Self-assessment of faecal pH and faecal bulk in epidemiological studies

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A high faecal pH and a low faecal bulk have been associated with an increased risk of colorectal cancer. The evidence is based on retrospective epidemiological studies (Glober et al., 1977; Jensen et al., 1982; Pietroiuistti et al., 1983). Prospective epidemiological studies comparing a large number of subjects require simple and inexpensive methods of data collection. Faecal pH and bulk probability can be determined by self-testing (Free et al., 1984).

We determined the accuracy of self-assessment of faecal pH and bulk in a study population, heterogeneously in dietary fibre intake, i.e. vegetarians (V) and non-vegetarians (NV). Dietary fibre has been shown to lower faecal pH and to increase faecal daily wet weight (van Dokkum et al., 1983).

Subjects were recruited during the pilot stage of the Dutch prospective cohort study on diet and cancer (van den Brandt et al., 1990). NV were recruited from an age (55–69 years) and gender stratified sample, drawn from 23 municipalities in The Netherlands. V were recruited by (advertisement in the magazine of) the Dutch Vegetarian Society.

Faecal pH self-assessed by the investigator (A.v.F.) and the volunteers using the Combur-3 test of Boehringer Mannheim, which shows a clear colour change in the pH range 6 to 9, while the colours can be distinguished from the brown faeces. Moreover, the test strip is easy to manipulate, because it has a plastic handle and the pH test is the patch nearest to this handle; the other two patches (for measurement of protein and glucose) were removed. Stool weights were estimated by the volunteers using black and white photographs (18 cm by 24 cm) of frozen stools weighing approximately 40, 120, 200, 300 and 400 g, representing the range of faecal weights among our Institute’s employees. The self-assessments were performed during 4 consecutive days. The stools were collected and mailed to the laboratory in a styrofoam box with dry ice (frozen carbon dioxide, –78°C). In the laboratory faecal bulk was measured by weighing. After thawing at 4°C the stools were mixed by kneading vigorously. The homogeneity of the mixture was transferred to plastic bowls and mixed again by stirring with a wooden spatula. Faecal pH was measured with a pH electrode for small samples on at least two positions and the mean pH was recorded. This measurement appeared to be feasible and reproducible. For a few samples it took about 1 min before the value had stabilised, but most samples could be measured immediately. The coefficient of variation as calculated from 20 duplicate measurements, was 2%.

Table I shows the variance of the laboratory measurements as explained by self-assessment. The percentage of variance explained in the laboratory pH measurement increased from the first to the third stool, suggesting a learning effect. For stool bulk the explained variance was higher than for faecal pH and remained stable from the first to the fourth stool. Figure 1 shows the regression line for pH as measured by the pH meter on pH measured by the volunteers in the third stool. The investigator (A.v.F.) achieved a considerably higher accuracy, although the regression coefficients were similar.

We evaluated the influence of misclassification due to the accuracy of self-assessments of faecal pH and bulk on hypothetical relative risks (R.R.) for developing colorectal cancer, in the way Copeland et al. (1977) described. Assuming the true R.R. for colorectal cancer of faecal pH > 6.5 compared to pH < 6.5 is 2, the observed R.R. in a case-control study nested in a cohort study would be 1.3. For faecal bulk < 145 g/day vs > 145 g, a hypothetical R.R. of 2 decreased to 1.7. This means we need a sample size approximately 4-fold larger than if no error existed (Willett, 1991). This makes the study very expensive (self-assessment of faecal pH by 1 million people will cost about 250,000 pounds).

Table I

| Order of stool | Number of volunteers | Explained variance (%) | pH | bulk |
|---------------|----------------------|------------------------|----|------|
| 1st           | 37                   | 5                      | 40 |      |
| 2nd           | 34                   | 19                     | 33 |      |
| 3rd           | 27                   | 19                     | 47 |      |
| 4th           | 9                    | 26                     | 49 |      |

Figure 1 Regression line obtained by the investigator in stools of the Institutes’ employees (--O--): y = 4.13 × 0.4x; s.e.(b) = 0.11; r2 = 0.58; residual s.d.(y) = 0.32. Regression line obtained by volunteers in third stool (--•--•): y = 4.00 × 0.41x; s.e.(b) = 0.10; r2 = 0.38; residual s.d.(y) = 0.45.
significantly (mean of 26 NV = 6.7; mean of 17 V = 6.8). As expected from the difference in dietary fibre intake, faecal bulk was significantly higher for V (mean = 189 g wet weight/24 h; n = 15) than NV (mean = 122 g/24 h; n = 33).

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References

VAN DEN BRANDT, P.A., GOLDBOHM, R.A., VANT VEER, P., VOLOVICZ, A., HERMUS, R.J.J. & STURMANS, F. (1990). A large-scale prospective cohort study on diet and cancer in the Netherlands. J. Clin. Epidemiol., 43, 285–295.

COPELAND, K.T., CHECKOWAY, H., MCMICHAEL, A.J. & HOLBROOK, R.H. (1977). Bias due to misclassification in the estimation of relative risks. Am. J. Epidemiol., 105, 488–495.

DOKKUM, W., VAN DE BOER, B.C.J., FAASSEN, A., VAN PIKAAR, N.A. & HERMUS, R.J.J. (1983). Diet faecal pH and colorectal cancer. Br. J. Cancer, 48, 109–110.

FREE, A.H. & FREE, H.M. (1984). Self testing, an emerging component of clinical chemistry. Clin. Chem., 30, 829–838.

GLOBER, G.A., KAMIYAMA, S., NOMURA, A., SHIMADA, A. & ABBA, B.C. (1977). Bowel transit-times and stool weight in populations with different colon-cancer risks. Lancet, ii, 110–111.

JENSEN, O.M., MACLENNAN, R. & WAHRENDORF, J. (1982). Diet, bowel function, fecal characteristics, and large bowel cancer in Denmark and Finland. Nutr. Cancer, 4, 5–19.

PIETROIUSTI, A., GIULIANO, M., VITA, P., CIARNIELLO, P. & CAPRILLI, R. (1983). Faecal pH and cancer of the large bowel. Gastroenterology, 84, 1273.

WILLETT, W.C. (1991). The use of biomarkers in nutritional epidemiology. In Biomarkers of Dietary Exposure, Kok, F.J. & van't Veer, P. (eds) p. 9. Smith-Gordon and Company Limited: London.