Best practice in symptom assessment: a review

E McColl

Assessment of symptoms is important both in the diagnosis of gastro-oesophageal reflux disease (GORD), and in monitoring response to therapeutic interventions. Key quality aims in assessing symptoms are to gather data that are valid, reliable, unbiased, discriminating, and responsive to change. Important considerations include who should assess symptoms, method of data collection, timing of assessment, and methods of assessing multiple symptoms.

SUMMARY
Assessment of symptoms is important both in the diagnosis of gastro-oesophageal reflux disease (GORD) and in monitoring response to therapeutic interventions. Key quality aims in assessing symptoms are to gather data that are valid, reliable, unbiased, discriminating, and responsive to change. Important considerations are: who should assess symptoms; method of data collection; timing of assessment; and methods of assessing multiple symptoms. A review of the substantial literature on these topics has led to the following conclusions. (1) Due to the subjective nature of symptoms, patient self report is more appropriate than clinician assessment. (2) Symptom diaries are generally considered to be the “gold standard” but well designed questionnaires with an appropriate recall period may be adequate. (3) Careful attention to instrument design and data collection protocol can enhance completion rates. (4) Use of a clearly defined reference period can improve the quality of response; the choice of reference period must take into account anticipated fluctuations over time in the symptoms of interest and how memorable those symptoms are likely to be—reference periods of one week to one month are most appropriate. (5) The duration of data collection must represent an appropriate compromise between capturing anticipated day to day variation and imposing excessive respondent burden. (6) There is no single optimal strategy with respect to collecting data on multiple symptoms. Decisions should be based on the objectives and analytic strategies of individual studies, on judgements about likely covariation in symptoms, and on whether respondents themselves can make composite ratings or whether the assumptions implicit in summated ratings are justified.

INTRODUCTION
Symptoms have been defined as “subjective experience of abnormal function, sensation, or appearance, generally indicating disorder or disease” (personal communication from Patient Reported Outcomes Harmonization Group, 2002). Since a significant proportion of patients with GORD do not have erosive oesophagitis,1 evaluation of GORD symptoms is a key part of diagnosis,2 while monitoring the response of symptoms to therapy is crucial in assessing the effectiveness of the intervention.

The key objectives, in terms of data quality, in measuring symptoms are to collect data that are:

- valid (that is, are a “pure” measure of the target symptom and are not distorted by other interfering variables);
- reliable (that is, are reproducible in the sense that the same measurement is obtained when there has been no change in symptoms);
- unbiased (that is, symptoms are not measured in a way that systematically under or over-estimates the true value);
- discriminating (that is, the measure can distinguish adequately between individuals who actually do differ to an important degree in terms of the variable measured—such as symptom severity);
- responsive to change (analogous to discriminating, but referring to the ability to detect true change over time or in response to an intervention).

Important considerations are: who should assess symptoms (patient versus clinician); the method of data collection (questionnaires versus diaries); the timing of assessment (reference period and duration); and the number of symptoms to be assessed (in particular, the relative merits of single item measures, multiple items analysed as a profile, summated scores, or global assessments).

WHO SHOULD ASSESS SYMPTOMS
Previous findings on clinician versus patient ratings of symptoms
There is some evidence from studies in which independent measures of symptoms could be made—for example, patient reports of palpitations compared with documented arrhythmias from 24 hour ambulatory electrocardiograph monitoring3—of a lack of concordance between subjective reports of the incidence and severity of symptoms and objective measures of disease activity. In general, however, the subjective

Abbreviations: GORD, gastro-oesophageal reflux disease

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nature of symptoms, as implied by the Patient Reported Outcomes Harmonization Group’s definition above, suggests that self report is the most appropriate method of gathering data on symptom experience. Findings from studies across a range of conditions, including GORD,6–10 show a general pattern of only weak to moderate agreement between patient and clinician ratings, both for symptoms and for quality of life impairment. Although discrepancies in both directions have been reported, the most common finding is for clinicians to underestimate the incidence and severity of symptoms, relative to patient ratings.6 Mismatches generally appear to be greater among patients with less severe symptoms. Agreement on the presence versus the absence of symptoms is usually better than agreement about symptom intensity.11

**Empirical evidence of mismatches between clinician and patient ratings of symptoms in GORD**

Previous research into mismatch between patient and clinician ratings of symptoms was borne out in secondary analyses of data on file at AstraZeneca, which were carried out specifically for this workshop. The data related to a large sample of both male and female patients, aged 18–80 years, whose dominant symptom was heartburn. These patients were included in a multicentre, double blind, randomised, parallel group study, comparing four week courses of three drug regimens; for the purposes of these secondary analyses, data from all three arms of the trial were combined. Clinician and patient ratings, at three points in time (baseline, two, and four weeks), of the severity of three symptoms of GORD (heartburn, epigastric pain, and regurgitation) were obtained. Our analyses showed that the level of agreement was poorest at baseline; clinicians underestimated symptom severity relative to patients’ ratings of heartburn in 34% of cases; underestimates of the severity of epigastric pain and regurgitation were observed in 55% and 46% of cases, respectively. At baseline, clinicians judged that 40% of patients were free from epigastric pain but only 14% of patients considered themselves to be free of this symptom; the corresponding values for regurgitation were 32% and 14%. Low clinician or investigator ratings of symptom severity at baseline represent a potential “floor effect” with little scope for improvement over time.

Levels of agreement between clinician and patient ratings improved over time. At four weeks, patients and clinicians agreed on heartburn severity in 78% of cases; agreement in respect of severity of epigastric pain and regurgitation was observed in 58% and 76% of cases, respectively. This convergence of patient and clinician ratings may be, in part, an artefact of the low investigator ratings at baseline coupled with patients’ perceptions of improvement over time, thus bringing the two ratings more closely in line.

**Implications of findings regarding clinician versus patient ratings**

The implications of our findings and those of previous researchers are that when clinicians’ ratings are used, estimates of absolute treatment effects at any given point in time are likely to be upwardly biased (that is, overestimate the impact of treatment) because symptom severity is underestimated. Conversely, by using clinicians’ ratings, comparisons of changes in symptom severity over time or in response to a therapeutic intervention are likely to underestimate the impact of treatment because of the floor effects and trend towards convergence noted above. In a well designed randomised controlled trial, there is no reason to believe that the mismatch between clinician and patient ratings will vary across the different arms of the trial and, therefore, estimates of the effects of one treatment relative to another should not be affected. However, estimates of absolute treatment effect and, therefore, calculations of number needed to treat, are likely to be biased, as described above.

**Potential explanations for mismatches between clinician and patient ratings**

Several possible explanations have been proposed for the lack of concordance between patient and clinician ratings. Lack of interrater reliability between clinicians has been demonstrated.11 Holmes and colleagues12 have suggested that clinicians may focus primarily on frequency and intensity of symptoms in their history taking while patients also consider symptom related disability and the impact on quality of life in judging the severity of their symptoms. Lack of a common vocabulary may also underpin discrepancies. Agreus and colleagues13–14 reported that the terms “pain” and “discomfort” do not adequately reflect the full range of descriptors used by the general public in describing dyspepsia. Stanghellini15 suggests that patients may attribute different meanings to “pain” and “discomfort”. Similarly, ambiguity of the term “heartburn” has been highlighted16,17; Carlsson and colleagues18 have shown how a “word picture” in which heartburn was defined as “a burning feeling rising from the stomach or lower chest up towards the neck” facilitated elicitation of experience of this symptom.

Patients may underreport (or indeed exaggerate) symptoms to clinicians. Underreporting may occur when patients wish to please the clinician or researcher10 and thus represents a social desirability bias. Overreporting may occur if there is a perceived benefit to the patient (for example, being certificated as unable to work).

Patients and clinicians may also use different frames of reference in assessing symptoms.19 For example, patients are likely to use an internal frame of reference—either in relation to their own past experiences of symptoms or to an ideal state of health—while health care professionals may implicitly compare a given patient to other patients whom they have treated.18 The use of a structured record or questionnaire16 may facilitate elicitation of symptoms by a clinician but even this will not guarantee consistency in characterising or assessing symptoms.11

**METHOD OF DATA COLLECTION**

In collecting self report data on symptoms, the two main methods of data collection are questionnaires (most usually self completed, paper and pencil questionnaires, but electronic questionnaires and interviewer administered instruments may also be used) and diaries. Approximately 25% of all clinical trials include diaries as a method of data collection.19 Several diaries have been developed and used with GORD patients.20,21 Structured self completion questionnaires have also been developed and validated in GORD patients.22–30 Both diaries and questionnaires have advantages and disadvantages.

**Advantages and disadvantages of diaries as a mode of data collection**

Diaries, in theory at least, allow for data to be recorded contemporaneously, thus reducing the risk of recall bias. They allow day to day fluctuations in symptoms to be readily captured and, therefore, facilitate calculation of symptom free days. Therapeutic intervention can be matched with exposure to potential exacerbating factors (such as ingestion of certain foods) can be captured more easily through contemporaneous accounts, as can details of self management behaviour.19 However, completion of a daily diary imposes considerable respondent burden, and several studies19,22–30 present objective evidence of non-adherence to data collection protocols, although the diary respondents...
themselves reported good compliance. In particular, the problem of “hoarding” (retrospective completion of diary entries) has been highlighted; with such retrospective data recording, recall errors may occur. Unfortunately, hoarding is almost impossible to detect unless electronic devices, which can unobtrusively record the timing of data entries, are used. The spectre of fatigue and attrition effects—a drop off in diary completion rates over time—has also been cited as a potential weakness of this method of data collection but this criticism is not supported by findings on participation and completion rates for diary studies. Burman reports that typical participation rates are in excess of 80% while Rogmann and Haggerty and Verbrugge have shown very typical participation rates are in excess of 80% while resulting in higher rates of reporting as time goes by.

Diary methods require respondents to remember whether and when they experienced symptoms. Types of recall error include omission (completely forgetting that the symptom was experienced in the time frame of interest) and “telescoping” (misplacing an event in time, most usually by recalling it as occurring more recently than it actually did). Dahlquist and colleagues have demonstrated that errors of recall in respect of minor symptoms are likely to occur when the time frame of reference exceeds one week, with minor symptoms being forgotten. Recall biases have also been noted in studies in which estimates of pain relief rely on recollections of past pain, leading to overestimation of treatment effects.

When symptoms fluctuate over time, the need to summarise may present patients with a complex cognitive task; they may employ short cuts or heuristics in producing their response, again threatening the validity of the answer. Steen and colleagues experimented with one and three month reference periods in assessing symptoms of asthma and diabetes. They found that patients commonly used an “anchor and adjust” heuristic in respect of the three month recall period; respondents first estimated how often they had experienced the symptom of interest in the preceding month and then adjusted the three month estimate upwards or downwards from that initial anchor. The risk of social desirability bias—for example, under-reporting consumption of certain foods that may trigger symptoms—may also be greater when data are collected by means of questionnaires, especially when these are interviewer administered.

**Advantages and disadvantages of questionnaires as a mode of data collection**

Questionnaire methods require respondents to remember whether and when they experienced symptoms. Types of recall error include omission (completely forgetting that the symptom was experienced in the time frame of interest) and “telescoping” (misplacing an event in time, most usually by recalling it as occurring more recently than it actually did). Dahlquist and colleagues have demonstrated that errors of recall in respect of minor symptoms are likely to occur when the time frame of reference exceeds one week, with minor symptoms being forgotten. Recall biases have also been noted in studies in which estimates of pain relief rely on recollections of past pain, leading to overestimation of treatment effects.

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**Diaries or questionnaires—whch mode of data collection is best**

It is generally held that diaries are the “gold standard” method of collecting symptom data; rates of reporting minor symptoms are generally higher in diaries than in questionnaires, and the implicit assumption is that “higher reporting is more accurate”. However, adequate agreement between patterns of symptoms, as recorded in diaries and reported in questionnaires, has been demonstrated in a number of studies. Questionnaires may, therefore, be an adequate method of data collection in situations where a contemporaneous account is not essential or where a measure of “symptom free” days is not required.

**Good practice in diary design**

Two main styles of diaries are used in health studies. In a ledger style diary, the occurrence of a symptom or other health related event (such as an encounter with a health professional) is recorded and dated as the event occurs; thus entries relate only to days on which the phenomenon of interest occur. In a journal style diary, an entry is made each day (or defined time period), regardless of whether an event occurs. In this style of diary, provision must be made for recording the absence of the phenomenon on that day (for example, that the symptom of interest was not experienced). Ledger style diaries reduce respondent burden but there is a greater risk of the respondent forgetting to make an entry when it would be appropriate to do so and, therefore, there is greater difficulty in distinguishing between the absence of the symptoms and missing data.

For either style of diary, a further choice lies between a highly structured format, using closed questions, and a less structured approach, with open ended items. In symptom diaries, the structured approach, using a predefined list of symptoms and associated response categories (for example, presence/absence, or rating of symptom severity) ensures equivalence of stimulus (thus increasing the reliability of the response), reduces respondent burden, and facilitates data coding and entry. Most diaries used with GORD patients to date have been highly structured based, for example, on the gastrointestinal symptoms rating scale. Well designed diaries and clear instructions on how they are to be completed enhance both the quantity and quality of data collected. Face to face instruction on diary protocols is desirable and telephone or electronic prompts seem to be more effective than postal reminders. Electronic diaries have been shown to be an effective and well accepted alternative to traditional paper based methods.

**Good practice in questionnaire selection and design**

Considerations in selecting a questionnaire for use with GORD patients include: the purpose for which the questionnaire was designed; evidence of validity, reliability, and responsiveness to change; the population on which it was tested and validated; the time frame of reference; and practical considerations of cost and copyright. Kirshner and Guyatt have distinguished between instruments designed for purposes of discrimination (diagnosis and screening), prediction (of response to treatment or of prognosis), and evaluation (assessing response to an intervention). They have highlighted that the relative weight accorded to the psychometric properties of validity, reliability, discriminatory power, and responsiveness to change will depend on the purpose. Although their focus was on quality of life instruments, the same distinctions are likely to be relevant with respect to symptom questionnaires. An instrument designed primarily as a diagnostic tool may not be sufficiently responsive to change for use in the context of evaluation.

Moreover, properties of validity, reliability, and so on are not universal. For this reason, a questionnaire developed and validated in, say, a secondary care population in the USA may need to be adapted and revalidated for application in a primary care setting or in another culture. Issues of cross cultural adaptation are discussed elsewhere in this issue
(Wyrvich and Staebler Tardino in this supplement (see page iv5–iv48).

Some symptom questionnaires are in the public domain and may be freely used by researchers and practitioners. In other cases, access may require completion of a copyright or users' agreement, possibly with explicit conditions of use and/or a charge for reproduction of the questionnaire and access to data processing instructions. Practical barriers to usage include unduly restrictive conditions of use or excessive costs.

As with diaries, careful attention to questionnaire design and administration can enhance both the quantity of response and the quality of that response. Jenkins and Dillman provide guidelines for questionnaire design and layout, based on principles of cognitive psychology. A comprehensive review of methods to enhance the rate of response to questionnaire surveys is provided by Edwards and colleagues.

**TIMING OF ASSESSMENTS**

The duration and frequency of symptom assessment must take into account the purpose of data collection and the natural history of the disease and anticipated response to any therapeutic intervention. There is often a need to strike a balance between what is desirable and what is feasible (for example, in terms of keeping respondent burden to an acceptable level).

**Use of a reference time period**

Dillman defines behaviour to include “what has happened in people's lives”, and this definition includes experience of symptoms. Sudman and Bradburn advise that the reliability of responses to behaviour questions can be improved by asking about a specific time period. In relation to asking about symptoms, Steen and colleagues state that the choice of reference period should take into account their prevalence and incidence. If the point prevalence is low and the reference period in question is very short, few patients may respond positively. Low item endorsement may reduce the utility of the measure in detecting differences between individuals and change over time.

Booth and colleagues have reported significant daily fluctuations in symptoms of GORD. Their findings suggest that asking only about “today” (or, indeed, any single day) may produce an incomplete picture of overall symptom burden, and that longitudinal data, reflecting day to day variability in symptoms, may be required. However, as already noted, recall of minor symptoms is likely to be poor when the reference period exceeds one week. Booth and colleagues found that one month was the maximum period over which patients could provide reliable data on frequency of symptoms of asthma or diabetes, and on the impact of those symptoms. The time frames referred to in questionnaires appropriate to GORD patients vary, although reference periods of 2–4 weeks are common. The interaction between the choice of reference period and timing of assessment must also be considered. Clearly, a questionnaire referring to symptoms “in the last four weeks” would be inappropriate for use two weeks after initiation of therapy.

**Duration of symptom recording**

Burman recommends that, in diary studies, the period of recording should represent a balance between the avoidance of excessive respondent burden and the necessity to capture adequately the anticipated fluctuations in the phenomenon of interest, as discussed above. Carp and Carp found that problems reported on a selected day in their diary study were not an adequate predictor of problems across the other days of that week, and concluded that short diary periods could be problematic. However, protracted periods of data recording (in excess of six months) pose significant respondent burden and are, therefore, likely to be associated with poorer compliance with diary protocol. Burman indicates that typical diary periods tend to be 2–4 weeks when continuous data recording is required. An alternative strategy to reduce respondent burden when longer term data are required is for diaries to be completed only on a random selection of days in the data collection period although with this strategy there is the risk that respondents will forget to fill in the diary on the designated dates.

**ASSESSMENT OF MULTIPLE SYMPTOMS**

In patients with GORD, the most characteristic symptom is heartburn but patients also report epigastric pain, regurgitation, belching, early satiety, acid reflux, dysphagia, and other symptoms. Moreover, for some individuals heartburn is not the dominant symptom. Furthermore, therapeutic interventions may themselves give rise to symptomatic side effects, both within and beyond the digestive tract. For these reasons, consideration must be given to the number of symptoms to be measured, whether multiple items (questions) are needed to measure each symptom, whether it is appropriate to produce an overall assessment (for example, of symptom severity), and, if so, how that score should be computed.

**How many symptoms, and how many dimensions**

The issue of how many symptoms to assess will depend on the purpose of assessment. For diagnostic purposes, it is the combination and coincidence of symptoms that may serve to distinguish those patients who are more likely to be experiencing GORD from those with other disorders of the gastrointestinal system. In therapeutic interventions, the reduction of heartburn may be the primary objective but patients will generally also expect reduction of other symptoms and will not welcome the development of new symptoms as a side effect of therapy. In these situations, it may be desirable to assess the response of multiple symptoms to the intervention.

The experience of symptoms is likely to be multifaceted, including timing, frequency, duration, intensity, effects on role function, and distress caused by symptom occurrence. Depending on the purpose of assessment, one or more facets of symptom experience may need to be measured. Keeffe has reported that pain intensity measures are flawed by focusing “only on one dimension of multidimensional experience”.

**Challenges of multiple measures**

Although justified by the multifaceted experience of symptoms, the use of more than one measure presents problems of multiple end points in statistical analysis and interpretation. Similar problems arise in relation to other multidimensional outcome measures, in particular health related quality of life. One solution is to define the symptom of greatest relevance as the primary end point (and in clinical trials or surveys to base power calculations on that item) and to treat the remaining symptoms as secondary end points. Another is to combine scores on individual symptoms to produce an overall measure of, say, symptom severity. The simplest way of doing this is simply to sum the individual responses. However, there are two assumptions implicit in this approach. The first is that symptoms are being measured on an interval scale and that the “distance” between each pair of points on the scale is the same (for example, if symptoms are categorised as “none”, “mild”, “moderate”, and “severe”, that the difference between “none” and “mild” is the same as between “mild” and “moderate” or between “moderate” and “severe”). The second is that all symptoms are given the same weight (that is, are of equal importance). Neither of these assumptions may be warranted.
Global assessments of symptoms

An alternative to asking individually about each symptom, and combining responses as described above, is to ask respondents to make a global assessment of their symptoms, using either a visual analogue scale or a set of verbal descriptors. This approach is also used in quality of life research. Global assessments may, however, be cognitively demanding for respondents, requiring them to: judge which symptoms to include in the global rating; retrieve from memory specific and relevant information on individual symptoms; synthesise the information retrieved into a single overall judgement; and to “map” the adjudged response onto one of the response categories offered. Respondents are likely to use a range of cognitive strategies and heuristics in making their assessment, and may not use comparable frames of reference, thus threatening the validity and reliability of responses. Making a global rating may also be difficult when some symptoms are improving while others are unchanged or deteriorating—Zielbland and colleagues observed that “the global transition item ... obscures a feature of (rheumatoid arthritis) activity whereby some functions may improve while others are in decline”. None the less, several studies in which global ratings have been compared with summed symptom scores have demonstrated reasonable agreement between the two approaches.

Global assessments of change over time or in response to an intervention have also been successfully used in quality of life research.

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