dl | 198 (5.4) |
| HDL-C $$<$$ 40 mg/dl | 1678 (45.7) |
| Triglycerides $$>$$ 150 mg/dl | 908 (24.7) |
| Exercise, number of times per week |         |
| $$>$$ 5       | 1013 (27.6)         |
| 3–4           | 1585 (43.2)         |
| $$<$$ 2        | 1071 (29.2)         |
| Sleep quality |                    |
| Good          | 1600 (43.6)         |
| Fair          | 1564 (42.6)         |
| Poor          | 505 (13.8)          |
| Symptoms of GORD | 413 (11.3) |
| Short-term use of PPI or H2RA | 587 (11.4) * |

*Serum uric acid concentration $$>$$ 7.5 mg/dl.
†From 5145 transition periods.

**Table 2 Aggregate counts of transitions between states during the three study epochs**

| Epochs | No. of subjects (%) |
|--------|---------------------|
|        | NE                  | Class A–B | Class C–D |
|        | (n = 3066)          | (n = 586) | (n = 17)  |
| Baseline | Endoscopy 2         | 2985 (8.8) | 350 (11.4) | 23 (8.8) |
|         | (n = 3066)          |           | 249 (42.5) | 304 (51.3) | 33 (5.6) |
| Class A–B (n = 586) | 0 (0) | 0 (0) | 17 (100) |
| Class C–D (n = 17) | 0 (0) | 0 (0) | 12 (100) |
| Endoscopy 2 | Endoscopy 3        | 791 (85.1) | 136 (14.6) | 3 (0.3) |
| NE (n = 930) | 74 (37.3) | 109 (55.1) | 15 (7.6) |
| Class A–B (n = 198) | 0 (0) | 0 (0) | 12 (100) |
| Class C–D (n = 12) | 0 (0) | 0 (0) | 12 (100) |
| Endoscopy 3 | Endoscopy 4        | 207 (82.1) | 45 (17.9) | 0 (0) |
| NE (n = 252) | 27 (34.6) | 39 (50) | 12 (15.4) |
| Class A–B (n = 78) | 0 (0) | 0 (0) | 6 (100) |
| Class C–D (n = 6) | 0 (0) | 0 (0) | 6 (100) |

NE, non-erosive state.
oesophagitis increased from 5.6% (95% CI 3.7% to 7.5%) in epoch 1 to 15.4% (95% CI 7.4% to 23.4%) in epoch 3; the probability of regression from class A–B to NE decreased from 42.5% (95% CI 38.5% to 46.5%) in epoch 1 to 34.6% (95% CI 24.8% to 44.4%) in epoch 3. However, no statistically significant increase in progression or decrease in regression across epochs was noted (p>0.05). Because no class C–D subjects showed regression to class A–B or NE, the annual regression rate from C–D to A–B (ie, \(l_3\)) was set to zero.

Annual progression rates from NE to A–B (ie, \(l_1\)) and from A–B to C–D (ie, \(l_3\)) were 0.151 (95% CI 0.156 to 0.165) and 0.079 (95% CI 0.063 to 0.094) per person year, respectively, and the regression rate from A–B to NE (ie, \(l_2\)) was 0.481 (95% CI 0.425 to 0.556) per person year. The corresponding figures were 0.159 (95% CI 0.126 to 0.152), 0.084 (95% CI 0.065 to 0.101) and 0.346 (95% CI 0.294 to 0.398) when we excluded the transition histories of being administered with short-term PPI or H2RA treatment preceding the next transition.

**Kinetic curves**

Model fitting was assessed by comparing predicted with observed transitions using \(x^2\); the lack of a significant difference indicated a good fit for the model (p = 0.415). The observed transition histories were still compatible with the predicted values using the cross-validation method (p = 0.876). Figure 2A shows the cumulative risk of developing class A–B (middle curve) and class C–D oesophagitis (bottom curve) and the probability of staying in the NE disease state (upper curve) for subjects free of erosive disease at baseline. After a 10-year follow-up, 68% of patients have NE disease, 19% class A–B, and 13% class C–D. Since disease regression is common, only 8% of patients would stay in the NE state throughout the follow-up period without any transition histories. Figure 2B shows the cumulative risk of progressing to C–D, of remaining in class A–B and of regressing to NE for subjects with A–B at baseline, respectively. Of these, 60% would undergo regression to NE within 10 years, 17% would remain as A–B, and 23% would progress to C–D.

**Effects of risk factors on GORD’s natural course**

**Univariate analysis**

Table 3 shows the results when we introduced covariates one by one into our model. We only evaluated effects on transitions between NE and A–B because there were too few transitions from A–B to C–D to allow stable parameter estimation.

Being male (RR = 2.36) and having a BMI \(\geq 27\) kg/m\(^2\) (RR = 1.23) both raised the likelihood of progressing from NE to erosive disease and lowered the likelihood of regression from erosive to NE disease (RR = 0.55 and 0.70). Smokers and heavy drinkers had a significant risk of erosive disease (RR = 2.27 and 1.52). Subjects with metabolic risk factors, including hypercholesterolaemia, hyperuricaemia, enlarged waist circumference, hypertension, low HDL cholesterol level, hypertriglycaemia and metabolic syndrome, were more likely to progress from NE to erosive disease and/or less likely to regress from erosive to NE states. Short-term PPI or H2RA use increased the likelihood of regression from erosive to NE states (RR = 2.85). GORD symptoms increased the risk of erosive disease (net RR = 1.64).

**Multivariate analysis**

We used forward selection to evaluate the additive effects of covariates on disease onset and regression. The initial model included only gender (the most significant factor in univariate analysis). We then added variables until they stopped adding significantly to the model. The final model included gender, smoking, metabolic syndrome and short-term PPI or H2RA usage (see table 4 for resulting risk estimates).

The clinical weight each risk factor contributes to (the net effect of regression coefficients) was 1.46 (natural logarithm of 4.31) for male gender, 0.18 for smoking, 0.56 for metabolic syndrome and 0.61 for short-term PPI or H2RA. The predicted risk score at time \(t\) based on the clinical weight together with risk factor was:

\[
\text{Risk score} = (1.46 \times \text{male}) + (0.18 \times \text{smoking}) + (0.56 \times \text{metabolic syndrome}) - (0.61 \times \text{short-term use of PPI or H2RA})
\]
### Table 3: Relative risk of transition and corresponding 95% CIs by factors from the univariate three-state Markov model

| Variables*   | RR (95% CI)† |
|--------------|--------------|
|              | NE→class A–B | Class A–B→NE | Net effect          |
| Age ≥65 years| 1.19 (0.95 to 1.48) | 1.02 (0.78 to 1.34) | 1.17 (0.91 to 1.50) |
| Male         | 2.36 (1.79 to 3.13)‡ | 0.55 (0.40 to 0.74)‡ | 4.33 (3.30 to 5.66)‡ |
| Body mass index ≥27 kg/m² | 1.28 (1.01 to 1.65)‡ | 0.70 (0.52 to 0.96)‡ | 1.81 (1.36 to 2.41)‡ |
| Smoker       | 2.27 (1.68 to 3.06)‡ | 1.38 (0.96 to 1.97) | 1.65 (1.24 to 2.18)‡ |
| Alcohol use  | 1.32 (1.08 to 1.62)‡ | 0.99 (0.78 to 1.26) | 1.34 (1.07 to 1.69)‡ |

*Factors were dichotomised (no/yes) as follows: age ≥65 years, male, body mass index ≥27 kg/m², smoker, alcohol consumed ≥ once per week, metabolic syndrome, exercise more than twice per week, poor sleep quality, symptoms of GORD and use of short-term PPI or H2RA. The “no” group constitutes the baseline comparator.

†The RR for evaluating the role of each factor was arrived at by taking the exponential of the regression coefficient (β) of the Markov regression—that is, exp(β) for progression, exp(β1–β2) for regression and exp(β1−β2) for the net effect.

‡p < 0.05.

GORD, gastro-oesophageal reflux disease; H2RA, histamine-2 receptor antagonist; PPI, proton pump inhibitor.

These dichotomous variables were coded as described in table 4.

Kinetic curves can be stratified by classifying predicted risk score into four categories, as shown in fig 3A–D. A female with reflux, for example, who does not smoke or have metabolic syndrome but received short-term PPI treatment (total risk score of 0.56) would have the lowest probability of progressing to erosive disease (fig 3A, low risk group). An untreated male smoker with metabolic syndrome (total risk score of 2.2) would have the highest probability of progressing from NE to erosive disease (fig 3D, high risk group). The higher the risk score, the higher the probability of developing erosive oesophagitis.

### DISCUSSION

We quantified the natural history of GORD by fitting a large longitudinal follow-up database of patients undergoing endoscopy. The step-by-step transitions are a solid demonstration of GORD’s dynamic nature. The predicted risk score may enable clinicians to develop individually tailored preventive strategies.

In addition to cross-validation, several studies support our model’s credibility on external predictive validity. Among patients with NE reflux disease, 53% (15%) developed erosive changes within 6 months; 16 of 18 (89%) became erosive 10 years later. A large database of 2306 GORD patients showed that oesophageal mucosa over 7.6 months was unchanged in 67% of patients (our model predicted 67%), improved in 21% (predicted, 27%) and worsened in 11% (predicted, 6%). Among 3894 GORD patients followed for 2 years, 25% with NE reflux disease progressed to A–B (predicted, 18%) and 0.6% progressed to C–D (predicted, 1.5%); 57% of those with A–B remained A–B (predicted, 39%) and 61% regressed to NE (predicted, 51%). Five years later, 72% of those with NE reflux disease at baseline...
remained NE (predicted, 74%), 22% progressed to A–B (predicted, 20%) and 6% progressed to C–D or Barrett’s oesophagus (predicted, 6%). For patients with class A–B at baseline, 57% regressed to NE (predicted, 63%), 31% stayed A–B (predicted, 21%) and 12% became C–D disease or Barrett’s oesophagus (predicted, 16%).

A plausible link can be established between category and continuum theories. The most significant factor affecting vulnerability to erosive oesophagitis is gender. Hence, a slim female who does not smoke or drink alcohol may remain in the NE state for a long time with little chance of developing erosive disease. An obese male, in contrast, who smokes, drinks heavily and has metabolic syndrome (again a typical picture) would probably progress to erosive disease. The probability of changes in disease status being detected at endoscopy would also increase. Thus, different combinations of risk factors lead to different severities of intraoesophageal damage and the disease appears as a continuum upon endoscopic inspection.

The pathogenesis of reflux symptoms is complicated and cannot be explained solely by intraoesophageal damage. Enhanced peripheral and central neural perceptions of stimuli may be crucial in symptom generation. As the above illustrates, demographic and endoscopic findings can vary markedly in symptomatic patients and may explain why GORD seems categorical under a symptom-oriented approach. Treating cases of reflux as categorical entities according to the mechanisms of symptom generation is worthwhile; however, the value of identifying risk factors and protecting the oesophageal mucosa from irreversible damage cannot be overemphasised. Since spontaneous regression is still possible in patients with low-grade erosive disease without pharmacological treatment, evaluation of individual risk at this stage would give patients opportunities to modify their behaviour (weight reduction, giving up smoking) and enable clinicians to select patients most likely to develop irreversible changes for endoscopic screening and offer them early pharmacological treatment. This argument has been supported by our findings.

Figure 3  The kinetic curves (cumulative probabilities) of transition from non-eroseive to class A–B or class C–D disease, and of remaining in the non-erosive disease state, stratified by individual risk score. A to D illustrate, respectively, the cumulative probability of transition between the above states for those with a risk score of <0, of between 0 and 1, of between 1 and 2, and of >2.
After removing the effect of short-term treatment, the regression rate has been attenuated from 0.481 to 0.346. This suggests the short-term treatment effect accounts for 27% (1–0.35/0.48) and other significant risk factors are responsible for 73%. The possibility of disease regression, making allowance for short-term treatment effect, remained and may be related to the modification of other risk factors.

Obesity significantly increased the risk of GORD symptoms, erosive oesophagitis, Barrett’s oesophagus and oesophageal adenocarcinoma. However, obesity cannot be the sole reason. Hypertension was found, after adjusting for BMI, to be associated with erosive oesophagitis. In a Japanese population, male sex, obesity, hyperglycaemia and hypertension were independent risk factors for erosive oesophagitis. Investigating a database of a Korean population, the presence of metabolic syndrome and a higher visceral adipose tissue area were risk factors. Consistent with the above, we found an association between several metabolic risk factors and accelerated progression to or attenuated regression from erosive disease, which suggests a potential benefit of treating metabolic disorders in GORD patients. The recommendation to abstain from smoking and alcohol is justified.

Our results are credible for several reasons. First, we had numerous cases of NE and were able to assess progression to erosive disease. The simultaneous evaluation of symptomatic and asymptomatic subjects also enabled us to observe the entire disease spectrum. Secondly, all our endoscopists completed the same training programme using a standardised rating protocol. This substantially reduced heterogeneity amongst observers and strengthened our ability to model natural history. Thirdly, we found no spontaneous regressions from high-grade erosive states, which conflicts with findings that 42% of patients with class C–D disease regresses to A–B and 50% to NE disease within 2 years. These latter results were obtained from symptomatic patients with GORD who were participating in a therapeutic trial and are potentially confounded by pharmacological effects. The fact that most of our participants remained asymptomatic, even in high-grade erosive states, lowered their incentive to seek treatment and allowed us to assess the unInterrupted natural history of the disease.

Because progression of GORD is orderly, the Markov approach was appropriate for modelling. However, our target group tended to reflect the general population, so there were few transitions from low- to high-grade oesophagitis and we were unable to investigate the effects of covariates on this stage. Secondly, information about H pylori infection was not available. For subjects with antrum-predominant gastritis, H pylori may increase gastric acid secretion and thus increase the risk of GORD. This is supported by our finding that peptic ulcer disease (mostly duodenal ulcer) at baseline was associated with a higher risk of progression to an erosive state. However, the effect of H pylori eradication on GORD should be investigated. Finally, a short course of acid-suppressing treatment is common among patients with minor erosive disease, and considering this factor in the Markov model may improve its ability to predict disease progression. However, the exclusion of long-term users of PPIs or H2RAs may limit the generalisability of the model to patients with earlier onset disease or more severe disease. The evaluation of a large and longitudinal database is warranted in order to update our parameter estimates and extend our model to include patients who require long-term treatment with acid-suppressing medication and who are diagnosed with Barrett’s oesophagus.

Our findings suggest that intraoesophageal damage develops as a dynamic process over time. Risk factors in susceptible individuals modulate the likelihood of state-to-state transitions, resulting (upon endoscopy) in the appearance of a continuous spectrum of disease. GORD can therefore be staged with respect to the extent of progression, as with many other chronic diseases. The translation of the quantified knowledge of GORD’s natural history into predicted risk score together with the kinetic curve will be vital for developing individually tailored prevention and screening programmes, to identify candidates for potential interventions, and to determine optimal timing of proposed interventions.

Competing interests: None.

Ethics approval: The ethics committee at the National Taiwan University Hospital approved the study protocol.

REFERENCES

1. Sandler RS, Everhart JE, Donowitz M, et al. The burden of selected digestive diseases in the United States. Gastroenterology 2002;122:1500–11.
2. Fass R. Distinct phenotypic presentations of gastroesophageal reflux disease: a new view of the natural history. Dig Dis Sci 2004;52:100–7.
3. Fass R, Olufman J.J. Gastroesophageal reflux disease—should we adopt a new conceptual framework? Am J Gastroenterol 2002;97:1901–9.
4. Labenz J, Nocon M, Lind T, et al. Prospective follow-up data from the ProGERD study suggest that GERD is not a categorical disease. Am J Gastroenterol 2006;101:2457–62.
5. Pace F, Bianchi Paro G. Gastroesophageal reflux disease: a typical spectrum disease (a new conceptual framework). Am J Gastroenterol 2004;99:946–8.
6. Hampil H, Abraham NS, EI-Senig HB. Meta-analysis: obesity and the risk for gastroesophageal reflux disease and its complications. Ann Intern Med 2005;143:199–211.
7. Corley DA, Kubo A, Levin TR, et al. Abdominal obesity and body mass index as risk factors for Barrett’s esophagus. Gastroenterology 2007;133:34–41.
8. Wu JC, Mui LM, Cheung CM, et al. Obesity is associated with increased transient lower oesophageal sphincter relaxation. Gastroenterology 2007;132:883–9.
9. Jacobson BC, Somers SC, Fuchs CS, et al. Body-mass index and symptoms of gastroesophageal reflux in women. N Engl J Med 2006;354:2440–8.
10. de Vries DR, van Henwaarden MA, Smout AJ, et al. Gastroesophageal pressure gradients in gastroesophageal reflux disease: relations with histal hernia, body mass index, and esophageal acid exposure. Am J Gastroenterol 2008;103:1349–54.
11. Pandolfino JE. The relationship between obesity and GERD: “big or overweight?”. Am J Gastroenterol 2008;103:1355–7.
12. Taiwan Cancer Registry. Age-specific incidence rates of major cancers in Taiwan, Department of Health, Executive Yuan: Taiwan; 2005. Available at: http://crs.cph.nthu.edu.tw/. Accessed 10 Jun 2008.
13. Tsubono Y, Hisamichi S. Screening for gastric cancer in Japan. Gastric Cancer 2000;3:9–18.
14. Lee YC, Lin JT, Wu HM, et al. Cost-effectiveness analysis between primary and secondary preventive strategies for gastric cancer. Cancer Epidemiol Biomarkers Prev 2007;16:875–85.
15. Bureau of National Health Insurance. Laws & Regulations. Bureau of National Health Insurance, Taiwan. Available at: http://www.nhi.gov.tw/. Accessed 7 Jan 2008.
16. Shu BS, Cheng HC, Chang WL, et al. The impact of body mass index on the application of on-demand therapy for Los Angeles grades A and B reflux esophagitis. Am J Gastroenterol 2007;102:2387–94.
17. Lee YC, Wang HP, Chiou HM, et al. Comparative analysis between psychological and endoscopic profiles in patients with gastroesophageal reflux disease: a prospective study based on screening endoscopy. J Gastroenterol Hepatol 2006;21:798–804.
18. Locke GR, Talley NJ, Weaver AL, et al. A new questionnaire for gastroesophageal reflux disease. Mayo Clin Proc 1994;69:539–47.
19. Tan CE, Mia S, Wai D, et al. Can we apply the National Cholesterol Education Program Adult Treatment Panel definition of the metabolic syndrome to Asians? Diabetes Care 2004;27:1182–6.
20. Chiu HM, Lin JT, Shun CT, et al. Association of metabolic syndrome with proximal and synchronous colorectal neoplasms. Clin Gastroenterol Hepatol 2007;5:221–9.
21. Dent J, Brun J, Fendrick AM, et al. on behalf of the Genval Workshop Group. An evidence-based appraisal of reflux disease management—the Genval Workshop Report. Gut 1999;44(Suppl 2):S1–16.
22. Kuo MC, Lin LT, Yang TC, et al. Interobserver and intraobserver variation in endoscopic assessment of GERD using the “Los Angeles” classification. Gastrointest Endosc 1999;49:700–4.
23. Lee YC, Lin JT, Chiou HM, et al. Interobserver and intraobserver consistency for grading esophagitis with narrow-band imaging. Gastrointest Endosc 2007;66:230–6.
24. Chen TH, Kuo HS, Yen MF, et al. Estimation of sojourn time in chronic disease screening without data on interval cases. Biometrics 2000;56:167–72.
25. Cox DR, Miller HD. The theory of stochastic processes. London: Methuen, 1965.
26. Kalbfleisch JD, Lawless JF. The analysis of panel data under a Markov assumption. J Am Stat Assoc 1985;80:863–71.
27. Duffy SW, Chen HH, Tabar L, et al. Estimation of mean sojourn time in breast cancer screening using a Markov chain model of both entry to and exit from the preclinical detectable phase. Stat Med 1995;14:1531–43.
28. Wu HM, Yen MF, Chen THH. SAS macro program for non-homogeneous Markov process in modeling multi-state disease progression. Comput Methods Programs Biomed 2004;75:95–105.
29. Yen AMF, Chen THH. Mixture multi-state Markov regression model. J Appl Stat 2007;34:11–21.
30. Tseng PH, Lee YC, Chiu HM, et al. Prevalence and clinical characteristics of Barrett’s esophagus in a Chinese general population. J Clin Gastroenterol 2008;42:1074–9.
31. Pace F, Santalucia F, Bianchi Porro G. Natural history of gastro-oesophageal reflux disease without oesophagitis. Gut 1991;32:945–8.
32. Pace F, Bollanti S, Molteni P, et al. Natural history of gastro-oesophageal reflux disease without oesophagitis (NERD)—a reappraisal 10 years on. Dig Liver Dis 2004;36:111–5.
33. Sontag SJ, Sonnenberg A, Schnell TG, et al. The long-term natural history of gastroesophageal reflux disease. J Clin Gastroenterol 2006;40:396–404.
34. Malfertheiner P, Nocon M, Hicker MR, et al. How does GERD evolve during a 5-year follow-up period? The ProGERD study [abstract]. Gastroenterology 2008;134:A592.
35. Fass R, Tougas G. Functional heartburn: the stimulus, the pain, and the brain. Gut 2002;51:885–92.
36. Gudlaugsdottir S, Verschuren W, Dees J, et al. Hypertension is frequently present in patients with reflux esophagitis or Barrett’s esophagus but not in those with non-ulcer dyspepsia. Eur J Intern Med 2002;13:369–75.
37. Moki F, Kusano M, Mizuide M, et al. Association between reflux oesophagitis and features of the metabolic syndrome in Japan. Aliment Pharmacol Ther 2007;26:1069–75.
38. Chung SJ, Kim D, Park MJ, et al. Metabolic syndrome and visceral obesity as risk factors for reflux oesophagitis: a cross-sectional case–control study of 7078 health check-up Koreans. Gut 2008;57:1360–5.
39. El-Omar EM, Dinken E, El-Nujumi A, et al. Helicobacter pylori infection and chronic gastric acid hyposecretion. Gastroenterology 1997;113:15–24.
40. Raghunath A, Hungin AP, Woolf D, et al. Prevalence of Helicobacter pylori in patients with gastro-oesophageal reflux disease: systematic review. BJM 2003;326:737–43.

APPENDIX 1

Model specification and the likelihood function for data on endoscopic examination

The transition rates in our three-state model can be expressed in an intensity matrix as follows:

\[
\begin{pmatrix}
 \lambda_1 & 0 & \lambda_2 \\
 0 & \lambda_3 & 0 \\
 \lambda_4 & 0 & \lambda_4
\end{pmatrix}
\]

where states 1 and 2 represent states NE, LA class A–B and LA class C–D, respectively. The four transition parameters of \(\lambda_1, \lambda_2, \lambda_3, \lambda_4\) have been defined in the text and fig 1. Using the Kolmogorov equation, the corresponding transition probabilities can be expressed as follows:

\[
\begin{align*}
\text{Current state} & \quad 1 & 2 & 3 \\
\text{Previous state} & \quad 1 & -\lambda_1 & \lambda_1 & 0 \\
& \quad 2 & 0 & -\lambda_2 & \lambda_2 \\
& \quad 3 & 0 & 0 & -\lambda_4 & \lambda_4
\end{align*}
\]

where \(P_{ij}(t) = Pr\{X(t) = j | X(t_0) = i\} \) is a function of \(\lambda_1, \lambda_2, \lambda_3, \lambda_4\) and fi. Each individual has his/her own history of endoscopic examination. The likelihood function is established based on the transition probabilities with the use of the data on the history of examinations, in order to estimate the transition parameters \(\lambda_1, \lambda_2, \lambda_3, \lambda_4\).

APPENDIX 2

Kinetic curves based on the estimated transition rates

Given the estimated annual progression rates from NE to class A–B oesophagitis and from A–B to class C–D oesophagitis of 0.151 \((\lambda_1)\) and 0.079 \((\lambda_2)\) per person year, respectively, and the regression rate from class A–B to NE and from C–D to class A–B of 0.481 \((\lambda_3)/\) per person year and 0 \((\lambda_4)\), respectively, the transition rate matrix can be expressed as follows:

\[
\begin{pmatrix}
 \lambda_1 & 0 & 0 \\
 0 & \lambda_2 & 0 \\
 0 & 0 & \lambda_3 \\
 0 & 0 & \lambda_4
\end{pmatrix}
\]

The cumulative risk (probability) over time can be calculated using an SAS/IML procedure. For example, the first year cumulative risk is demonstrated as follows:

\[
\begin{pmatrix}
 1 & 0 \\
 2 & 0.481 \\
 3 & 0
\end{pmatrix}
\]

where the \(\nu\) denotes the number of individuals who are in state \(i\) at time \(t\) and in state \(j\) at time \(t\). The maximum likelihood estimates of the transition rates can be obtained using the Newton–Raphson procedure.