

# Epidemiology and impact of multimorbidity in primary care: a retrospective cohort study

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## ABSTRACT

### Background

In developed countries, primary health care increasingly involves the care of patients with multiple chronic conditions, referred to as multimorbidity.

### Aim

To describe the epidemiology of multimorbidity and relationships between multimorbidity and primary care consultation rates and continuity of care.

### Design of study

Retrospective cohort study.

### Setting

Random sample of 99 997 people aged 18 years or over registered with 182 general practices in England contributing data to the General Practice Research Database.

### Method

Multimorbidity was defined using two approaches: people with multiple chronic conditions included in the Quality and Outcomes Framework, and people identified using the Johns Hopkins University Adjusted Clinical Groups (ACG<sup>®</sup>) Case-Mix System. The determinants of multimorbidity (age, sex, area deprivation) and relationships with consultation rate and continuity of care were examined using regression models.

### Results

Sixteen per cent of patients had more than one chronic condition included in the Quality and Outcomes Framework, but these people accounted for 32% of all consultations. Using the wider ACG list of conditions, 58% of people had multimorbidity and they accounted for 78% of consultations. Multimorbidity was strongly related to age and deprivation. People with multimorbidity had higher consultation rates and less continuity of care compared with people without multimorbidity.

### Conclusion

Multimorbidity is common in the population and most consultations in primary care involve people with multimorbidity. These people are less likely to receive continuity of care, although they may be more likely to gain from it.

### Keywords

chronic disease; comorbidity; family practice; primary health care; outcome and process assessment (healthcare); prevalence.

## INTRODUCTION

The practice of medicine is becoming increasingly specialised, both in hospitals and in general practice. For example, many practices now offer chronic disease management clinics for conditions such as diabetes. This approach of treating each condition in isolation has serious limitations. It is important to recognise that many people have multiple coexisting chronic medical conditions, or 'multimorbidity'.<sup>1</sup> People with multimorbidity are likely to have complex needs for health care and to account for a high proportion of the healthcare workload. Increased understanding of the epidemiology and impact of multimorbidity is needed to inform the way in which health care is organised and delivered.

The basis of primary care is that generalists manage all health problems commonly occurring in the population, identifying and referring those problems needing specialist care, and coordinating care for patients with complex health problems.<sup>2</sup> However, efforts to improve quality of care have fuelled a move towards specialisation within general

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practice, and an emphasis on improving access has led to a multiplicity of providers, with patients being less likely to consult the same professional on each occasion.

It is important to consider how the balance should be struck between care provided by professionals with generalist training, able to tackle a wide range of problems at one consultation, or care provided by a wide range of specialists with cross-referral between themselves. This debate needs to be informed by data about the extent and nature of multimorbidity in patients consulting in primary care.

Several studies have examined the prevalence of multimorbidity in different countries. These have reached varied conclusions as a result of differences in setting, the range of health conditions included, and data sources.<sup>3-5</sup> Some studies have been based on surveys or administrative data, or have been restricted to older populations.<sup>6,7</sup> Most studies based on medical records have involved relatively small numbers of practices and/or patients,<sup>3</sup> while larger studies have focused on the determinants and prevalence of multimorbidity,<sup>8-12</sup> but few have related this to process or outcome variables in primary care.<sup>13,14</sup> Earlier research on multimorbidity has been limited by problems with case definition and with the reliability of data recording within routine practice records.<sup>15</sup> There is little information about the epidemiology of multimorbidity in the UK.

The almost universal computerisation of general practice records in the UK over the last 20 years has been accompanied by the development of research databases that combine quality-assured data from a large and representative range of practices. In addition, the introduction of the Quality and Outcomes Framework (QOF) has led to the consistent recording of diagnoses of many important chronic health conditions. These factors enable population-based research into multimorbidity. The aims of this study were to gain a detailed and reliable understanding of the epidemiology of multimorbidity in England, and relationships between multimorbidity, consultation rate, and longitudinal continuity in primary care.

## METHOD

### Sample

This study is based on the anonymised records of a random sample of 99 997 adult patients (aged 18 years or over) from 182 practices in England contributing data to the General Practice Research Database (GPRD). These were all the practices in the GPRD that had provided 'research standard' data continuously from 1 April 2005 to 31 March 2008, and had given consent to link their data to measures

## How this fits in

Several studies have described the prevalence and determinants of multimorbidity, but these have produced varied results. There is little published information on the prevalence of multimorbidity in the UK. Computerisation of general practice records in the UK along with reliable coding of many chronic diseases, due to the Quality and Outcomes Framework, have made it possible to obtain reliable estimates of multimorbidity. This study shows that multimorbidity is very common in the population, particularly in older people and those living in deprived areas. The majority of consultations in general practice involve people with multimorbidity. People with multimorbidity are frequent users of primary care and are less likely to receive continuity of care. These findings support the importance of people having access to a generalist primary care service that is able to coordinate care for a wide range of problems in one individual.

of area deprivation. Patients were eligible for inclusion if they were aged 18 years or over and were registered with one of the participating practices on the index date of 1 April 2005. Sample selection was stratified by practice, age group, and sex. Data were obtained with regard to all diagnoses, consultations, and prescriptions entered until 31 March 2008, including both diagnoses entered contemporaneously and earlier diagnoses that had been entered retrospectively by practices from earlier paper notes.

### Defining multimorbidity

Multimorbidity was operationalised in two ways. The primary approach was to define it as a patient who, on the index date, had more than one of 17 important chronic conditions for which care is incentivised under the QOF (Box 1). The QOF business rules<sup>16</sup> specify the Read Codes which are used to define each condition. These rules were applied to the diagnostic codes within the GPRD data to determine whether or not individuals had each chronic condition.

As in the QOF itself, patients were included if they had ever had the condition unless there was a code to indicate that it had resolved. The advantage of this approach is that recording of QOF conditions is likely to be reliable and complete, as recording is linked to payments to practices, and case definition is well defined. However, although these 17 conditions include many of the most important chronic conditions, many others are not included (for example, skin disease and liver disease). Therefore, this approach underestimates the true prevalence of multimorbidity.

As a secondary approach, the Johns Hopkins University Adjusted Clinical Groups (ACG<sup>®</sup>) Case-Mix System<sup>17</sup> was used to identify whether or not each patient had one or more of a much wider list of

**Box 1. Chronic conditions included in the Quality and Outcomes Framework and counted within the definition of multimorbidity.**

- Asthma
- Atrial fibrillation
- Cancer
- Coronary heart disease
- Chronic kidney disease
- Chronic obstructive airways disease
- Dementia
- Depression
- Diabetes
- Epilepsy
- Heart failure
- Hypertension
- Learning disability
- Mental health problem (psychosis, schizophrenia, or bipolar affective disorder)
- Obesity
- Stroke
- Thyroid disease

chronic conditions. One problem with operationalising multimorbidity based on a count of chronic diseases entered in routine medical records is that the same disease may be coded in different ways and therefore counted twice in the same individual. The ACG system uses a software programme to collapse a wide range of diagnostic codes found within patients' records into 260 mutually exclusive clinically homogeneous 'expanded diagnostic clusters' (EDCs).

Three investigators, all of whom are GPs, independently assessed whether each of the diagnostic clusters should be included as a chronic condition. Differences were resolved by consensus. A chronic condition was defined as one that normally lasts 6 months or more, including past conditions that require ongoing disease or risk management, important conditions with a significant risk of recurrence, or past conditions that have continuing implications for patient management. In this way, 114 of the 260 diagnostic clusters were defined as chronic conditions (Appendix 1).

The ACG software was used in conjunction with the diagnostic data in the GPRD dataset to determine whether or not each patient had ever been diagnosed with each condition, and an EDC-based chronic condition count per patient was calculated.

### Analysis

The prevalence of multimorbidity was estimated in relation to age, sex, and deprivation. Age-standardised prevalence estimates were calculated using the European Standard Population.<sup>18</sup> Deprivation was based on Townsend scores derived from the patient's postcode and national quintiles using 2001 census data. Based on previous research, it was anticipated that multimorbidity would show a positive association with age, female sex, and deprivation.<sup>3,8,12,19,20</sup>

The study sought to investigate associations between multimorbidity, consultation rates, and continuity of care. Consultation rates were calculated over 3 years beginning 1 April 2005, and were based on consultations with GPs or practice nurses, including face-to-face and telephone consultations. Annual rates were calculated, taking account of patients who left the practice or died during the period studied. Using the total number of consultations (rather than the number of patients) as the denominator, the proportion of consultations that involved patients with different levels of multimorbidity was also explored.

Longitudinal continuity of care over 3 years was calculated, making use of the fact that each consultation in the GPRD dataset has identifiers to indicate type of consultation, which clinician conducted it, and that clinician's profession. The usual provider continuity index<sup>21</sup> and the continuity of care index<sup>22</sup> were calculated. The usual provider continuity index represents the proportion of all consultations with the clinician who had been consulted most often, and the continuity of care index is a measure of the concentration of consultations with different doctors, adjusting for the number of consultations. Face-to-face and telephone consultations with GPs or practice nurses were included in continuity analyses. Patients with fewer than two consultations in the 3-year period were excluded.

### Statistical analysis

For both the QOF and the ACG/EDC approaches, all analyses were repeated treating multimorbidity as a binary variable (none or one chronic condition versus more than one) and also as a discrete variable (number of conditions). Descriptive statistics, including 95% confidence intervals, were used to describe relationships between multimorbidity and other variables. Independent relationships between age, sex, deprivation, and multimorbidity when treated as a binary or discrete variable, were examined using logistic and Poisson regressions respectively. Consultation rate data were skewed and were therefore log transformed

before being included in multiple linear regression analyses with multimorbidity, age, sex, and deprivation as explanatory variables. Relationships between multimorbidity and continuity of care were similarly examined. All analyses were conducted in Stata (version 11.0). The 'svy' survey commands were used throughout to take account of the sampling methods.

## RESULTS

### Epidemiology of multimorbidity

Initial examination of the patient sample in comparison with census data indicated that it was representative of the population of England in terms of age–sex distribution, although with slightly lower levels of deprivation (mean Townsend score =  $-1.04$ ).

By the index date, 16% of the sample (16 030/99 997) had been diagnosed with more than one of the conditions included in the QOF, but 58% (58 115/99 997) had been diagnosed with more than one EDC chronic condition. The age-standardised prevalence was 14% using the QOF approach and 56% using the ACG/EDC approach.

Figure 1 demonstrates that multimorbidity is strongly related to age, and slightly more common in females than males below the age of 65 years. Patients aged 75 years or over had mean = 1.56 (95% CI = 1.52 to 1.59) QOF conditions and mean = 5.63 (95% CI = 5.52 to 5.74) EDC conditions. Figure 2 illustrates that 77% of patients aged 75 years or over had at least one QOF condition, 44% had more than one QOF condition (multimorbidity), and 9% had four or more conditions.

Table 1 shows that increasing age, female sex, and living in a deprived area were all, as hypothesised, independently associated with increased odds of having multimorbidity. Patients in the most deprived quintile for deprivation were almost twice as likely to have multimorbidity as those in the least deprived quintile (odds ratio 1.91 [95% CI = 1.78 to 2.04] adjusted for age and sex). Similar results were observed with the ACG/EDC approach, although the relationship with deprivation was less marked.

### Multimorbidity and consultation rates

The overall consultation rate was 4.63 (95% CI = 4.58 to 4.69) consultations per patient per annum. Consultation rates showed the anticipated relationships with age, sex, and deprivation, with increasing rates with age, higher consultation rates for females than males among those below the age of 65 years, and higher rates in more deprived areas. Patients with multimorbidity (based on QOF) had 9.35 (95% CI = 9.21 to 9.49) consultations per

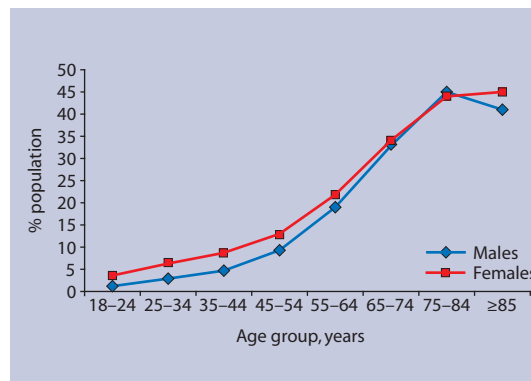


Figure 1. Percentage of population with more than one chronic condition in the Quality and Outcomes Framework, by age and sex.

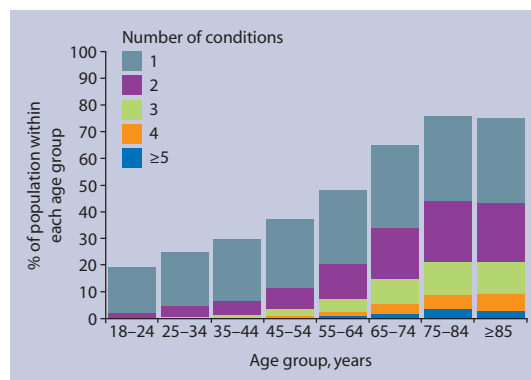


Figure 2. Number of conditions included in the Quality and Outcomes Framework, per patient, by age group.

Table 1. Independent associations between multimorbidity<sup>a</sup> and age, sex, and deprivation.

Variable	Odds ratio	95% confidence interval	P-value
Age (per 10-year increase; range $\geq 18$ years)	1.78	1.76 to 1.81	<0.001
Sex (reference category: male)	1.23	1.18 to 1.28	<0.001
Deprivation (per 10-unit increase in Townsend score)	2.08	1.95 to 2.22	<0.001

<sup>a</sup>Multimorbidity is defined as having more than one condition in the Quality and Outcomes Framework. All variables are adjusted for each other.  $n = 99\ 624$  due to missing deprivation data for 373 individuals.

annum compared with 3.75 (95% CI = 3.71 to 3.80) among those without multimorbidity.

Table 2 shows the independent relationships between consultation rate and age, sex, and deprivation before and after adjusting for multimorbidity. It demonstrates that the relationship between age and consultation rate is reduced, and that between deprivation and consultation rate almost disappears, after adjustment for multimorbidity. This suggests that the main reason that older patients and those in deprived areas consult more often is because they have more chronic health conditions.

### Proportion of consultations involving patients with multimorbidity

Using all consultations between 1 April 2005 and 31

**Table 2. Independent relationships between age, sex, deprivation, and consultation rate, before and after adjusting for multimorbidity.**

Explanatory variable	Consultation rate					
	Before adjustment for multimorbidity $R^2 = 0.19$			After adjustment for multimorbidity $R^2 = 0.30$		
	Coefficient	95% CI	P-value	Coefficient	95% CI	P-value
Age (per 10-year increase; range $\geq 18$ years)	0.17	0.17 to 0.18	<0.001	0.09	0.09 to 0.10	<0.001
Sex (reference category: male)	0.48	0.46 to 0.50	<0.001	0.42	0.40 to 0.44	<0.001
Deprivation (per 10-unit increase in Townsend score)	0.12	0.10 to 0.15	<0.001	0.03	0.01 to 0.05	0.01
Multimorbidity (number of QOF chronic conditions)	–	–	<0.001	0.37	0.36 to 0.38	<0.001

All variables were adjusted for each other.  $n = 99\ 624$  due to missing deprivation data for 373 individuals. QOF = Quality and Outcomes Framework.

March 2008 as the denominator ( $n = 1\ 430\ 773$ ), rather than the number of patients, the proportion of primary care consultations that involved people with multimorbidity was explored. Although patients with multimorbidity (QOF) represented only 16% of the population, they accounted for 32% of all consultations. Using the ACG/EDC approach, 58% of patients had multimorbidity, and these individuals accounted for 78% of all consultations.

Figure 3 shows the number of different chronic conditions encountered by primary care practitioners in an average consultation, in relation to patients' age. Consultations with patients aged over 75 years involved individuals with mean = 1.86 (95% CI = 1.82 to 1.91) QOF conditions and 6.68 (95% CI = 6.53 to 6.82) EDC conditions.

#### Multimorbidity and continuity of care

Levels of longitudinal continuity of care were low for all patients (mean = 0.25 on both indices). QOF multimorbidity was inversely associated with the usual provider continuity index (coefficient  $-0.08$  [95% CI =  $-0.09$  to  $-0.08$ ] after adjusting for age, sex, and deprivation). The equivalent model using the continuity of care index (which takes account of the number of consultations) showed a much smaller but

still inverse relationship between multimorbidity and longitudinal continuity (coefficient =  $-0.008$  [95% CI =  $-0.011$  to  $-0.004$ ]).

## DISCUSSION

### Summary of main findings

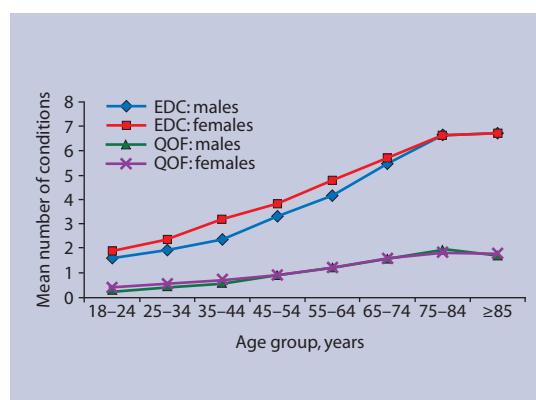
Most consultations in primary care involve patients with multimorbidity. These people have reduced longitudinal continuity of care, largely because of their high consultation rates. Although multimorbidity is common, the estimated prevalence varies considerably, depending on how it is measured. Prevalence increases with age, but affects patients of all age groups. Multimorbidity is more common among patients living in deprived areas.

### Strengths and limitations of the study

This article is based on a larger and more representative sample than earlier studies of the epidemiology of multimorbidity. As almost everyone in the UK is registered with just one general practice, participants are broadly equivalent to a population sample. The introduction of the QOF means that the coding of important diseases is likely to be consistent across participating practices, and the development of case-mix software such as the ACG system overcomes some of the coding problems encountered by previous studies.<sup>23</sup> The comprehensive nature of the data recorded with the GPRD makes it possible to explore relationships between multimorbidity and health processes and outcomes. This paper also builds on previous research by highlighting the high proportion of consultations that involve people with multimorbidity, which is important from a health-system perspective.

The study has several limitations. The large sample means that it is important to consider the magnitude of associations rather than  $P$ -values. The difference between the findings obtained using the

**Figure 3. Number of chronic conditions in patients consulting, per consultation, by age group. EDC = expanded diagnostic clusters from the Johns Hopkins University Adjusted Clinical Groups (ACG) Case-Mix System.<sup>17</sup> QOF = Quality and Outcomes Framework.**



QOF or ACG/EDC approaches illustrates the difficulty of providing precise estimates of the prevalence of multimorbidity, as this depends on the range of conditions included. Because the QOF approach only includes a limited number of conditions, it underestimates prevalence. The ACG/EDC approach includes a comprehensive list of chronic conditions, but in the absence of an internationally recognised list of conditions defined as chronic, the authors had to generate their own, and other investigators may have identified a different list.<sup>24</sup> In addition, not all of the 'chronic' conditions included may have been active or relevant in a particular patient. The prevalence of multimorbidity should therefore always be stated in relation to the measure used.

Both the QOF and the ACG/EDC approaches represent disease counts, with each disease counted equally. Case-mix adjustment methods that weight diseases differentially to estimate the burden of illness (such as the Charlson index<sup>25</sup> or the full ACG system<sup>17</sup>) are likely to predict outcomes more effectively,<sup>1</sup> but, unlike a disease count, they do not provide a direct measure of multimorbidity as usually defined in terms of multiple conditions. Furthermore, this paper is based on a definition of multimorbidity as the coexistence of multiple diseases. This approach provides a limited and medicalised perspective, which may not reflect patients' understanding of their problems.<sup>1</sup>

Estimates of multimorbidity in this study are based on diagnoses recorded in medical records. Different estimates of prevalence are obtained using different sources of data, such as GP records, patient surveys, or studies involving examination of patient cohorts.<sup>5,26</sup> Studies based on medical records will underestimate multimorbidity because some diseases are undiagnosed, and because they will not identify people who do not consult. Conversely, the relationship between multimorbidity and consultation rate has a risk of circularity, in that people who consult more often may have more conditions diagnosed.<sup>27</sup>

### **Comparison with existing literature**

This study supports previous research with regard to the strong relationships between multimorbidity and increasing age<sup>3,8,12</sup> and social disadvantage.<sup>8,12,20</sup> Previous studies have examined the relationship between multimorbidity and hospitalisation, but few have explored the relationship with utilisation of primary care, and these studies are not directly comparable.<sup>28-30</sup> Apart from a small study in just one practice,<sup>31</sup> it was not possible to identify any previous research examining the relationship between multimorbidity and continuity of care.

This paper adds to previous literature because of the size and generalisability of the sample, the reliability of the data sources, and the comparison of different approaches to measuring multimorbidity. It is also the first comprehensive published study of multimorbidity in the UK.

### **Implications for clinical practice and future research**

Most consultations in primary care involve patients with multiple conditions, and these may need to be taken into account when making decisions about patient management. Interactions between the conditions may further complicate decision making.<sup>1</sup> Guidelines are available to inform the management of many chronic diseases but these often fail to offer clear guidance about patients with comorbidities.<sup>32</sup> Therefore, practitioners working in primary health care need to have a broad knowledge base, be able to coordinate care across a wide range of different specialist services, have excellent communication skills to manage complex consultations in a limited time, and have good judgment to balance competing priorities.

Given that so many patients have multiple chronic conditions, healthcare systems that are based on first-contact specialist care are likely to result in patients having frequent consultations in different locations or with different providers and often seeing practitioners who have to cross-refer to other specialists for advice about comorbidities. Systems that fragment care across multiple primary care providers face challenges in coordinating care for the many patients with multiple problems.

Continuity of care is particularly important to people with multimorbidity,<sup>33,34</sup> but this study demonstrates that these people are less likely to receive it. Longitudinal continuity, as measured here, is closely related to relational continuity,<sup>34</sup> and although better coordination between providers and shared records may enhance 'management continuity' and 'information continuity',<sup>35</sup> it should not be assumed that these can substitute for the relationship with a particular doctor that many people with multimorbidity value.<sup>33,34,36</sup>

Multimorbidity should be considered as a possible confounding factor in studies comparing the process or outcomes of care in different settings. Although multimorbidity is very common, it is not known how best to organise health services in order to optimise care for these people. A better understanding of the problems that multimorbidity generates is needed to develop and test interventions to improve care.

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#### Ethics committee

Studies based on the GPRD are covered by ethics approval granted by Trent Multicentre Research Ethics Committee, reference 05/MRE04/87.

#### Competing interests

The authors have stated that there are none.

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#### Discuss this article

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**Appendix 1. Expanded diagnostic clusters<sup>a</sup> within the ACG<sup>®</sup> system, designated by the authors as chronic conditions.**

Expanded diagnostic cluster code	Label
Not an EDC. Itemised separately within the ACG software	Asthma
Not an EDC. Itemised separately within the ACG software	Hypertension
Not an EDC. Itemised separately within the ACG software	Diabetes
Not an EDC. Itemised separately within the ACG software	Arthritis
ADM02	Surgical aftercare (e.g. heart valve replacement, colostomy)
ADM03	Transplant status (e.g. heart transplant, bone marrow transplant)
ALL06	Disorders of the immune system
CAR03	Ischemic heart disease (excluding acute myocardial infarction)
CAR04	Congenital heart disease
CAR05	Congestive heart failure
CAR06	Cardiac valve disorders
CAR07	Cardiomyopathy
CAR09	Cardiac arrhythmia
CAR10	Generalised atherosclerosis
CAR11	Disorders of lipid metabolism
CAR12	Acute myocardial infarction
CAR16	Cardiovascular disorders, other (e.g. sub-acute bacterial endocarditis)
EAR08	Deafness, hearing loss
END02	Osteoporosis
END04	Thyroid disease
END05	Other endocrine disorders (including diabetes insipidus)
EYE02	Blindness
EYE03	Retinal disorders (excluding diabetic retinopathy)
EYE06	Cataract, aphakia
EYE08	Glaucoma
EYE13	Diabetic retinopathy
FRE03	Endometriosis
FRE12	Utero-vaginal prolapse
GAS02	Inflammatory bowel disease
GAS05	Chronic liver disease
GAS08	Gastro-oesophageal reflux
GAS09	Irritable bowel syndrome
GAS10	Diverticular disease colon
GAS12	Chronic pancreatitis
GAS13	Lactose intolerance
GSU06	Chronic cystic disease of the breast
GSU08	Varicose veins of lower extremities
GSU11	Peripheral vascular disease
GSU13	Aortic aneurysm
GTC01	Chromosomal anomalies



**Appendix 1 continued. Expanded diagnostic clusters<sup>a</sup> within the ACG<sup>®</sup> system, designated by the authors as chronic conditions.**

Expanded Diagnostic Cluster code	Label
GTC02	Inherited metabolic disorders
GUR01	Vesicoureteral reflux
GUR03	Hypospadias, other penile anomalies
GUR04	Prostatic hypertrophy
GUR09	Renal calculi
GUR10	Prostatitis
HEM01	Hemolytic anaemia
HEM02	Iron deficiency, other deficiency anaemias
HEM05	Aplastic anaemia
HEM03 or HEM 06	Thromboplebitis and DVT (deep vein thrombosis)
HEM07	Hemophilia, coagulation disorder
HEM08	Hematologic disorders, other (including secondary polycythaemia)
INF01	Tuberculosis
INF04	HIV, AIDS
MAL01	Malignant neoplasms of the skin
MAL02	Low impact malignant neoplasms
MAL03	High impact malignant neoplasms
MAL04	Malignant neoplasms, breast
MAL05	Malignant neoplasms, cervix, uterus
MAL06	Malignant neoplasms, ovary
MAL07	Malignant neoplasms, esophagus
MAL08	Malignant neoplasms, kidney
MAL09	Malignant neoplasms, liver and biliary tract
MAL10	Malignant neoplasms, lung
MAL11	Malignant neoplasms, lymphomas
MAL12	Malignant neoplasms, colorectal
MAL13	Malignant neoplasms, pancreas
MAL14	Malignant neoplasms, prostate
MAL15	Malignant neoplasms, stomach
MAL16	Acute leukemia
MAL18	Malignant neoplasms, bladder
MUS06	Kyphoscoliosis
MUS07	Congenital hip dislocation
MUS11	Congenital anomalies of limbs, hands, and feet
MUS13	Cervical pain syndromes
MUS14	Low back pain
NUR03	Peripheral neuropathy, neuritis
NUR05	Cerebrovascular disease
NUR06	Parkinson's disease
NUR07	Seizure disorder
NUR08	Multiple sclerosis
NUR09	Muscular dystrophy
NUR11	Dementia and delirium
NUR12	Quadriplegia and paraplegia
NUR16	Spinal cord injury/disorders

**Appendix 1 continued. Expanded diagnostic clusters<sup>a</sup> within the ACG<sup>®</sup> system, designated by the authors as chronic conditions.**

Expanded Diagnostic Cluster code	Label
NUR17	Paralytic syndromes, other
NUR18	Cerebral palsy
NUR19	Developmental disorder
NUR21	Neurologic disorders, other (e.g. Huntingdon's chorea)
NUT03	Obesity
PSY01	Anxiety, neuroses
PSY02	Substance use
PSY04	Behaviour problems
PSY05	Attention deficit disorder
PSY07	Schizophrenia and affective psychosis
PSY08	Personality disorders
PSY09	Depression
REC01	Cleft lip and palate
REC03	Chronic ulcer of the skin
REN01	Chronic renal failure
REN04	Nephritis, nephrosis
REN05	Renal disorders, other
RES03	Cystic fibrosis
RES04	Emphysema, chronic bronchitis, COPD (chronic obstructive pulmonary disease)
RES06	Sleep apnea
RES08	Pulmonary embolism
RES09	Tracheostomy
RES11	Respiratory disorders, other (e.g. pneumoconiosis)
RHU01	Autoimmune and connective tissue diseases
RHU02	Gout
RHU03	Arthropathy (e.g. pyogenic arthritis)
SKN02	Dermatitis and eczema
SKN12	Psoriasis
SKN13	Disorders of hair and follicles (e.g. alopecia)

<sup>a</sup>Details of expanded diagnostic clusters (EDCs) obtained from the Johns Hopkins University Adjusted Clinical Groups (ACG) Case-Mix System Reference Manual version 7.<sup>17</sup> Bloomberg: John Hopkins University, 2005. Copyright, used by permission.