Evaluating meta-ethnography: systematic analysis and synthesis of qualitative research

R Campbell, P Pound, M Morgan, G Daker-White, N Britten, R Pill, L Yardley, C Pope and J Donovan

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Evaluating meta-ethnography: systematic analysis and synthesis of qualitative research

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Abstract

Evaluating meta-ethnography: systematic analysis and synthesis of qualitative research

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Background: Methods for reviewing and synthesising findings from quantitative research studies in health care are well established. Although there is recognition of the need for qualitative research to be brought into the evidence base, there is no consensus about how this should be done and the methods for synthesising qualitative research are at a relatively early stage of development.

Objective: To evaluate meta-ethnography as a method for synthesising qualitative research studies in health and health care.

Methods: Two full syntheses of qualitative research studies were conducted between April 2002 and September 2004 using meta-ethnography: (1) studies of medicine-taking and (2) studies exploring patients’ experiences of living with rheumatoid arthritis. Potentially relevant studies identified in multiple literature searches conducted in July and August 2002 (electronically and by hand) were appraised using a modified version of the Critical Appraisal Skills Programme questions for understanding qualitative research. Candidate papers were excluded on grounds of lack of relevance to the aims of the synthesis or because the work failed to employ qualitative methods of data collection and analysis.

Results: Thirty-eight studies were entered into the medicine-taking synthesis, one of which did not contribute to the final synthesis. The synthesis revealed a general caution about taking medicine, and that the practice of lay testing of medicines was widespread. People were found to take their medicine passively or actively or to reject it outright. Some, in particular clinical areas, were coerced into taking it. Those who actively accepted their medicine often modified the regimen prescribed by a doctor, without the doctor’s knowledge. The synthesis concluded that people often do not take their medicines as prescribed because of concern about the medicines themselves. ‘Resistance’ emerged from the synthesis as a concept that best encapsulated the lay response to prescribed medicines. It was suggested that a policy focus should be on the problems associated with the medicines themselves and on evaluating the effectiveness of alternative treatments that some people use in preference to prescribed medicines. The synthesis of studies of lay experiences of living with rheumatoid arthritis began with 29 papers. Four could not be
synthesised, leaving 25 papers (describing 22 studies) contributing to the final synthesis. Most of the papers were concerned with the everyday experience of living with rheumatoid arthritis. This synthesis did not produce significant new insights, probably because the early papers in the area were substantial and theoretically rich, and later papers were mostly confirmatory. In both topic areas, only a minority of the studies included in the syntheses were found to have referenced each other, suggesting that unnecessary replication had occurred.

**Limitations:** We only evaluated meta-ethnography as a method for synthesising qualitative research, but there are other methods being employed. Further research is required to investigate how different methods of qualitative synthesis influence the outcome of the synthesis.

**Conclusions:** Meta-ethnography is an effective method for synthesising qualitative research. The process of reciprocally translating the findings from each individual study into those from all the other studies in the synthesis, if applied rigorously, ensures that qualitative data can be combined. Following this essential process, the synthesis can then be expressed as a ‘line of argument’ that can be presented as text and in summary tables and diagrams or models. Meta-ethnography can produce significant new insights, but not all meta-ethnographic syntheses do so. Instead, some will identify fields in which saturation has been reached and in which no theoretical development has taken place for some time. Both outcomes are helpful in either moving research forward or avoiding wasted resources. Meta-ethnography is a highly interpretative method requiring considerable immersion in the individual studies to achieve a synthesis. It places substantial demands upon the synthesiser and requires a high degree of qualitative research skill. Meta-ethnography has great potential as a method of synthesis in qualitative health technology assessment but it is still evolving and cannot, at present, be regarded as a standardised approach capable of application in a routinised way.

**Funding:** Funding for this study was provided by the Health Technology Assessment programme of the National Institute for Health Research.
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# List of abbreviations

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<tr>
<td>ADR</td>
<td>adverse drug reactions</td>
</tr>
<tr>
<td>AZT</td>
<td>azidothymidine (zidovudine)</td>
</tr>
<tr>
<td>CASP</td>
<td>Critical Appraisal Skills Programme</td>
</tr>
<tr>
<td>CINAHL</td>
<td>Cumulative Index to Nursing and Allied Health Literature</td>
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<tr>
<td>CIS</td>
<td>critical interpretive synthesis</td>
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<tr>
<td>ESRC</td>
<td>Economic and Social Research Council</td>
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<tr>
<td>HIV</td>
<td>human immunodeficiency virus</td>
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<tr>
<td>HTA</td>
<td>health technology assessment</td>
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<tr>
<td>NatCen</td>
<td>National Centre for Social Research</td>
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<tr>
<td>PPI</td>
<td>proton pump inhibitor</td>
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<tr>
<td>RA</td>
<td>rheumatoid arthritis</td>
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<tr>
<td>RCT</td>
<td>randomised controlled trial</td>
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<td>WHO</td>
<td>World Health Organization</td>
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All abbreviations that have been used in this report are listed here unless the abbreviation is well known (e.g. NHS), or it has been used only once, or it is a non-standard abbreviation used only in figures/tables/appendices, in which case the abbreviation is defined in the figure legend or in the notes at the end of the table.
Executive summary

Background

Methods for reviewing and synthesising findings from quantitative research studies in health care are well established. Although there is recognition of the need for qualitative research to be brought into the evidence base, there is no consensus about how this should be done and methods for synthesising qualitative research are at a relatively early stage of development.

Aim

Our aim was to undertake methodological research to evaluate meta-ethnography as a method for synthesising qualitative research studies in health and health care.

Methods of synthesis for qualitative research

A review of methods of qualitative synthesis was undertaken to examine the ways in which meta-ethnography, first described by Noblit and Hare (Noblit G, Hare R. *Meta-ethnography: synthesising qualitative studies*. 11th edn. London: Sage Publications; 1988), was being used, and to identify what other methods of qualitative synthesis were available. A range of methods for synthesising qualitative research was identified; none has become established, but meta-ethnography was the most widely cited method. Methods of qualitative synthesis could be broadly categorised as integrative or interpretative. In integrative syntheses, data in the primary studies are considered comparable and can therefore be pooled or aggregated. Methods falling into this category include numeric methods that involve the systematic pooling of qualitative data as a precursor to a quantitative analysis, narrative methods and cross-case techniques. Interpretative syntheses entail an emic approach with concepts and explanatory frameworks emerging through a process of induction. Meta-ethnography and grounded theory are examples of methods included in this second category.

Meta-ethnography has been applied in ways consistent with the approach originally described by Noblit and Hare (1988), but it has also been used as a procedure within an all-embracing form of synthesis called meta-study. Critical interpretative synthesis has also been proposed as a new method which evolved from an attempt to use meta-ethnography to bring together findings from a large and methodologically diverse group of studies.

Identification and selection of studies for synthesis

For this research, two full-qualitative syntheses were conducted using meta-ethnography: the first a synthesis of qualitative studies of medicine-taking and the second a synthesis of studies exploring patients’ experiences of living with rheumatoid arthritis (RA). Searching the literature to identify studies for possible inclusion in a qualitative synthesis remains problematic. Our findings suggest that multiple search strategies should be employed and that hand-searching key journals is particularly worthwhile. The yield of relevant papers produced by electronic databases appeared to be topic dependent, suggesting that a variety should be searched. Similarly, the effectiveness of the electronic search strategy depended on the subject of the search, making it advisable to try more than one approach. Potentially relevant studies identified in literature searches, conducted in July and August 2002, were appraised using a modified version of the
Critical Appraisal Skills Programme [Critical Appraisal Skills Program. (CASP) collaboration for qualitative methodologies. 1998 URL: www.casp-uk.net] questions for understanding qualitative research. Candidate papers for both of our syntheses were excluded only on the grounds of lack of relevance to the aims of the synthesis or because the work failed to employ qualitative methods of data collection and analysis. Papers were not excluded on quality grounds alone. Quantifiable data from the appraisal process showed inter-rater agreement to be low.

Reproducibility

Before proceeding with the full syntheses, the first four papers to be synthesised in each of the two topic areas were independently synthesised by two researchers to explore the reproducibility of the meta-ethnographic method. The findings from these two preliminary syntheses produced mixed evidence of reproducibility. There were broad similarities in interpretation, but differences of detail. It is not possible to predict whether the differences would have become more or less pronounced as more papers were drawn into the syntheses.

Medicine-taking synthesis

Thirty-eight studies were entered into the synthesis, one of which did not contribute to the final synthesis. Most of the papers were about medicine-taking for chronic illness. Studies were initially synthesised by reciprocal translation within groups of papers defined by the type of medication. This process produced diagrams and textual summaries of medicine-taking within each group. The textual summaries were then synthesised thematically across the medicine groups as a lines-of-argument synthesis and the models were synthesised to produce a general model of medicine-taking. The synthesis revealed a general caution about taking medicine, and that the practice of lay testing of medicines was widespread. People were found to take their medicine passively or actively or to reject it outright. Some, in particular clinical areas, were coerced into taking it. Those who actively accepted their medicine often modified the regimen prescribed by a doctor, without the doctor’s knowledge. This synthesis concluded that people do not take their medicines as prescribed because of concerns about the medicines themselves. ‘Resistance’ emerged from the synthesis as a concept that best encapsulates the lay response to prescribed medicines. It was suggested that a future policy focus should be on the problems associated with the medicines themselves and on evaluating the effectiveness of alternative treatments that some people use in preference to prescribed medicines.

Synthesis of studies of lay experiences of living with rheumatoid arthritis

This synthesis began with 29 papers. Four of these could not be synthesised, leaving 25 papers (describing 22 studies) contributing to the final synthesis. Most of the papers were concerned with the everyday experience of living with RA. Papers were synthesised chronologically. The earliest paper in the series was used as an index paper and each subsequent paper entered into the synthesis was compared with it and all preceding papers. The smaller number and coherence of papers in this synthesis permitted the reciprocal translation of findings for all papers to be tabulated. These reciprocal translations were then subject to a process of reordering and reanalysis. The final synthesis was presented as a textual distillation of the findings supported by novel tabular summaries of the needs of people with RA and the general and specific coping strategies that they deployed to accommodate the disease. This synthesis did not produce significant new insights, probably because the early papers in the area were substantial and theoretically rich, and later papers were mostly confirmatory.
Conclusions

This methodological research, conducted between April 2002 and September 2004, has shown that meta-ethnography is an effective method for synthesising qualitative research. The meta-ethnographic method enables a body of qualitative research to be drawn together in a systematic way. The process of reciprocally translating the findings from each study into those from all the other studies in the synthesis, if applied rigorously, ensures that qualitative data can be combined. Following this essential process, the synthesis can then be expressed as a 'line of argument' that can be presented as text and in summary tables and diagrams or models. Meta-ethnography can produce significant new insights, as was achieved in the medicine-taking synthesis. However, as the synthesis of studies about lay experiences of arthritis showed, not all meta-ethnographic syntheses will lead to new insights. This does not signal failure, however, because, as our RA synthesis showed, the meta-ethnographic method is able to identify fields in which saturation has been reached and in which no theoretical development has taken place for some time, an outcome that can ensure that resources are not wasted in future. Both of our syntheses found that only a minority of the studies included referenced each other, suggesting that unnecessary replication occurred. In addition to being able to demonstrate what the cumulative knowledge is in a particular area, meta-ethnographic syntheses are also able to identify absences of knowledge and can reveal aspects of a body of literature that may have been obscured.

Meta-ethnography as described by Noblit and Hare (1988) is a highly interpretative method, and considerable immersion in the individual studies is necessary to achieve a synthesis. It is a method that places substantial demands upon the synthesiser and requires a high degree of qualitative research skill. Experience to date suggests that the inclusion of > ~40 papers would result in a trade-off between the breadth of included papers and the depth of scrutiny and thought invested in each stage of the synthesis. Meta-ethnography has great potential as a method of synthesis in qualitative health technology assessment, but it is still evolving and cannot, at present, be regarded as a standardised approach capable of application in a routinised way.

Funding

Funding for this study was provided by the Health Technology Assessment programme of the National Institute for Health Research.
Chapter 1

Introduction

Although it is widely acknowledged that science is cumulative, people have only very recently begun to acknowledge that scientists have a responsibility to cumulate scientifically.

Chalmers et al.1

Evidence-based medicine and a science of synthesis

Recognition of the need for research syntheses has existed for over two centuries,1 but a ‘science of synthesis’ has been slow to advance. In the field of health care, much of the momentum for the development of methods of synthesis has come from the move towards evidence-based medicine: a shift that has had a variety of antecedents. One was apprehension about the rapidity of technological change and the impossibility of individual clinicians reading and appraising all the journal articles relevant to their specialty.2 Another factor was unease about the quality of primary research testing the effectiveness of clinical and health-care interventions.3 Foremost, however, was recognition that all the best available scientific evidence should primarily inform clinical decision-making and that changes in medical practice should not simply rely on expert opinion.

Narrative literature reviews have been the traditional method for bringing together existing knowledge in a particular area. From the mid-1980s onwards, however, a number of questions about the adequacy of this method for compiling research evidence on health-care interventions were raised. The narrative review was criticised for being unscientific and, as a consequence, it was suggested ‘the results and conclusions would often be susceptible to and reflect the biases of the reviewer’.4 In addition, seminal studies such as that by Antman et al.5 provided powerful demonstrations of the value of a systematic approach to all the components of a review including the selection of studies for inclusion, the appraisal of the quality of the research and the application of relevant statistical methods for pooling data from a number of studies. Using meta-analysis to synthesise randomised controlled trial (RCT) evidence about the effectiveness of treatments in preventing death following heart attack, and contrasting this with expert recommendations in textbooks and review articles, these authors showed that some treatments were still being promoted when there was clear evidence that they were harmful, and that thrombolytic drugs were still not being recommended for use by more than half of clinical experts 13 years after cumulative meta-analyses showed them to be highly effective in reducing mortality.5

In the late 1980s, international collaborations of researchers, responding both to Archie Cochrane’s6 championing of the value of the RCT in health-care evaluation and to his stinging criticism of health-care professionals’ failure to use scientific evidence as the primary basis for making choices about which preventive measures, diagnostic tests and treatments to use, began to publish the first systematic reviews of health care. Foremost among these was a two-volume publication containing hundreds of reviews about different aspects of the care of women and infants during pregnancy and childbirth.7 There were two notable features of this review. Firstly, it was acknowledged that there would be different audiences for such reviews, and so a much shorter, more accessible, summary was also published.8 Secondly, it was recognised that such a
review would need to be regularly updated and that electronic publishing could facilitate this requirement. Rapid adoption of systematic reviews followed this pioneering work and global systematic reviewing networks have gradually developed, with the Cochrane Collaboration being the best known.

**Health technology assessment**

By the late 1990s, with the systematic review firmly established as a scientific method for bringing together findings from quantitative studies of effectiveness, interest was beginning to shift towards how qualitative data could be brought into the evidence base and a 'science of synthesis' developed for qualitative research. Whereas bringing together research information on effectiveness, costs and acceptability has always been fundamental to the health technology assessment (HTA) enterprise, the use of qualitative research methods within HTA has been a more recent development. In the mid-1990s, the HTA Methodology programme commissioned a review of the literature on the use of qualitative research methods in HTA, which noted that application of conventional systematic review methodology to qualitative research presented both philosophical and practical challenges. Following publication of this report, the HTA recognised that further work was required to enable 'users of qualitative research to be able to both appraise and to synthesise qualitative studies in a rigorous, replicable and formalised way'.

**Synthesis of qualitative research**

The possibility of synthesising qualitative research immediately raises a range of important epistemological, methodological and practical questions. The most fundamental of these is what 'synthesis' means in this context. Strike and Posner have suggested that 'synthesis is usually held to be activity or the product of activity where some set of parts is combined or integrated into a whole... [Synthesis] involves some degree of conceptual innovation, or the invention or employment of concepts not found in the characterization of the parts as means of creating the whole.' Thus, synthesis of qualitative research could be envisaged as the bringing together of findings on a chosen theme, the results of which should, in conceptual terms, be greater than the sum of parts. This implies that qualitative synthesis would go beyond the description and summarising usually associated with a narrative literature review, as it would involve conceptual development and would be distinct from a quantitative meta-analysis in that it would not simply entail the aggregation of findings from individual, high-quality research studies. Qualitative synthesis should involve reinterpretation but, unlike secondary analysis, it would be based on published findings rather than primary data.

The impetus for developing methods of qualitative synthesis has arisen from recognition of the importance of qualitative evidence in complementing quantitative research and in particular its ability to provide a more complete understanding of phenomena, especially in terms of processes involved in the organisation and provision of services and influences on behaviours. Another driver has been acknowledgement that the considerable expansion of the qualitative research literature over recent years has produced little accumulated understanding, highlighting a need to bring together isolated studies. Thus, if qualitative research is to be made accessible to health policy-makers and planners, in a manageable form, then developing and evaluating methods of qualitative synthesis, as in this project, would seem to be an important and timely endeavour.

Despite recognition of the potential value of qualitative synthesis, it is a contentious enterprise. From a social science perspective, the notion of qualitative research synthesis best agrees with what is termed a 'subtle realist' position. This stance maintains that phenomena exist
independently of the investigators' claims about them, but also acknowledges the possibility of multiple, non-competing valid descriptions and explanations of the same phenomena. Subtle realism, therefore, enables the different constructions people make of reality to be studied, without accepting that particular beliefs are true; thus, qualitative synthesis can be regarded as potentially promoting such understanding. By contrast, an extreme relativist or radical constructionist perspective is based on a belief that reality is only what we make it and it is therefore not possible to have any knowledge of phenomena apart from our own experience of them. The aim of qualitative research arising from within this influential tradition is to describe unique particularities and present an ideographic account. The process of synthesis could thus be regarded as destroying the integrity of individual studies in the pursuit of some unattainable more 'complete' or 'true' account. As Sandelowski et al. explained, from this perspective the summary of qualitative findings is regarded as thinning out the desired thickness of particulars, which may therefore 'lose the vitality, viscerality, and vicariism (sic) of the human experiences represented in the original studies'. From a radical constructionist perspective, the aim of qualitative research is to produce fit descriptions of unique cases and leave it to the reader to engage in 'naturalistic generalisation' using these individual descriptions. However, as Hammersley observed, in practice it is rare for researchers not to hint at general conclusions from the unique cases that they study.

There are no standard or agreed methods for conducting syntheses of qualitative research. The review presented in Chapter 2 of this report illustrates a number of possible methods that have been identified and applied. Meta-ethnography, developed and used first in educational research, is one approach. It involves taking relevant empirical studies to be synthesised, reading them repeatedly and noting down key concepts (interpretive metaphors). These key concepts are the raw data for the synthesis. Noblit and Hare suggested that the process by which a synthesis is achieved is one of translation. This entails examining the key concepts in relation to others in the original study and across studies. The way of translating key concepts or interpretive metaphors from one study to another involves an idiomatic rather than a word-for-word translation. The purpose of the translation is to try to derive concepts that can encompass more than one of the studies being synthesised. The synthesised concepts may not have been explicitly identified in any of the original empirical studies. As perhaps the best developed method for synthesising qualitative data, and one which clearly had its origins in the interpretive paradigm from which most methods of primary qualitative research evolved, this was the method selected for evaluation in this HTA project.

Prior to the research reported on here, this research group had undertaken a pilot synthesis of four papers concerned with the lay meanings of medicines, which demonstrated that it was possible to use meta-ethnography to synthesise qualitative research. A feasibility study followed that included the formative evaluation of a set of criteria for assessing qualitative research studies followed by a synthesis of selected studies on patient experiences of diabetes mellitus. This second pilot synthesis proved to be successful and illuminating; as none of the research papers included it in contained any references to each other, it demonstrated the need for primary researchers to search and read the existing literature more carefully and indicated that it would be helpful to bring the findings of qualitative research together. This second synthesis also confirmed the effectiveness of meta-ethnography as a method of synthesis. In addition, a practical method of qualitative research assessment and data extraction evolved from it. However, this process required further testing and evaluation before it could be recommended for widespread adoption by those undertaking HTA, as a number of important questions remained to be answered, for example, in terms of the selection of studies, how many studies can be included and how reproducible syntheses are. It is this more detailed evaluation that is the subject of this report.
Aims and objectives

Aim

To appraise and synthesise qualitative health research for HTA using a meta-ethnographic approach.

Objectives

Primary

1. To conduct, using the meta-ethnographic method, syntheses of qualitative research studies in two applied health-care contexts:
   (a) living with rheumatoid arthritis (RA) (patient experiences of a chronic illness)
   (b) lay beliefs about medicine-taking in chronic disease.
2. To test a modified version of the Critical Appraisal Skills Programme (CASP)\(^1^8\) criteria for appraising qualitative research for appropriateness, ease of use and inter-reviewer agreement.
3. To evaluate the reproducibility of the meta-ethnographic method of synthesis.

Secondary

4. To complete a review of the methods available for appraising and synthesising qualitative research.
5. To document the effectiveness of different elements in the search strategies in identifying relevant qualitative research studies.

Content of this report

This report continues in Chapter 2 with a review of the methods of qualitative synthesis. We did not complete a review of methods of appraisal as two other projects commissioned by other bodies at a similar time had such a review within their remit.\(^1^9,2^0\) To avoid unnecessary duplication, we focused our efforts on reviewing methods for the synthesis of qualitative research. Chapter 3 describes the literature searching strategies employed in the two meta-ethnographic syntheses undertaken. An evaluation of the methods of appraisal used in the syntheses is reported in Chapter 4. The precise methods used to conduct the syntheses are described in Chapter 5 together with an assessment of the reproducibility of the meta-ethnographic method. The two syntheses are presented in Chapters 6 and 7. Chapter 8 contains a discussion of all the issues raised by this evaluation of meta-ethnography, and a number of conclusions are drawn about its value as a method of qualitative synthesis and its potential role in HTA.
Chapter 2
Methods of qualitative synthesis

The relative inattention towards integrating qualitative findings stands in sharp contrast to the considerable attention given to the development of techniques for conducting syntheses of quantitative research.

Sandelowski et al.13

Introduction

Methods for the synthesis of qualitative research have been slow to develop, as the above quote suggests, but a number have emerged in recent years. These methods are still evolving, but are doing so in the shadow of widely used and well-developed methods for quantitative research synthesis. The result is that there is sometimes an expectation or assumption that qualitative syntheses will proceed in a manner similar to their quantitative counterparts, despite the very obvious differences between the quantitative and qualitative research traditions. This chapter begins by considering the distinctions between different types of research syntheses before going on to review methods of qualitative synthesis.

One of the confusing features of the research synthesis literature is that the terms ‘review’ and ‘synthesis’ are often used interchangeably, yet a distinction can be made between these two activities. A process of review can be said to be one of seeking out, sifting through, reading, appraising and describing relevant research evidence. The synthesis of evidence, on the other hand, involves a process of extracting data from individual research studies and interpreting and representing them in a collective form. According to this distinction, conventional literature reviews are generally reviews, whereas systematic reviews usually combine elements of review and synthesis.

Within systematic reviews, Hammersley14 identified three types of synthesis: aggregative (involving the accumulation and generalisation of evidence); comparative or replicative (determining the extent to which different sources of evidence reinforce each other by comparison between sources); and developmental (overarching theory development). Although these have evolved within a quantitative research tradition, Hammersley14 suggested that none of these types of synthesis is incompatible with qualitative synthesis. Hammersley14 also suggested convenience mapping synthesis as a fourth category. This category drew on Howard Becker’s notion of a mosaic of different studies being put together in order to see the bigger picture, with each study providing a context for the next.

Noblit and Hare (p. 15)15 also made a distinction between integrative reviews, in which data from different studies are pooled or aggregated, and interpretive reviews, which bring together the findings from different studies using induction and interpretation to gain deeper understandings of a particular phenomenon. They noted how integrative reviews require an etic approach (working with a pre-existing frame of reference), thereby enabling the aggregation of findings by ensuring that there is ‘a basic comparability between phenomena’.15 Drawing on work by Spicer,22 interpretative reviews were contrastingly characterised as using an emic approach with concepts and explanatory frameworks emerging through a process of induction.
This chapter examines the assumptions and procedures of different methods of synthesising qualitative research drawing on the framework proposed by Hammersley as a means of distinguishing different methods of qualitative synthesis. Particular attention is given to Noblit and Hare’s meta-ethnography, as this forms the main approach to qualitative synthesis employed in the health field and is the focus of this report. Methods for the formal integration and synthesis of quantitative and qualitative studies are beyond the scope of this report and have been considered in detail elsewhere.

Types of qualitative synthesis

Methods of qualitative synthesis are at an early stage of development and considerable methodological work is ongoing, particularly within health and educational research. The rapid expansion of the field has resulted in a lack of standard terminology, with the terms ‘meta-ethnography’, ‘meta-interpretation’, ‘meta-analysis’, ‘narrative synthesis’, ‘meta-synthesis’ and other descriptors being widely used to describe similar approaches. Conversely, the same terms are frequently employed to describe different approaches. This suggests a need to look beyond labels when searching and reviewing this area of work and for developing an agreed terminology.

The term ‘synthesis’ also has varying meanings and fuzzy boundaries in relation to qualitative research and involves a range of activities. At present, there is no single agreed classification of different types of qualitative synthesis and different authors adopt varying frameworks. Based on their main aims and approaches to synthesis, this chapter identifies the different types of synthesis that may be undertaken in relation to qualitative studies (i.e. secondary data) as numeric, narrative and interpretive (Box 1). In practice, however, these different methods may overlap and can be viewed as forming part of a continuum that ranges from numeric syntheses at one end to interpretive approaches based on a qualitative paradigm at the other. Some

<table>
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<tr>
<th>Secondary data</th>
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<tbody>
<tr>
<td>Numeric synthesis</td>
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<td>Case survey method</td>
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<td>Bayesian methods</td>
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<td>Narrative synthesis</td>
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<td>Thematic analysis</td>
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<td>Cross-case analysis</td>
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<td>Interpretive synthesis</td>
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<td>Meta-ethnography</td>
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<td>Meta-study</td>
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<td>Grounded theory</td>
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<td>Realist synthesis</td>
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<tr>
<td>Primary data</td>
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<tr>
<td>Secondary data analysis</td>
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forms of secondary analysis of primary data across studies can also be regarded as a form of qualitative synthesis.

**Aggregative synthesis**

It seems paradoxical to begin this review of methods for synthesising qualitative research with numeric approaches. Nevertheless, a variety of methods have been proposed for converting data from qualitative research studies into a quantitative form for the purposes of analysis and synthesis. An early example is Yin's case survey method, which was developed for the synthesis of case studies based on a qualitative or mixed-method approach. This method involves the initial application of a closed-ended coding instrument to each qualitative case study to abstract and record relevant data. The collective data from these codes are then tallied and analysed in much the same way as traditional survey data. Examples are analyses by Yin and colleagues of citizen participation in urban services and of innovations in urban services. This approach to the synthesis of qualitative data mimics quantitative meta-analysis in which findings (usually from individual RCTs) are aggregated in order to have sufficient statistical power to detect a cause and effect relationship between a particular treatment and specific health outcomes.

Current interest in combining qualitative and quantitative syntheses has led to other types of numeric methods, with a key example being the Bayesian synthesis of qualitative and quantitative evidence. This involves pooling of the data and is illustrated by Roberts et al.'s investigation of childhood immunisation and the probable effects of a variety of factors on uptake. Their approach involved the initial extraction of factors influencing the uptake of childhood immunisation identified from qualitative studies. These factors were then classified in terms of broader descriptive categories (e.g. 'lay beliefs' was used to capture such concepts as 'parents' beliefs' and 'opinions of parents and other family members about the value of immunisation'). Rankings of the relative importance of each factor by each reviewer yielded a probability of that factor being important in determining uptake. These factors were then combined with evidence from the quantitative data to form a posterior probability that each factor identified (e.g. lay beliefs) was important in determining uptake of immunisation. This synthesis, therefore, provided a more complete enumeration of factors associated with the uptake of immunisation through combining quantitative and qualitative studies. Bayes factor methods were then employed to compare two possible meta-regression models in explaining the log-odds of immunisation uptake for each possible factor. This approach forms an important attempt to combine qualitative and quantitative evidence using numeric methods and pooling data, but like other numeric methods is mainly concerned with what Hammersley identified as aggregative purposes in terms of the accumulation and generalisation of evidence.

Other approaches to the systematic analysis/synthesis of studies derive from Miles and Huberman’s cross-case techniques, which involve developing summary tables based on content analysis and noting commonalities and differences between studies.

**Narrative synthesis**

The aims of narrative synthesis are generally to achieve an aggregation of findings or check the comparability and replication of findings, based on narrative rather than numeric methods. A narrative synthesis may also lead to new insights and form the basis for further interpretive analysis and conceptual development, although this is not the primary focus.

There are two broad approaches to qualitative narrative synthesis. One is based on the traditional literature review and involves the use of informal but critical and reflective methods. This approach, lately described as ‘narrative review’, unusually aims to provide a commentary and summary of the main features and findings of a body of literature using methods that are not explicitly pre-defined and transparent. This method has been distinguished from 'narrative
Methods of qualitative synthesis

Narrative synthesis is an approach that has evolved largely from within the quantitative systematic reviewing tradition. Indeed, the Centre for Reviews and Dissemination at York University now encourages reviewers to first undertake a ‘descriptive or non-quantitative synthesis’ of studies to be included in a review, in other words a narrative synthesis, to help them think about what the ultimate method of synthesis should be. In order to further assist this development, a report has recently been compiled to provide detailed guidance on how to undertake narrative synthesis in systematic reviews.

**Interpretive synthesis**

This is generally regarded as ‘the’ approach to the synthesis of qualitative research, and has as its specific aim the achievement of the developmental goal of qualitative synthesis in terms of producing interpretations that go beyond individual studies and thus contribute to conceptual and theoretical development in the field and accords with Strike and Posner’s notion of qualitative synthesis as achieving ‘conceptual innovation’ or the ‘invention or employment of concepts not found in the characterization of the parts as a means of creating the whole’ (p. 346). The output is, therefore, a new interpretation or theory that goes beyond the findings of any individual study.

A number of approaches to the conduct of interpretive synthesis have been developed. These include meta-ethnography, meta-study, critical interpretive synthesis, realist synthesis and grounded formal theory. Each of these approaches is examined in detail in *Approaches to interpretive synthesis*, but a general feature is that the synthesis builds interpretation from original studies by firstly identifying interpretations offered by the original researchers (second-order constructs) and, secondly, enabling the development of new interpretations (third-order constructs) that go beyond those offered in individual primary studies.

**Secondary data analysis**

This approach is concerned with the analysis and interpretation of primary (i.e. ‘raw’) data rather than the findings of published studies. Secondary data analysis may be regarded as a form of interpretive synthesis if it fulfils two requirements: it involves multiple qualitative data sets rather than a single data set; and the aim is to provide a new perspective or conceptual focus rather than to undertake an additional in-depth or subset analysis of existing data.

There are currently only a few examples of secondary analysis of multiple qualitative data sets. These have mainly involved the re-analysis by researchers of their own data. An example is Bloor and Macintosh’s study of techniques of client resistance to surveillance based on two earlier studies they conducted. Interest in the secondary analysis of qualitative data is growing, and the capacity to undertake such work is considerably enhanced by the establishment by the Economic and Social Research Council’s (ESRC’s) UK Data Archive of an archive of data from qualitative studies.

Secondary analysis of qualitative data raises a number of ethical and methodological issues, including issues of consent in relation to the reuse of another researcher’s data for a different
purposes,\textsuperscript{36,37} and methodological questions such as the effects of distance from the data collection and of possible variations in study design on interpretation.\textsuperscript{38} It is likely that the greater availability of qualitative data sets and increasing interest in secondary analysis will lead to new protocols and strategies to address these issues, thus, enhancing this approach to the synthesis of qualitative research.

\section*{Approaches to interpretive synthesis}

This section provides a more detailed consideration of four approaches to interpretive synthesis: meta-ethnography, meta-study, grounded formal theory and realist synthesis.

\subsection*{Meta-ethnography}

Noblit and Hare's\textsuperscript{15} method of meta-ethnography was published in 1988 and is described as 'an attempt to develop an inductive and interpretive form of knowledge synthesis'. Noblit and Hare\textsuperscript{15} developed meta-ethnography in response to the perceived failure of a synthesis of five ethnographic studies of educational desegregation that were undertaken to convey information to policy-makers. This educational synthesis took an aggregative, thematic approach that involved abstracting data and isolating factors in each study that appeared to be responsible for the failure of schools to desegregate. This process of abstraction de-emphasised the uniqueness of each site. The context therefore merely became a confounding variable in the search for common findings rather than contributing to an explanation of these findings. As a result, the synthesis did not provide researchers or policy-makers with an understanding of what went wrong and what could be done about it. Noblit and Hare\textsuperscript{15} aimed to overcome these limitations through developing a distinct method for the synthesis of qualitative studies that was informed by Turner's\textsuperscript{39} theory of social explanation and is interpretive rather than aggregative. As they stated:

\begin{quote}
The nature of interpretive explanation is such that we need to construct an alternative to the aggregative theory of synthesis entailed in integrative research reviews and meta-analysis and be explicit about it.
\end{quote}

Noblit and Hare\textsuperscript{15} (p. 18)

This aim of constructing adequate interpretive explanations required developing a way of 'reducing' and deriving understanding from multiple cases, accounts, narratives or studies while retaining the sense of the account. Noblit and Hare\textsuperscript{15} were themselves ethnographers who were concerned with long-term intensive studies that employed observation, interviews and documents, and termed the approach that they developed 'meta-ethnography'. However, they described meta-ethnography as being applicable to qualitative research generally and as forming 'a rigorous procedure for deriving substantive interpretations about any set of ethnographic or interpretive studies' (p. 9). Noblit and Hare\textsuperscript{15} also noted that their particular approach was 'a' meta-ethnography and that it formed 'but one of many possible approaches' (p. 25).

Noblit and Hare\textsuperscript{15} identified seven phases in undertaking meta-ethnography (\textit{Box 2}), but observed that in practice these phases may occur in parallel and overlap. The phases broadly correspond with other methods of synthesis, but differ in the assumptions and procedures involved. One difference is that the sample for research is purposively selected in relation to the topic of interest (and may involve maximum variation sampling), rather than being exhaustive. This reflects the general approach of qualitative methods and the aim of achieving interpretive explanation. A second difference is that the interpretations and explanations contained in the original studies are treated as data through the selection and analysis of key 'metaphors' (i.e. the themes, perspectives or concepts revealed by qualitative studies), with the aim of reducing
accounts while preserving the sense of the account. Preparation for comparison between studies requires listing and juxtaposing the key metaphors, phases, ideas and/or concepts used in each account but retaining, as far as possible, the terminology used by the authors to remain faithful to the original meanings (phase 4). A third difference is that comparison between studies involves processes of ‘translation’, with the metaphors/concepts and their interrelationships in one account being compared with those in another account. This process of translation is idiomatic and focuses on translating the meaning of the text rather than a literal translation, with the aim of preserving original meanings and contextualisation. Noblit and Hare\textsuperscript{15} identified three possible types of relationship that guide translation and subsequent synthesis:

1. Reciprocal: when studies are about similar things, they can be synthesised as direct translations (i.e. in an iterative fashion each study is translated into the metaphors of the others – see Chapter 5 for a detailed description and illustration of this process). ‘These reciprocal translations may reveal that the metaphors of one study are better than those of others in representing both studies, or that some other set of metaphors not drawn from these studies seems reasonable.’ Noblit and Hare\textsuperscript{15} pointed out that the ‘uniqueness of studies may not make it possible for a single set of metaphors to adequately express the studies’, in which case more is often learned from the process of translation than from the metaphors alone.

2. Refutational: this is undertaken when studies refute each other. It requires a more elaborate set of translations ‘involving’ translations of both the ethnographic account and the refutations to examine the implied relationship between competing explanations.

3. Lines-of-argument: many studies suggest a lines-of-argument or inference about some larger issue or phenomenon. This involves first translating studies into each other and then constructing an interpretation (‘lines-of-argument’) that may serve to reveal what was hidden in individual studies (discovering a ‘whole’ among a set of parts). This is achieved

### BOX 2 Noblit and Hare’s\textsuperscript{15} phases for conducting a meta-ethnography

- **Phase 1: Getting started** – ‘identifying an intellectual interest that qualitative research might inform’. This may be changed/modified as interpretive accounts are read.
- **Phase 2: Describing what is relevant to initial interest** – an exhaustive search for relevant accounts can be undertaken followed by selection of research relevant to the topic of interest (they observe that employing all studies of a particular setting often yields trite conclusions).
- **Phase 3: Reading the studies** – the repeated reading and noting of metaphors is required and continues as the synthesis develops.
- **Phase 4: Determining how the studies are related** – the task of putting together the studies requires creating a list of key metaphors, phrases, ideas or concepts (and their relations) used in each account, and juxtaposing them. This leads to initial assumptions about relations between studies.
- **Phase 5: Translating the studies into one another** – the metaphors and/or concepts in each account and their interactions are compared with the metaphors and/or concepts and their interactions in other accounts. These translations are one level of meta-ethnographic synthesis.
- **Phase 6: Synthesizing translations** – ‘the various translations can be compared with one another to determine if there are types of translation or if some metaphors/concepts are able to encompass those of other accounts. In these cases, a second level of synthesis is possible, analysing types of competing interpretation and translating them into each other’ to produce a new interpretation/conceptual development.
- **Phase 7: Expressing the synthesis** – for the proposed synthesis to be communicated effectively it needs to be expressed in a medium that takes account of the intended audience’s own culture and so uses concepts and language they can understand.
through the use of the comparison and theory generation aspects of grounded theory as described by Glaser and Strauss, and involves the detailed study of differences and similarities among studies to be synthesised with the aim of producing an integrated scheme and new interpretive context.

How translations are synthesised (phase 6), and the product of this process, depends on how studies relate to each other. Both translation and synthesis involve a continuous comparative analysis of texts until a comprehensive understanding of the phenomena is realised and the synthesis is then complete. Noblit and Hare did not specify this process in detail, but regarded it as akin to the general processes of qualitative research. The main difference is that the ‘data’ of meta-ethnography are the substance of several qualitative studies.

The final stage of meta-ethnography is ‘expressing the synthesis’ or communicating with an audience. This was given considerable emphasis by Noblit and Hare, who stated that ‘the worth of any synthesis is in its comprehensibility to some audience’ (p. 82). They described the needs of the audience (e.g. researchers, policy-makers) as influencing both the form and substance of the synthesis. Some understanding of the audience's culture is therefore required to ensure that the translation of studies for the synthesis uses intelligible concepts to inform the final presentation of synthesis. They observed that if the data are inadequate or if the audience cannot see the connection between data and the argument then the study becomes unbelievable. Comprehensibility and believability are thus central to determining worth.

Noblit and Hare's approach to meta-ethnography was based on a literary tradition of interpretivism and was, therefore, driven by a desire to achieve adequate interpretive explanations rather than by technical interests. This means that the authors were critical of the emphasis given by some qualitative researchers to the explicitness of processes employed to analyse data and refer to what Marshall described as the 'bureaucratization of data analysis'. Instead they emphasised the fluidity and interpretive aspects, and described the conduct of meta-ethnography as an ongoing process in which substantive interests may change as the synthesis proceeds. Similarly, translations were regarded as emergent and interactive, with the stages of translation and synthesis often interweaving rather than forming distinct phases.

Noblit and Hare illustrated the methods of meta-ethnography using brief excerpts of material from educational studies. Following publication of their monograph, there has been considerable interest in the application of meta-ethnography to a variety of fields including education, public policy and health. The greatest number of recent meta-ethnographic syntheses have come from nursing, reflecting the large number of qualitative studies in nursing and the particular interests and contributions of schools of nursing in the USA and Canada.

The present review involved a search for examples of meta-ethnography and other methods of interpretive synthesis in the health field. This initially involved a computer search undertaken in spring 2003 and updated in the summer of 2006 using MEDLINE, Cumulative Index to Nursing and Allied Health Literature (CINAHL), EMBASE, Web of Science (Social Science Citation Index and Science Citation Index), PsychINFO and Zetoc (British Library's Electronic Table of Contents) electronic databases. A simple search strategy was used that combined the term ‘qualitative research’ with the terms ‘meta analysis’, ‘meta synthesis’, ‘meta ethnography’, ‘systematic review’, ‘data synthesis’ and ‘meta study’. Hyphenated terms (e.g. meta-ethnography) were also used. Further syntheses were identified via a published textbook of methods of qualitative meta-synthesis and later compared with Finfgeld's review. However, the aim was not so much to achieve an exhaustive list as to identify representative syntheses using meta-ethnography and to examine their practical applications.
A total of 41 reports were identified that appeared to be qualitative syntheses.\textsuperscript{16,17,44–82} The authors used various terms to describe the methods employed and occasionally used more than one term. We therefore broadly grouped syntheses according to what appeared to be their main approach. Syntheses employing meta-ethnography prior to the beginning of the present project demonstrated the potential value of this approach in producing conceptual developments. This is illustrated by the development of a wellness–illness model,\textsuperscript{61} a model of caring,\textsuperscript{74} models of diabetes management\textsuperscript{17,70} and explanations of states and behaviours such as moral distress among nurses, non-adherence among patients, the experience of post partum depression and help-seeking behaviours (\textit{Table 1}). The studies included in these syntheses mainly collected data through interviews with only a few including focus groups or observational methods. They also exhibited a limited variety of theoretical perspectives, reflecting the dominant approaches to qualitative research in the field of health and illness.

Altogether we identified six syntheses that were clearly labelled as employing the methods of meta-ethnography. These included two papers on preliminary work published by the present team;\textsuperscript{16,17} a synthesis of 43 papers concerned with the experience of diabetes;\textsuperscript{70} a meta-ethnographic synthesis by Smith \textit{et al.}\textsuperscript{45} examining cancer patients’ help-seeking experiences;

\begin{table}[h]
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\begin{tabular}{|l|l|l|l|l|}
\hline
\textbf{Authors and title} & \textbf{Number of reports included} & \textbf{Authors’ description} & \textbf{Appraisal criteria} & \textbf{Main concepts/explanations} & \textbf{Comments} \\
\hline
\textbf{Meta-ethnography} & & & & & \\
Britten \textit{et al.}, 2002\textsuperscript{16} & Four papers, arbitrarily chosen (included three by authors) & Use of Noblit and Hare’s\textsuperscript{15} meta-ethnography & No & Developed a line of argument that accounts for patients’ medicine-taking behaviour and communication with health professionals in different settings based on notions of self-regulation and selective disclosure & Aimed to assess benefits of meta-ethnography through a worked example \\
Campbell \textit{et al.}, 2003\textsuperscript{17} & 10 reports, purposively selected & Use of Noblit and Hare’s\textsuperscript{15} meta-ethnography & Questions from CASP\textsuperscript{18} Two papers excluded because of methods and one because of an overlap with another paper & Identified a model of diabetes management involving strategic non-compliance which was associated with being in control of diabetes, ‘coping’, achieving a balance between quality of life and illness, improved glucose levels and a feeling of well-being & Aimed to provide a formative assessment of meta-ethnography and of a qualitative assessment tool \\
Feder \textit{et al.}, 2006\textsuperscript{44} & 29 articles, computer search & Meta-analysis of qualitative studies. Use of Schulz \textit{et al.’s}\textsuperscript{31} framework of constructs for meta-analysis. Meta-ethnography method used by Campbell \textit{et al.},\textsuperscript{17} and first used by Noblit and Hare\textsuperscript{15} & No & Use of first-, second- and third-order constructs to identify desirable characteristics of health-care professionals in consultations in which partner violence is raised & Table of third-order constructs in terms of recommendations to health-care providers by stage of interaction with abused women \\
\hline
\end{tabular}
\caption{Examples of syntheses in the health field that have employed meta-ethnography}
\end{table}
TABLE 1 Examples of syntheses in the health field that have employed meta-ethnography (continued)

<table>
<thead>
<tr>
<th>Authors and title</th>
<th>Number of reports included</th>
<th>Authors’ description</th>
<th>Appraisal criteria</th>
<th>Main concepts/explanations</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smith et al., 2005(^5) Delay in presentation of cancer: a synthesis of qualitative research on cancer patients’ help-seeking experiences</td>
<td>32 papers, computer search</td>
<td>Use of Noblit and Hare’s(^5) meta-ethnography</td>
<td>No</td>
<td>Use of second- and third-order constructs. Similarities in help-seeking in patients with different cancer types. Key concepts were recognition and interpretation of symptoms, and fear of consultation. Fear of embarrassment and fear of cancer. Patient’s gender and the sanctioning of help-seeking were important factors in prompt consultation</td>
<td>Provides international overview through the systematic synthesis of a diverse group of small-scale qualitative studies</td>
</tr>
<tr>
<td>Walter et al., 2004(^46) Lay understanding of familial risk of common chronic diseases: a systematic review and synthesis of qualitative research</td>
<td>11 qualitative articles, computer search and reference lists</td>
<td>Meta-analysis using meta-ethnographic methods and drawing on Schulz et al.(^3) Three stages: first-order constructs (key concepts from each article); second-order constructs (translating first-order constructs); third-order constructs (synthesising second-order constructs to produce overarching concepts)</td>
<td>Articles assessed using appraisal scoring system (CASP). None excluded</td>
<td>Second-order constructs included diseases running in my family; experiencing a relative’s illness; personal mental models; personalising vulnerability; and control of familial risk. Led to three main third-order constructs: salience; personalising process; and personal sense of vulnerability</td>
<td>Identifies third-order constructs for health professionals to explore with patients that may improve the effectiveness of communication about disease risk and management</td>
</tr>
<tr>
<td>Attree, 2005(^47) Parenting support in the context of poverty: a meta-synthesis of the qualitative evidence</td>
<td>12 studies, search methods not given</td>
<td>Use of Noblit and Hare’s(^5) meta-ethnographic methods to produce meta-synthesis of findings</td>
<td>Quality appraised using checklist that drew on earlier models of assessing qualitative research: Popay et al.(^5), Seale,(^44) Mays and Pope,(^48) CASP,(^18) Spencer et al.(^19) Studies graded A–D. A–C grades were included</td>
<td>Systematic review of qualitative studies of low-income parents to explore informal and formal support networks. Two main themes were found: informal and formal support</td>
<td>Difficult to identify any of the key components of a meta-ethnography, e.g. reciprocal translation, refutation or lines-of-argument</td>
</tr>
<tr>
<td>Barroso and Sandelowski, 2004(^48) Substance abuse in HIV-positive women</td>
<td>74 reports, search methods not given</td>
<td>Qualitative meta-synthesis of studies</td>
<td>No</td>
<td>Qualitative meta-synthesis of studies containing information on abuse among HIV-positive women. Three main themes found: diagnosis as a turning point; complications of motherhood for dually diagnosed women; and benefits of recovery – beyond stopping substance abuse</td>
<td>Constructs trajectory that describes events of women’s lives with regard to substance abuse and its intersection with HIV infection</td>
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</tbody>
</table>

a synthesis by Feder et al.\(^44\) of health-care professionals’ interactions with women in violent relationships; and another by Walter et al.\(^46\) which was concerned with lay understandings of familial risk in common cancers.
<table>
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<tbody>
<tr>
<td>Barroso and Powell-Cope, 2000</td>
<td>21 studies, computer search</td>
<td>Meta-synthesis</td>
<td>Burns’60 standards: led to ‘some exclusions’</td>
<td>Identified importance of finding meaning in HIV infection/AIDS which led to being able to establish human connectedness, focusing on the self, negotiating health care and dealing with stigma. Those for whom HIV infection had shattered meaning were unable to establish a framework for coping.</td>
<td>Conducted a post-review validity check on a random sample of a further 33 studies published 1996–8 to determine that their metaphors were still valid in these studies. Many extracts of interviews with HIV-positive people.</td>
</tr>
<tr>
<td>Beck, 2002&lt;sup&gt;24&lt;/sup&gt;</td>
<td>Six qualitative studies, computer search</td>
<td>Use of Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Synthesis produced five themes: bearing the burden; riding the emotional roller coaster; lifesaving support; striving for maternal justice; and acknowledging individuality.</td>
<td>Synthesis conducted to illustrate the method.</td>
</tr>
<tr>
<td>Beck, 2002&lt;sup&gt;23&lt;/sup&gt;</td>
<td>18 studies, computer search</td>
<td>Use of Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Identified four overarching themes with their subsumed metaphors in relation to the experience of post partum depression: incongruity between expectations and reality of motherhood; spiralling downward; pervasive loss; and making gains. Coping involved seeking professional help, adjusting unrealistic expectations and regaining control in their lives.</td>
<td>Described the end product as a comprehensive and thickly descriptive account that is the foundation for theory development.</td>
</tr>
<tr>
<td>Carrol, 2004&lt;sup&gt;22&lt;/sup&gt;</td>
<td>12 studies, computer search</td>
<td>Meta-synthesis following Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Found five overarching themes and divided them into two groups: characteristics of non-vocal ventilated patients’ communication experience; type of nursing care desired by non-vocal patients in order to be understood.</td>
<td>Data translated into second-order interpretations, but not constructed into third-order interpretations.</td>
</tr>
<tr>
<td>Clemmens, 2003&lt;sup&gt;23&lt;/sup&gt;</td>
<td>18 studies, computer search</td>
<td>Meta-synthesis following Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Five metaphors were found: reality of motherhood brings hardship; living in the two worlds of adolescence and motherhood; motherhood as positively transforming; baby as stabilising influence; and supportive context as turning point for future. Clinical implications of study.</td>
<td></td>
</tr>
<tr>
<td>Coffey, 2006&lt;sup&gt;24&lt;/sup&gt;</td>
<td>11 studies, computer search</td>
<td>Meta-synthesis following Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Two of the 11 studies were triangulated. Seven themes were found: living worried; staying in the struggle; carrying the burden; survival as a family; bridge to the outside world; critical times; and taking charge.</td>
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</tbody>
</table>

TABLE 1 Examples of syntheses in the health field that have employed meta-ethnography (continued)
### TABLE 1 Examples of syntheses in the health field that have employed meta-ethnography (continued)

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<tr>
<th>Authors and title</th>
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<tbody>
<tr>
<td>Coffman M, 2004&lt;sup&gt;55&lt;/sup&gt; Cultural caring in nursing practice: a meta-synthesis of qualitative research</td>
<td>13 studies, computer search</td>
<td>Meta synthesis following Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Findings reduced to six overall themes: connecting with the client; cultural discovery; the patient in context; in their world, not mine; roadblocks; and the cultural lens</td>
<td></td>
</tr>
<tr>
<td>Evans and FitzGerald, 2002&lt;sup&gt;56&lt;/sup&gt; The experience of physical restraint: a systematic review of qualitative research</td>
<td>Four qualitative studies, computer search</td>
<td>Using meta-ethnography Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-synthesis Jenson and Allen&lt;sup&gt;13&lt;/sup&gt; content analysis (Suikkala and Leino-Kilpi&lt;sup&gt;87&lt;/sup&gt;). Identify the key themes and compare these across studies</td>
<td>Locally developed tool to appraise: methodology; clear descriptions; adequate result information; information supported by exemplars from study participants</td>
<td>Summarises current evidence on the experience of being physically restrained, and the experience of having a relative subject to restraint in an acute or residential care facility</td>
<td>References a number of approaches to qualitative synthesis, but does not seem to apply them</td>
</tr>
<tr>
<td>Finfgeld-Connett, 2005&lt;sup&gt;57&lt;/sup&gt; Clarification of social support</td>
<td>47 studies, computer search (3 linguistic analyses and 44 qualitative studies)</td>
<td>Meta-synthesis. Use of the Template Verification and Expansion Model (Finfgeld&lt;sup&gt;43&lt;/sup&gt;) of concept development, based on meta-synthesis of findings from qualitative studies and linguistic analyses to inductively clarify and expand existing conceptual models and triangulate findings</td>
<td>No</td>
<td>Aimed to clarify the concept of social support. Findings placed in a matrix organised by Walker and Avant’s&lt;sup&gt;89&lt;/sup&gt; broad concept analysis categories: antecedents, critical attributes and consequences. Three themes found: types of social support; attributes of social support; and antecedents of social support</td>
<td></td>
</tr>
<tr>
<td>Finfgeld-Connett, 2006&lt;sup&gt;58&lt;/sup&gt; Meta-synthesis of presence in nursing</td>
<td>18 studies, computer search (14 qualitative studies and 4 linguistic-concept analyses)</td>
<td>Finfgeld’s&lt;sup&gt;43&lt;/sup&gt; meta-synthesis methods. Grounded theory (Strauss and Corbin&lt;sup&gt;90&lt;/sup&gt;)</td>
<td>No</td>
<td>Presence is an interpersonal process that is characterised by sensitivity, holism, intimacy, vulnerability and adaptation to unique circumstances. Created chart: ‘process of presence’</td>
<td></td>
</tr>
<tr>
<td>Goodman, 2005&lt;sup&gt;59&lt;/sup&gt; Becoming an involved father of an infant</td>
<td>10 articles, computer search</td>
<td>Meta-synthesis using Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography</td>
<td>No</td>
<td>Fathers of infants experienced four phases: entering with expectations and intentions; confronting reality; creating one’s role of involved father; and reaping rewards. Implications for theory development, research and clinical practice are discussed</td>
<td></td>
</tr>
</tbody>
</table>

<sup>continued</sup>
### Methods of qualitative synthesis

**TABLE 1** Examples of syntheses in the health field that have employed meta-ethnography (continued)

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<tr>
<td>Howell and Brazil, 2005&lt;sup&gt;51&lt;/sup&gt; Reaching common ground: a patient-family-based conceptual framework of quality EOL care</td>
<td>Seven studies, computer search</td>
<td>Descriptive meta-synthesis in systematic review. Linking metaphors in a Venn diagram, Miles and Huberman&lt;sup&gt;30&lt;/sup&gt;</td>
<td>Excluded if inductive reasoning not used/ lack of evidentiary support. Sandelowski and Barroso’s&lt;sup&gt;91&lt;/sup&gt; typology used for classifying studies on the basis of their interpretive analysis to determine their methodological comparability</td>
<td>Found eight themes: pain and symptom control; dying process not prolonged; prepared for death; support of family or friends; supported decision-making; spiritual support and meaning; holistic and individualised care; death in a supportive environment in location of choice</td>
<td>Grounded theory methods were used for developing metaphors and themes in four studies; unspecified content analysis was described in the remaining studies</td>
</tr>
<tr>
<td>Jensen and Allen, 1994&lt;sup&gt;43&lt;/sup&gt; A synthesis of qualitative research on wellness–illness</td>
<td>112 studies, search methods not given (included 63 dissertations and theses)</td>
<td>Use of Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; meta-ethnography and grounded theory</td>
<td>No</td>
<td>Derived overall model of wellness–illness based on lived experience involving abiding vitality (when one is healthy); transitional harmony (experience of unity disrupted by disease); rhythmical connectedness (disease results in detachment or disconnection from oneself, others and environment); unfolding fulfilment; active optimism; and reflective transformation</td>
<td></td>
</tr>
<tr>
<td>Kärkkäinen et al., 2005&lt;sup&gt;52&lt;/sup&gt; Documentation of individualized patient care: a qualitative metasynthesis</td>
<td>14 research reports, computer search</td>
<td>Qualitative meta-synthesis</td>
<td>Data management followed Gadamer’s hermeneutic empirical theory: a new understanding of investigated matter occurs when a concept conveyed by the studied text by means of hermeneutic dialogue is added to the researchers view. Results of analysed articles can be interpreted in a different way</td>
<td>Three themes emerged that affected the content of nursing-care documentation: reflecting the demands on an organisation; reflecting nurses attitudes and duties; and reflecting patients’ involvement in their care</td>
<td></td>
</tr>
<tr>
<td>Kennedy et al., 2003&lt;sup&gt;63&lt;/sup&gt; An exploration meta-synthesis of midwifery practice in the United States</td>
<td>Six studies, search methods not given</td>
<td>Meta-synthesis</td>
<td>No</td>
<td>Four themes identified, conceptually arrayed into a helix model to portray their dynamic and overlapping nature: the term midwife as an instrument of care; the woman as a partner in care; alliance in midwifery care; and the environment in the process of midwifery care</td>
<td>Uses some of the stages in meta-ethnography as described by Noblit and Hare&lt;sup&gt;15&lt;/sup&gt;</td>
</tr>
<tr>
<td>Kyllma, 2005&lt;sup&gt;64&lt;/sup&gt; Despair and hopelessness in the context of HIV infection – a meta-synthesis on qualitative research findings</td>
<td>Five qualitative studies by the same author, computer search</td>
<td>Meta-synthesis method: extraction, comparing extracted factors, coding, gathering categories together</td>
<td>No</td>
<td>Despair consists of two subprocesses – downwards and upwards. Downwards: being stuck in a situation, losing perspective. Upwards: fighting against sinking, rising up</td>
<td>Five studies were all by the same author</td>
</tr>
</tbody>
</table>
The largest group of qualitative syntheses that we identified were labelled as meta-synthesis. Qualitative meta-synthesis was defined by Sandelowski et al. as ‘the theories, grand narratives, generalizations, or interpretative translations produced from the integration or comparisons of qualitative synthesis’. Meta-synthesis, thus, seems to be, in effect, another term for qualitative synthesis and one that is not identified with any particular method. Many of the syntheses included within this category cited Noblit and Hare and claimed to be using meta-ethnographic methods, but what seemed to distinguish this group of studies from those labelling themselves

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<tr>
<td>Kylma, 2005</td>
<td>Five articles by the same author, computer search</td>
<td>Meta-synthesis: read articles, extraction, comparing extracted factors, coding, gathering categories together</td>
<td>No</td>
<td>Dynamically altering balance between interconnected hope, despair and hopelessness based on folding and unfolding possibilities with regard to the dynamics of hope in dealing with the changing self and life with HIV infection/AIDS</td>
<td>Five grounded theory studies – all by the same author</td>
</tr>
<tr>
<td>Lefler and Bondy, 2004</td>
<td>48 reports, computer search (39 descriptive studies, 4 experimental in design, 5 used qualitative methodology)</td>
<td>Meta-synthesis of the literature using Cooper’s five stages: problem formation; data collection; data evaluation; analysis and interpretation; and presentation of results</td>
<td>Articles representing opinions/discussions were excluded</td>
<td>Three factors explain why women delay in seeking treatment: clinical; sociodemographic; and psychosocial</td>
<td></td>
</tr>
<tr>
<td>Meadows-Oliver, 2003</td>
<td>18 studies, search methods not given</td>
<td>Meta-synthesis</td>
<td>No</td>
<td>Six reciprocal translations of homeless mothers caring for children in shelters: becoming homeless; protective mothering; loss; stressed and depressed; survival strategies; and strategies for resolution</td>
<td>Uses some of the stages in meta-ethnography as described by Noblit and Hare</td>
</tr>
<tr>
<td>Nelson, 2003</td>
<td>Nine studies, computer search</td>
<td>Meta-synthesis</td>
<td>No</td>
<td>Two processes inherent in maternal transition were identified: engagement, and growth and transformation. In addition, five thematic categories identified signifying areas of disruption present in the maternal transition: commitment; daily life; relationships; work; and self</td>
<td></td>
</tr>
<tr>
<td>Nelson, 2002</td>
<td>12 studies, search methods not given</td>
<td>Use of Noblit and Hare’s meta-ethnography</td>
<td>No (three exclusions as not qualitative and two not relevant)</td>
<td>13 themes/metaphors reduced to four steps common to the mothering experience: becoming a mother of a disabled child, negotiating a new kind of mothering, dealing with daily life, and the process of acceptance/denial</td>
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**TABLE 1 Examples of syntheses in the health field that have employed meta-ethnography (continued)**
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<tr>
<td>Paterson et al., 1998&lt;sup&gt;12&lt;/sup&gt; Adapting to and managing diabetes</td>
<td>43 papers (38 studies), computer search</td>
<td>Use of Noblit and Hare’s meta-ethnography</td>
<td>No</td>
<td>Identifies “balance” as the main metaphor in the experience of diabetes, and examines process and requirements for learning to balance</td>
<td>Use of prior organising framework: Curtin and Lubkin’s&lt;sup&gt;13&lt;/sup&gt; conceptualisation of the experience of chronic illness. Several methods used to achieve trustworthiness including other members of team reviewing/agreeing analysis; identifying negative/disconfirming cases; testing rival hypotheses</td>
</tr>
<tr>
<td>Roux et al., 2002&lt;sup&gt;21&lt;/sup&gt; Inner strength in women: meta-synthesis of qualitative findings in theory development</td>
<td>Five studies, search methods not given</td>
<td>Meta-synthesis using Walker and Avant.&lt;sup&gt;90&lt;/sup&gt; Meta-synthesis used is similar to qualitative analysis or constant comparison, the selected literature was read, reread and examined for patterns of similarities among study findings that could be categorised or grouped together</td>
<td>No</td>
<td>“Conceptual model of inner strength in women revealed in the following constructs: knowing and searching; nurturing through connection; dwelling in a different place by recreating the spirit within; healing through movement in the present; and connecting with the future by living a new normal.”&lt;sup&gt;71&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Sandelowski et al., 2004&lt;sup&gt;12&lt;/sup&gt; Stigma in HIV-positive women</td>
<td>93 studies, search methods not given</td>
<td>Meta-synthesis. Worked inductively to create a taxonomy to depict the conceptual range of findings</td>
<td>No</td>
<td>Key findings included pervasiveness of both felt and enacted stigma; gender-linked intensification of HIV infection-related stigma; and unending work and care of stigma management</td>
<td></td>
</tr>
<tr>
<td>Sandelowski and Barroso, 2005&lt;sup&gt;23&lt;/sup&gt; The travesty of choosing after positive prenatal diagnosis</td>
<td>17 studies, computer search</td>
<td>Meta-summary techniques included the calculation of frequency effect sizes, used to aggregate findings. Meta-synthesis techniques included the reciprocal translation of concepts (Noblit and Hare’s&lt;sup&gt;15&lt;/sup&gt; used to interpret the findings)</td>
<td>No</td>
<td>Emphasis in findings is on the termination of pregnancy following positive diagnosis. The thematic emphasis is on the dilemmas of choice and decision-making. For couples, positive prenatal diagnosis was an experience of chosen losses and lost choices</td>
<td></td>
</tr>
<tr>
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<tr>
<td>Sherwood, 1997</td>
<td>16 studies, search methods not given</td>
<td>Use of Noblit and Hare's meta-ethnography</td>
<td>Burns' and Roberts and Burke's applied to each study. None excluded</td>
<td>Developed a therapeutic model of caring that identifies relationships between caregivers and receivers of care in producing therapeutic outcomes (involves the interaction context, nurses' knowledge and nurses' response patterns)</td>
<td>Consultation with two independent caring experts to validate the findings</td>
</tr>
<tr>
<td>Steeman et al., 2006</td>
<td>33 articles (26 studies), computer search, manually searched reference lists</td>
<td>Meta-synthesis aimed at an integrative interpretation of findings from single qualitative studies to synthesise a more substantive description of the phenomenon</td>
<td>Studies meeting inclusion criteria were appraised using Sandelowski and Barroso guide for qualitative research</td>
<td>Living with dementia is described from the stage at which a person discovers the memory impairment, through the stage of being diagnosed with dementia, to that of the person's attempts to integrate the impairment into everyday life</td>
<td></td>
</tr>
<tr>
<td>Swartz, 2005</td>
<td>10 studies, computer search</td>
<td>Meta-synthesis. Use of Noblit and Hare's meta-ethnography</td>
<td>No</td>
<td>Five themes of parenting preterm infants emerged: adapting to risk; protecting fragility; preserving the family; compensating for the past; and cautiously affirming the future.</td>
<td></td>
</tr>
<tr>
<td>Varcoe et al., 2003</td>
<td>Three reports (based on own studies)</td>
<td>Qualitative meta-data analysis influenced by Noblit and Hare's meta-ethnography</td>
<td>No</td>
<td>Importance of 'moral distress' for nurses due to experience of structural and personal constraints (e.g. excessive workloads, absence of interdisciplinary team rounds, conflict between team members and conflict with patients and families) and may lead to coercive practices</td>
<td>All studies conducted within 5-year period in four acute hospitals in same geographical location in western Canada. Studies based on different theoretical perspectives</td>
</tr>
<tr>
<td>Werner, 2002</td>
<td>42 studies, computer search</td>
<td>Meta-synthesis</td>
<td>No</td>
<td>Use of Loomis and Wood's model of the seven human response systems which has three dimensions: health and illness; human-response systems; and nursing clinical decision-making. Findings organised into categories: physical human responses influenced by nurse caring; emotional human responses influenced by nurse caring; cognitive human responses influenced by nurse caring; family human responses influenced by nurse caring; social human responses influenced by nurse caring; spiritual human responses influenced by nurse caring</td>
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### TABLE 1 Examples of syntheses in the health field that have employed meta-ethnography (continued)

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<tr>
<td><strong>Meta-studies</strong></td>
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<tr>
<td>Paterson et al., 2003&lt;sup&gt;[29]&lt;/sup&gt; Embedded assumptions in qualitative studies of fatigue</td>
<td>35 primary research reports, computer search</td>
<td>Meta-study (analysis in three stages): meta-data analysis, meta-method and meta-theory. Synthesis is a creative interpretation of the primary research to produce new and expanded understandings of the phenomenon under study</td>
<td>No</td>
<td>Four key assumptions found in the data: fatigue as exclusively attributed to disease; fatigue as a unitary phenomenon across human experiences; fatigue as inherently and necessarily problematic; and fatigue as isolated from the context in which it occurs</td>
<td></td>
</tr>
<tr>
<td>Paterson, 2001&lt;sup&gt;[30]&lt;/sup&gt; The shifting perspectives model of chronic illness</td>
<td>292 studies that met inclusion criteria from 1000 computer and manual search and professional networks</td>
<td>Meta-study including meta-ethnography</td>
<td>Yes Burns standards&lt;sup&gt;[26]&lt;/sup&gt;</td>
<td>Depicts people with chronic illness as shifting between illness and wellness in the foreground. Also identifies major factors leading to shift to illness in the foreground and process of ‘bouncing back’ to wellness in the foreground</td>
<td>Data synthesised to create model</td>
</tr>
<tr>
<td>Thorne and Paterson, 1998&lt;sup&gt;[31]&lt;/sup&gt; Shifting images of chronic illness</td>
<td>158 reports that met inclusion criteria from 400 identified, computer and manual search (included dissertations)</td>
<td>Meta-study</td>
<td>Yes Burns standards&lt;sup&gt;[26]&lt;/sup&gt;</td>
<td>Describes shifting conceptualisations of individuals with chronic illness and parallel shifts in conceptualisations of health-care relationships appropriate to chronic illness from client-as-patient to client-as-partner</td>
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<tr>
<td><strong>Critical interpretative synthesis</strong></td>
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<tr>
<td>Dixon Woods et al., 2006&lt;sup&gt;[32]&lt;/sup&gt; Conducting a critical interpretive synthesis of the literature on access to healthcare by vulnerable groups</td>
<td>119 papers, computer search, websites, reference chaining, contacting experts</td>
<td>Development and use of the method of CIS. Identified their method as reciprocal translational analysis; refutational syntheses and lines of argument synthesis</td>
<td>NHS National Electronic Library for Health for the evaluation of qualitative research, to inform judgements on the quality of papers. 20 excluded on grounds of being ‘fatally flawed’</td>
<td>Created synthesis including critique of utilisation as a measure of access; candidacy; identification of candidacy; navigation; permeability of services; appearances at health services; adjudications; offers and resistance; operating conditions; and the local production of candidacy</td>
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AIDS, acquired immunodeficiency syndrome; EOL, end of life; HIV, human immunodeficiency virus.

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explicitly as meta-ethnographies was that the latter group adhered more closely to the approach described by Noblit and Hare,15 included a smaller number of studies (maximum of 32) and, in general, achieved a higher degree of conceptual development. Meta-study and critical interpretative synthesis, which have their origins in meta-ethnography, are more recent developments and to date have been used only by their creators. The general tendency among all these syntheses was not to formally assess the quality of the research prior to synthesis. Where this did occur it was liable to result in just a couple of exclusions or in none at all. The reports of most syntheses aimed to draw out the implications for policy-makers and practitioners, but there was little evidence of wider policy/practitioner responses to these findings and assessment of worth in these terms.

The next section considers a development of Noblit and Hare’s15 meta-ethnography through its use as one aspect of a wider meta-study.

**Meta-study**

Paterson et al.42 described an approach to synthesis that incorporates Noblit and Hare’s15 meta-ethnography as part of a meta-study.70 Meta-study aims to go beyond meta-ethnography (or other methods of what they term meta-data analysis), as Paterson and colleagues do not regard meta-ethnography as giving sufficient attention to the way that theories and methods shape knowledge and confer meaning on findings. The enlarged framework of meta-study that Paterson and colleagues present derives from earlier sociological work that was concerned about understanding the ways in which theoretical, methodological or societal contexts shape reported results and bodies of knowledge. They draw particularly on the writings of Ritzer99 and of Zhao,100 who introduced the term meta-study and described its components: meta-data analysis (study of the processed data based on meta-ethnography or other methods of data analysis), meta-method (study of the appropriateness and rigour of particular methods used in the research studies and the implications of a range of epistemologically sound approaches for emerging data and interpretations) and meta-theory (analysis of the philosophical, cognitive and theoretical perspectives underlying the research that influences emerging data and interpretations). Paterson et al.42 described each of these components as providing a unique angle of vision from which to deconstruct and interpret a body of qualitatively derived knowledge about a particular phenomenon, and suggests that meta-synthesis ‘… brings back together those ideas that have been taken apart or deconstructed in these three analytic meta-study processes’ (p. 13). Meta-synthesis, therefore, represents ‘the creation of a new interpretation of a phenomenon that accounts for the data, method and theory by which the phenomenon has been studied by others’ (p. 13) (Figure 1).

Paterson et al.42 outlined the assumptions and methods for conducting a meta-study in their book, *Meta-study of qualitative health research*,42 and illustrated this in terms of the conduct of their own synthesis of the subjective experience of chronic illness.70,101 Their initial phase of meta-data analysis was based on the methods of meta-ethnography, but whereas Noblit and Hare15

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**FIGURE 1** Components of meta-study. Adapted from Paterson et al.42
focused on the in-depth study of a small number of cases, Paterson et al.'s synthesis began with the identification of close to 1000 reports, books, dissertations and research articles published during a 16-year period. These initial publications were reduced to 292 through assessment of their relevance to the review and appraisal of the sampling, data analysis and interpretation that was undertaken using Burns' standards for qualitative research. This appraisal involved at least three members of the team reviewing each report and reaching a consensus. Paterson et al. stated that the number of studies was not reduced further as they regarded this number as necessary to achieve all components of the meta-study. The decision to include such a large number of studies required a systematised approach to meta-ethnography (i.e. meta-data analysis component) and was undertaken by the use of coded categories of themes, patterns, processes, etc. This was followed by a computer-based analysis in much the same way as the analysis of the original primary data. The use of computer software also facilitated comparisons between different groups (e.g. whether or not the disease was considered terminal, the method of sampling, gender, etc.). Paterson et al. acknowledged that the handling of large numbers of reports in this way meant that there was an element of context stripping at this level, although their overall aim was to achieve greater contextualisation through finally integrating the different elements of a meta-synthesis (see Figure 1).

Although Paterson et al. provided a clear and fairly detailed description of their approach to meta-data analysis, the stages of meta-method and meta-theory are less fully developed (particularly meta-theory). They also gave little guidance as to how the three components were to be integrated in the final stage of the meta-synthesis. Indeed, in terms of achieving the synthesis, they stated that they ‘… resist definitive procedural steps and encourage instead a dynamic and iterative process of thinking, interpreting, creating, theorising, and reflecting’.

Paterson et al.'s own experience of conducting a meta-study has been their work on the subjective experience of chronic illness. They acknowledged that the vast size of this undertaking presented considerable demands of organisation and data management, as well as requiring much time and a large team of researchers. What they termed the meta-data analysis component led to the depiction of the shifting perspective experienced by people with chronic illness from ‘wellness’ in the foreground to ‘illness’ in the foreground and the factors responsible for this shift and ‘bouncing back’. Their attempt to achieve a meta-synthesis based on the three components of meta-study has not, however, led to a major new understanding of chronic illness, although they described the changes that have occurred in the conceptualisation of chronic illness over time. The approach they adopted to meta-synthesis also raises issues of practicability, with considerable time and resources required by the processes involved in first deconstructing and then reconstructing and synthesising the distinct components of meta-study; their own study involved seven researchers who contributed time over several years. Nevertheless, their work is important in its emphasis on the need for syntheses to take account of wider contextual factors, including the influence of theories and methods on the questions addressed and the interpretations developed. This is of particular importance in relation to the syntheses of studies conducted over a long time period and involving a broad area of research, as with Paterson et al.’s own study of chronic illness. Their work also draws attention to the importance of meta-theory in terms of the study of the influence of cognitive, philosophical and theoretical perspectives on the questions addressed and interpretations as an aspect of study in its own right to complement more empirical data-based approaches.

Critical interpretive synthesis

More recently, Dixon-Woods et al. used meta-ethnography to synthesise the diverse quantitative and qualitative literature on access to health care by vulnerable groups. In doing so, they suggest that it was necessary to introduce a number of innovations that were judged to be
of sufficient substance to produce what they regard as a new methodological approach that they titled critical interpretive synthesis.

This synthesis was truly innovative in that it used meta-ethnography to synthesise a wide range of different paper types including primary quantitative, primary qualitative, mixed methods, editorial, review and theoretical papers. Books and reports were also included. Dixon-Woods *et al.*, citing Glaser and Strauss, present a compelling argument for there being no reason in principle why interpretive syntheses should not include different forms of evidence. The end result was that six syntheses were produced. The first was a synthesis that set out ‘a general taxonomic framework within which the literature on access to health care can usefully be organised’.* The remaining five syntheses comprised interpretive syntheses of access to health care for specific vulnerable groups (socioeconomically disadvantaged people, people of minority ethnicity, children and young people, and older people) and a synthesis examining the effects of gender. These syntheses led to a substantial number of findings and specific recommendations for policy and practice. In addition, concepts of candidacy and the porosity or permeability of services were proposed as useful aids to understanding access to health care. Candidacy was characterised as the processes of negotiation between people and health-care providers by which people’s eligibility for health care is established. Service porosity signalled ease of use and the number of qualifications for candidacy: the more porous a service the more comfortable it is for people to use and the fewer the qualifications required for candidacy.*

Much of the claimed novelty of critical interpretive synthesis (CIS) rests on the way in which the three different types of syntheses identified by Noblit and Hare* (reciprocal translation, refutational and lines-of-argument) were interpreted and understood within this particular synthesis. In terms of reciprocal translation, Dixon-Woods *et al.* reported two problems. Firstly, they found it difficult to operationalise this process with such a large collection of papers. Although not specifically mentioned, given the wide variation in the types of papers included it seems possible that incommensurability was a problem. In other words, it is likely that the very different kinds of papers included had incompatible aims and therefore could not be integrated within the same frameworks. As we indicated after our first worked example of meta-ethnography, a key assumption of the process is that the studies are commensurable.* A second issue raised by Dixon-Woods *et al.* was that their experience suggested that reciprocal translation ‘provides primarily a demonstration of the extent to which concepts reported in papers are “translatable” into each other, but offers little in the way of insight … it provides only a summary in terms that have already been used in the literature, and the extent to which demonstrating “reciprocity” between the various terms and concepts is a useful exercise is questionable’. However, the authors also noted that in producing a lines-of-argument synthesis it may be necessary to employ ‘found’ or second-order constructs that are already reported in the literature ‘… to code evidence that was not previously categorised using such a construct’. This use of second-order constructs from one study to code findings reported in another study which may not have been explicitly labelled in this way is, however, part of what we would recognise as reciprocal translation. In a more recent paper providing a reflexive account of the process of producing this synthesis, Dixon-Woods *et al.* reflected further on the difficulties that they experienced with reciprocal translation and suggested that it may not have worked for them because of the ‘large and diverse body of literature’ that they were attempting to synthesise.

A further part of the uniqueness claimed for CIS is that it produces a ‘synthesising argument’ that ‘integrates evidence from across the studies in the review into a coherent theoretical framework comprising a network of constructs and the relationships between them’.* It is difficult, however, to see how this differs substantially from Noblit and Hare’s eloquent description of a lines-of-argument synthesis as having as its goal ‘to discover a “whole” in a set of parts’ and being akin to
ethnographic accounts in that ‘it is emic … in its allegiance to the studies being synthesized; it is historical in that it uses time to give order and history-in-use to give context; it is comparative in that it constructs an analogy of relationships among studies and it is holistic in that it constructs an interpretation of all the studies; their interrelations, and contexts’.15

The final original aspect of CIS, and the main rationale for use of the term critical, relates to refutation synthesis. In this new approach it is suggested that the synthesiser should ‘consider contradictions in the evidence in the course of producing the synthesising argument, in the same way as deviant or negative cases would be considered as part of the analysis that produces the theoretical output in primary research’.103 Again this does not seem very different from Noblit and Hare’s15 suggestion that ‘the synthesis must “take into account” the implied relationship between the competing explanations. The implied refutation, then, is analyzed substantively and subsequently incorporated into the synthesis. Our approach treats the refutation itself as part of the interpretation to be synthesised’.15

In their monograph, Noblit and Hare15 did appear to suggest, through the examples that they provided, that any individual synthesis would take just one of the three forms (i.e. reciprocal translation, refutational or lines-of-argument). In practice what we, and Dixon-Woods et al.,103 have found is that one meta-ethnography may include elements of these different types of synthesis. The point at which variations on an existing method become sufficient to justify it being labelled as a new method is a matter or judgement.

The next sections describe two other approaches to meta-synthesis: grounded formal theory, that is closely linked with meta-ethnography; and realist synthesis, that describes an integrative theory-building approach developed in the health-policy field.

**Grounded formal theory**

Grounded formal theory is an approach to meta-synthesis that shares a common framework with Noblit and Hare’s15 meta-ethnography, but is specifically concerned with theory development and gives greater emphasis to the analysis and reporting of verbatim data. Examples of the synthesis of qualitative research using grounded formal theory are Kearney’s104 research aiming to develop a grounded theory of women’s addiction and recovery, and Finfgeld’s105 synthesis of courage among people with long-term threats to well-being that describes a grounded theory approach as forming the ‘epistemological and methodological basis’ for the study.

Kearney104 and Finfgeld105 both included dissertations in the reports that they synthesised because these include more first-hand data. Kearney104 described how she analysed the research reports as if each one was an interview transcript and coded descriptively (verbatim data) and theoretically. She developed a formal theory using a constant comparative approach that involved consulting the verbatim descriptions and the sociocultural and historical context. Following the requirements of grounded theory, once the formal theory was developed other kinds of qualitative reports on addiction and recovery were examined for relevance and to support or challenge the theory (e.g. personal accounts, autobiographies and ethnographies). This led to the development of a theory of truthful self-nurturing that regards the basic problem of addiction as self-destructive self-nurturing, and the basic process of recovery as truthful self-nurturing in which addiction gains meaning as a problem. Recovery work therefore involved three areas of psychological change: abstinence work, self-work and connection work. The analysis is illustrated by substantial reference to original data and the inclusion of verbatim material from the studies in terms of the respondent’s own words. Finfgeld’s105 approach gives a similar emphasis to coding verbatim data from the included reports. This material was then grouped into metaphorical codes and a process model was developed that consisted of the causal and contextual conditions of the phenomenon of becoming and being courageous.
Realist synthesis

This term was introduced by Pawson\textsuperscript{106} to describe a new methodology developed in the field of evidence-based policy to understand the changes introduced by programmes in terms of their generative mechanisms. Realist synthesis aims to achieve this through a process of theory building rather than assembling empirical generalisations, and draws on material regarding the workings of the same underlying mechanism (such as the public disclosure of information on performance) in different contexts. It therefore differs from other forms of synthesis in bringing together material from diverse sources that may include health, education, social policy or other sectors. In addition, the explicit aim of realist synthesis is theory development. Realist synthesis has so far not been widely employed, but forms one of the increasing armoury of approaches to developing new conceptualisations by drawing together the findings of individual studies and shares with other interpretive approaches the aim of going beyond individual studies to develop new theories and conceptualisations.

Summary

In this chapter we sought to identify the various approaches and methods that can and have been used to synthesise the findings from individual qualitative research studies. Both Hammersley\textsuperscript{14} and Noblit and Hare\textsuperscript{15} differentiated integrative syntheses, which involve the aggregation, pooling or comparison of findings from qualitative studies, from interpretative syntheses, whose methods involve induction and interpretation and whose aim is to achieve conceptual and theoretical development and, hence, greater insight into particular phenomena or issues.

The method that has received the greatest attention to date is meta-ethnography. This method has been applied by some in a way that appears to be fairly consistent with the approach to qualitative synthesis described by its architects, Noblit and Hare.\textsuperscript{15} Others, notably Paterson et al.\textsuperscript{42} have sought to use meta-ethnography as a procedure within a more all-encompassing form of synthesis which they call meta-study. In addition, Dixon-Woods et al.\textsuperscript{102} have tested whether or not meta-ethnography can be used to bring together a large and methodologically diverse group of studies and in the process have evolved what they consider to be a new method, CIS.

The subsequent chapters consist of a detailed evaluation of meta-ethnography as a method of qualitative synthesis. To accomplish this evaluation, two substantial syntheses were undertaken: the first was a synthesis of studies of medicine-taking and the second a synthesis of patients’ experiences of RA. In both cases the intention was to apply the method as depicted by Noblit and Hare.\textsuperscript{15} The next chapter describes how relevant qualitative studies on these topics were identified.
Chapter 3

Literature searching

Introduction

Although the potential relevance and importance of qualitative research is being increasingly acknowledged by policy-makers and those concerned with establishing evidence-based medicine and social care, there is consensus that searching for and identifying appropriate qualitative research remains frustrating and difficult. The problem becomes particularly acute when attempting a synthesis of the studies on a given topic – the task that we have set ourselves here. Naturally, one would wish to locate the maximum amount of primary research in the most efficient way before undertaking further assessment of the material. However, it is becoming ever clearer that it is no easy task to develop a search strategy that maximises recall and precision, while ensuring that expenditure of resources is kept within manageable limits.

Considerable effort has already been expended to produce guidelines and procedures for retrieval and, to a lesser extent, to evaluate their usefulness and relative efficiency. Nevertheless, it is apparent that even when the focus is on studies using quantitative methods reported in the medical literature (and hence perhaps more systematically indexed), retrieval rates can be surprisingly low. Limitations with searching bibliographic databases led the Cochrane Collaboration to search European general health-care journals by hand for reports of controlled trials: only 17% of those identified were indexed in MEDLINE as controlled trials and 30% were not indexed in MEDLINE at all. Accordingly, a multipronged approach is currently advocated: searching a range of databases; hand-searching of relevant journals; contacting experts in the field of enquiry for curricula vitae and information; and examination of the ‘grey literature’, conference proceedings and the like. Decisions have to be made at each stage depending on the resources available.

Whatever problems are encountered in searching for studies employing quantitative methods, they are multiplied when searching for relevant qualitative research. For example, Evans highlighted the particular difficulties of searching electronic databases for reports of qualitative studies arising because of the nature of the titles, the content of the abstracts and the indexing practices of those preparing the databases. Evans pointed out that the titles given to qualitative work, although often very descriptive of the study’s focus, may not easily lend themselves to the selection of specific key terms. Moreover, the content of the abstracts can also vary greatly and the research method may not be stated in the title or the abstract. Such studies run the risk of failing to be identified.

For this reason, it is important to include a range of databases as they vary in the extent to which they use methodological indexing terms that accurately describe qualitative study design. In addition, the more diffuse or broad the topic for review or synthesis is – and this may often be the case where qualitative research is involved – the more necessary it is to throw the net wider and include more databases. However, the more varied the databases, the more problematic the notion becomes of a ‘standard’ search filter suitable for application to all. Search strings to identify qualitative studies probably need to be adapted to the idiosyncrasies of each synthesis to achieve the best results. Moreover, qualitative research lacks the more standardised keywords associated with quantitative research in medicine. An evaluation of three different electronic
search strategies for finding qualitative research (using thesaurus, free-text or broad-based terms) by Shaw et al.\textsuperscript{114} found that relying on one strategy alone would have resulted in failure to identify some relevant records. This evaluation also highlighted the low level of precision of all three strategies, with only 4% of papers initially identified as potentially relevant actually proving to be so.

In addition to the need to include a range of databases and search strategies in order to maximise the yield of relevant qualitative research papers comes the further complication that social science literature is much more diverse in terms of its publication media than medical literature.\textsuperscript{115} This suggests a requirement for the effective qualitative literature search not simply to be confined to electronic searches for journal articles.

In the following sections we describe the choices made and the strategies adopted for the retrieval of material for the two synthesis studies undertaken, one being a relatively diffuse topic, whereas the other had a specific disease focus. In both cases, the decision was made at the beginning not to explore the ‘grey literature’ or to consider unpublished theses. The decision to restrict ourselves to published work was made in order to keep the number of papers manageable, as we anticipated that these would be high, particularly in the case of the medicines synthesis. It was also part of the purpose to investigate the usefulness of the methodology for HTA and health-services research in which published research would be easier and quicker to access. The literature searches for both syntheses were conducted in July and August 2002.

**Methods**

The research group formed into two subgroups, one for each of the syntheses. PP led the medicines synthesis and GDW led the RA synthesis. Each subgroup managed the process of searching, appraising and synthesising separately, although the whole group got together regularly to discuss progress and general issues arising from the projects.

**Medicines synthesis**

**Defining the topic**

We used the following definition when searching: ‘Papers whose primary focus is patients’ views of medicines prescribed and taken for the treatment of a long- or short-term condition (excluding medicines only taken for preventive purposes)’. The study had to use qualitative methods of data collection and analysis and be published in the English language.

**Sampling strategy**

We chose the 10-year period from 1 January 1992 to 31 December 2001 (the study began in spring 2002), believing that most of the relevant studies would be found within this period. We were aware that a small number of important early studies would be missed, e.g. Arluke,\textsuperscript{116} Trostle et al.,\textsuperscript{117} and Conrad,\textsuperscript{118} but needed to set limits on what was a potentially large number of available studies.

**Electronic search strategies**

We used the following electronic databases: MEDLINE, EMBASE, CINAHL, Web of Science, PsycINFO and Zetoc. The initial MEDLINE search was very detailed (Table 2) and produced a large pool of papers. Hoping for a more sensitive and focused search, we tried simply using the terms ‘patient compliance’, ‘adherence’, ‘medicine’, ‘medication’ and ‘qualitative’ (Table 3). The second, simpler, MEDLINE search produced a much smaller pool of potential papers, but all the relevant papers that had been identified in the first search (this simpler search produced
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continued
We repeated these two searches in EMBASE. Again, the second, simpler, search produced all the relevant papers that had been identified in the first, more detailed EMBASE search. The same strategy in CINAHL resulted in the second, simpler, search producing an additional three relevant papers to the first. Thus, the simple search was just as sensitive as the more comprehensive search in MEDLINE and EMBASE, and more sensitive in CINAHL.

For Zetoc, the search terms used were ‘prescri*’ and ‘qualitative’. For Web of Science, the following search was conducted: (medicine* or medication* or prescription*) and (compliance or adherence or concordance) and (qualitative or grounded same theory or patient* same interview* or patient* same perception* or patient* same experience* or content same analysis or ethnography). The databases searched were SCI-EXPANDED, Social Sciences Citation Index, Arts and Humanities Citation Index. The following search was conducted for PsycINFO: (medicine* or medication* or prescription*) and (compliance or adherence or concordance) and (qualitative or (grounded adj theory) or (patient* adj interview*) or (patient* adj perception*) or (patient* adj experience*) or (content adj analysis) or ethnography) and (PO= HUMAN).

### Hand-searching strategies

For the hand-searches we asked colleagues and team members to suggest any papers they might think relevant, we searched the departmental Reference Manager (version 8; Thomson Research Soft, Carlsbad, CA, USA) database (which includes a record of all references held by

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**TABLE 2** Initial detailed MEDLINE search for medicines synthesis (continued)

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<td>17 and 42 and 56</td>
<td>2105</td>
</tr>
<tr>
<td>59</td>
<td>57 and 58</td>
<td>305</td>
</tr>
<tr>
<td>60</td>
<td>limit 59 to yr=1997 to 2001</td>
<td>268</td>
</tr>
</tbody>
</table>

**TABLE 3** Simple MEDLINE search for medicines synthesis

<table>
<thead>
<tr>
<th>#</th>
<th>Search history</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Patient Compliance/or patient compliance.mp.</td>
<td>7154</td>
</tr>
<tr>
<td>2</td>
<td>adherence.mp. [mp=title, abstract, cas registry/ec number word, mesh subject heading]</td>
<td>6563</td>
</tr>
<tr>
<td>3</td>
<td>medicin$.mp.</td>
<td>37,736</td>
</tr>
<tr>
<td>4</td>
<td>medication$1.mp. [mp=title, abstract, cas registry/ec number word, mesh subject heading]</td>
<td>23,725</td>
</tr>
<tr>
<td>5</td>
<td>qualitative$.mp. [mp=title, abstract, cas registry/ec number word, mesh subject heading]</td>
<td>16,190</td>
</tr>
<tr>
<td>6</td>
<td>1 or 2</td>
<td>12,812</td>
</tr>
<tr>
<td>7</td>
<td>3 or 4</td>
<td>60,560</td>
</tr>
<tr>
<td>8</td>
<td>5 and 6 and 7</td>
<td>37</td>
</tr>
<tr>
<td>9</td>
<td>Limit 8 to yr=1997-2001</td>
<td>32</td>
</tr>
</tbody>
</table>
academic and research staff in the School of Social and Community Medicine at University of
Bristol), we checked reference lists of obtained papers and we searched the Medicines Partnership
website. We also hand-searched the Arts and Humanities Library and the Medical Library of
University of Bristol. All books in the RA–RC (medical) section and in the HM–HV (general
sociology) section were searched, as were short-loan collections for the same class marks.

The key journals were hand-searched in case the electronic searches were not sufficiently
sensitive or in case the individual articles had not been given adequate keywords. Hand-
searching of journals was conducted for either the online or paper versions of journals and was
shared between members of the team. The journals we hand-searched were Qualitative Health
Research; Sociological Review; Psychology and Health; Journal of Health Psychology; Journal of
Advanced Nursing; Medical Anthropology Quarterly; Culture, Medicine and Psychiatry; Medical
Anthropology; Patient Education and Counselling; Family Practice; Sociology of Health and Illness;
Social Science and Medicine; and Health.

The process of selecting/rejecting
For the electronic searching, PP was responsible for deciding whether or not to include studies.
She examined titles and abstracts and obtained full copies of papers that looked promising. For
hand-searching, the decision-making process was shared among team members and occurred as
we went along. The process of selecting/rejecting was lengthy but was conducted thoroughly at
this stage (rather than being initially overinclusion as with the RA synthesis), because of the much
larger number of studies involved.

Results of searches
The electronic searches produced 21 studies for appraisal and the hand-searches 21, giving
a total of 42 studies to be appraised for inclusion in the synthesis. (Further exclusions could
be made following the appraisal.) Figure 2 gives a breakdown of the yield from each of the
different databases.

Reproducibility of electronic search strategy
A second researcher, GDW, conducted the electronic searches independently in order to assess
the reproducibility of the selection/rejection process. GDW used the same search strategy with
the same databases and examined titles and abstracts before drawing up a list of papers for
possible inclusion (Box 3). Of the 21 papers that PP found electronically, GDW found 18. He also
identified one additional useful paper that PP had missed, bringing the final number of studies to
be appraised to 43 (22 produced electronically and 21 from hand-searching) (Box 3). (GDW also

![FIGURE 2](image-url)
Literature searching

identified 22 other papers which PP had rejected. This may be accounted for by the fact that PP had time to obtain papers that she was not sure about and look at them in more detail, but for the purposes of this exercise, GDW was working with titles and abstracts only.)

**Success of search strategy**

As indicated above, the double screening of titles and abstracts ensured that one relevant study that might otherwise have been missed was included. Time limitations meant that only one person hand-searched any one journal. Late in the course of conducting the synthesis, we came across a study that could have been included and which should have been identified during the hand-searching. Given that this discovery was made at a point at which the synthesis was well advanced, we did not feel it appropriate to try and incorporate it post hoc.

**Rheumatoid arthritis synthesis**

**Defining the topic**

Papers were sought that reported the results of qualitative studies concerned with lay experiences of RA, or lay perceptions of aetiology, treatment or management. The study had to use both qualitative methods of data collection and analysis. Papers were not restricted to English language only, because initial scoping exercises suggested that the number of papers produced for the RA synthesis would be manageable.

**Sampling strategy**

We searched intensively for papers published between January 1992 and December 2001, using the strategies below. However, because we knew that important work had been published before 1992, and because initial searches suggested that the number of potential papers would be manageable, we decided to further include so-called ‘classic’ studies in the field that were published prior to 1992. To decide which ‘classic’ papers to include, we consulted relevant experts in medical sociology, nursing, occupational therapy and rheumatology, including authors of some of the papers identified as a result of our searches below.

**Electronic search strategies**

We used the following electronic databases: MEDLINE, EMBASE, CINAHL, Web of Science, PsycINFO and Zetoc. The MEDLINE searches are detailed in Tables 4 (the detailed search) and 5 (the simpler one). The two different searches were conducted in order to maximise yield. Although they produced some of the same papers, they also each produced a distinct list of papers that met the inclusion criteria. The same searches were run in EMBASE and CINAHL. For PsycINFO and the Web of Science (Science Citation Index and Social Science Citation Index), the search terms were ‘arthritis and (qualitative or focus group or grounded theory or lay or

---

**BOX 3 Searching and selection process for medicines synthesis**

<table>
<thead>
<tr>
<th>Electronic searches (PP)</th>
<th>21</th>
</tr>
</thead>
<tbody>
<tr>
<td>Additional paper identified by GDW as a result of reproducibility exercise (electronic)</td>
<td>1</td>
</tr>
<tr>
<td>Hand-searches (whole team)</td>
<td>21</td>
</tr>
<tr>
<td>Total papers for appraisal</td>
<td>43</td>
</tr>
</tbody>
</table>

identified 22 other papers which PP had rejected. This may be accounted for by the fact that PP had time to obtain papers that she was not sure about and look at them in more detail, but for the purposes of this exercise, GDW was working with titles and abstracts only.)

**Success of search strategy**

As indicated above, the double screening of titles and abstracts ensured that one relevant study that might otherwise have been missed was included. Time limitations meant that only one person hand-searched any one journal. Late in the course of conducting the synthesis, we came across a study that could have been included and which should have been identified during the hand-searching. Given that this discovery was made at a point at which the synthesis was well advanced, we did not feel it appropriate to try and incorporate it post hoc.

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perception or experience)’ for the period 1992–2001. For Zetoc, six separate simple searches were conducted:

1. arthritis qualitative
2. arthritis grounded theory
3. arthritis focus group
4. arthritis lay
5. arthritis interview
6. arthritis perception.
**Hand-searching strategies**

For the RA synthesis, the hand-searches were conducted in the same way as for the medicines synthesis, the only differences being that the journal *Family Practice* was not hand-searched for the RA synthesis, nor was the Medicines Partnership website.

**The process of selecting/rejecting**

For the electronic searching, GDW was responsible for deciding whether or not to include studies. He examined titles and abstracts and obtained full copies of papers that looked promising. For the hand-searching, decisions were shared among team members and were made as we went along. Because of the manageable number of papers, we erred on the side of inclusivity at this stage.

**Results of searches**

The electronic searches produced 38 papers for appraisal and the hand-searches four. A further nine ‘classics’ were obtained, giving a total of 51 studies to be appraised. (Further exclusions could be made following the appraisal.) Figure 3 provides a breakdown of the yield from each of the different databases.

**Reproducibility of electronic search strategy**

The electronic searches were conducted independently by a second researcher, RC, in order to assess the reproducibility of the selection/rejection process. RC used the same search strategy with the same databases and examined titles and abstracts before drawing up a list of papers for possible inclusion (Box 4). RC’s searches did not produce any additional relevant papers.

**Conclusion**

When searching for qualitative reports, the policy of a ‘belt and braces’ strategy was certainly vindicated for both syntheses. Perhaps the most striking feature of the results of our searches was the demonstration of the importance of hand-searching and consultation with experts, and the difference between the two topic areas. For the medicines synthesis, a more diffuse topic, these sources contributed as many papers as did the electronic searching. For the disease-based synthesis on RA, electronic searching identified three-quarters of the papers considered for appraisal. The yield from the various electronic databases also differed between the two topic areas, with MEDLINE being most successful for the medicines synthesis and Zetoc the most successful for RA.

Our strategy included a limited search of library books, but excluded theses. One book was included in the RA synthesis and one book chapter, and two book chapters were included in the medicines synthesis. Searching for books and theses is challenging as they are not indexed in the same way as journal papers, and may be accessible only from experts who know the field. Qualitative researchers have often chosen to publish their work in book form as this is far less constraining than publishing in journal articles. Searches confined to journal articles may

**Box 4 Searching and selection process for RA synthesis**

| Electronic searches (GDW) | 38 |
| Hand-searches (whole team) | 13 (of which 9 ‘classics’) |
| Reproducibility exercise | 0 |
| Total papers for appraisal | 51 |
produce a ‘truncation’ bias as the full details of a descriptive qualitative study are unlikely to be published in a short article. The deciding factor is likely to be the resources available in any individual project. If resources are available to include theses, one strategy would be to identify recent theses on the grounds that they are less likely to have been published in journal form.

This experience has reinforced our commitment to an eclectic approach to the issue of searching and the need to remain alert to the distinct possibility that what worked last time will not necessarily work for the next retrieval exercise. What is clear is that this is never going to be a process capable of being regulated and therefore totally delegated. This vital initial part of any synthesis requires experienced skills of judgement and persistence, just as much as at the later stages.

Although it is too early to make authoritative recommendations, we can make some observations that may be helpful to others attempting a similar task. A multiple search strategy is more likely to identify relevant qualitative research than one relying solely on electronic searching. However, the purpose of the synthesis will determine the most appropriate search strategy. In the synthesis of quantitative research, the purpose is usually to be comprehensive, and to identify all the published and grey literature on a particular question in order to calculate a precise and unbiased estimate. In qualitative synthesis, the aims may be different. One aim might be the mapping out of key conceptual developments in a particular field. For this purpose, a grounded theory and/or purposeful sampling approach may be appropriate. If the aim is not to produce an aggregative synthesis, the omission of some papers is unlikely to have a dramatic effect on the results. Our experience suggests that a maximum of about 40 papers is realistic because it is difficult to maintain sufficient familiarity with > 40 papers when trying to synthesise them all and that it is, therefore, best not to be overambitious in terms of numbers.

FIGURE 3 Search yields (RA synthesis) for 51 papers. *Papers could appear on more than one electronic database. Zet, Zetoc; Cin, CINAHL; Med, MEDLINE; Emb, EMBASE; WoS, Web of Science; Hnd, hand-searching; Psy, PsycINFO.
Chapter 4

Appraisal of papers

Introduction

Conventional models for conducting quantitative systematic reviews first require the application of criteria for assessing the quality of the research in the literature. Study quality in this context largely equates to methodological quality. The highest quality quantitative research is usually deemed to be that in which the study design minimises bias (systematic error) and, hence, maximises internal validity. The more internally valid a study is, the argument runs, the more likely its results are ‘to approximate to the truth’.121 Within quantitative systematic reviews, a minimum quality threshold is normally set to ensure the exclusion of studies so methodologically weak that their results are likely to be very biased. The quality grade of studies included can also be used to weight the findings of studies contributing to cumulative meta-analyses and be used to explore heterogeneity in study results.

Over the past 30 years, there has been considerable debate about quality assessment in qualitative research. This debate centres on two related questions: firstly, whether or not there is a plausible philosophical rationale for undertaking such assessments; and, secondly, if such a rationale exists, what criteria should be used to inform judgements about quality. Fundamental issues of ontology, epistemology and methodology underpin this, and debates about the nature and practice of qualitative research have been described, explained and analysed in considerable detail by Murphy et al.9 Quality assessment of qualitative research was part of the brief for this methodological research, making it necessary to include a brief summary of the main positions taken in this debate.

Taking the view that there is no single, unequivocal social reality or truth independent of the researcher or the research process, extreme relativists argue that it is impossible to judge research against external criteria. They contend that the multiple realities produced by different styles of research, and from different studies, provide unique but equally valid perspectives. Lincoln and Guba,122 although rejecting the conventional quantitative criteria of reliability, validity and generalisability, took a less extreme stance when they maintained that it was possible to develop alternative criteria better suited to the assessment of qualitative work. They suggested a focus on credibility (as confirmed by the participants or subjects through member validation); transferability (how far the ideas are transferable to another context); consistency or dependability (how well the researchers have taken into account ways in which the phenomenon studied have changed over time); and confirmability (the ability to follow how the conclusions were reached). Later they argued for qualitative work to be judged on its authenticity, which rests on the researchers themselves demonstrating the fairness and strength of the interpretation and the impact on the participants.123

Against these relativist perspectives, others124,125 have argued that the same criteria can be used to assess quantitative and qualitative research. Hammersley126 proposed subtle realism as a variant of this position, accepting that similar criteria apply, but suggesting that some of the methods used to assess quality in quantitative research styles may need to be adapted for qualitative work. He argued that the central issues for all research are validity and relevance. Validity can be judged from the clarity of the account of data collection and analysis; the researchers’ sensitivity
to, and reflection on, the ways in which the researcher/research has shaped the data; fairness or balance in the account; and attention to negative or deviant cases that disprove or develop the interpretation. Relevance is concerned with whether or not the research adds to the sum of knowledge and the extent to which findings can be generalised beyond the setting in which they were generated.\textsuperscript{126}

Taking account of this debate, but also as a pragmatic response to requests from publishers, funding bodies and those in the health service who wanted guidance on how to assess qualitative research, a substantial number of authors have produced sets of criteria and checklists.\textsuperscript{83,127–132} Indeed, research to examine how systematic reviews can incorporate qualitative research identified > 100 such sets of criteria.\textsuperscript{20} To our knowledge none of these lists of criteria has, however, been designed specifically as a precursor to conducting any secondary analysis or synthesis of qualitative research. The Joanna Briggs Institute at the University of Adelaide, Australia, has, however, developed systematic review software that includes a module ‘designed to manage, appraise, extract and synthesise qualitative data as part of a systematic review of evidence’ [www.joannabriggs.edu.au/Appraise%20Evidence/JBI%20SUMARI%20(systematic%20review%20software)%20FREE].

As the review in Chapter 2 of qualitative research syntheses employing meta-ethnography or related methods illustrates, only a small minority of researchers have undertaken an appraisal of the quality of individual research papers as a precursor to the synthesis. Sandelowski et al.\textsuperscript{13} argued against assessing papers for methodological quality because of concern that relatively minor methodological flaws may result in insightful studies being excluded from syntheses. They suggested that the qualitative synthesiser needs to apply careful judgement in order to distinguish between superficial mistakes and flaws of such magnitude that the findings are rendered invalid. In their recent review of vulnerable groups and access to health care, Dixon-Woods et al.\textsuperscript{82} followed this prescription and used a very brief appraisal tool to identify papers that were ‘fatally flawed’ and therefore unsuitable for inclusion.

The recent drive within the UK government towards evidence-based policy-making has also focused attention on the quality of the qualitative research because government policy-makers are substantial consumers of qualitative social research. As a result, the Strategy Unit in the Cabinet Office commissioned the National Centre for Social Research (NatCen) to produce guidance on quality assessment in qualitative research.\textsuperscript{19} After conducting a number of review exercises, the NatCen team proposed a new generic assessment framework for qualitative research. Drawing on 29 sets of previously suggested assessment criteria, it comprises a series of 18 appraisal questions underpinned by four ‘guiding principles’. For each question, a set of points for consideration is suggested. This new framework was still under development when we began our evaluation of meta-ethnography and so we continued to use the modified version of the CASP criteria for quality appraisal\textsuperscript{18} that we had used in our pilot synthesis on lay experiences of diabetes.\textsuperscript{17} We originally chose the CASP criteria\textsuperscript{18} because they had been developed and tested over a period of time and because they were widely recommended for use within the NHS.\textsuperscript{18}
Prior to the quality appraisal, two screening questions were asked:

1. Does the paper report on findings from qualitative research and did that work involve both qualitative methods of data collection and analysis?
2. Is the research relevant to the synthesis topic?

If one of the answers to these questions was ‘no’, the paper was excluded. The first question effectively excluded those papers that could be regarded as ‘fatally flawed’. If both answers were ‘yes’, the appraisal could proceed. This involved answering questions (both yes/no answers and an elaboration on these) about each paper for each of the following categories: aims, methodology, theoretical perspective, sampling, data collection, data analysis, research relations, data interpretation, transferability, relevance and usefulness. The data recorded were both quantitative (the yes/no answers) and qualitative (the elaboration on those answers). A key part of the appraisal was to outline, in detail, the main findings and concepts. These were recorded in large free-text boxes on the database.

Assessing reproducibility of appraisals

A second reviewer from the research team also appraised each of the papers, blind to the first appraisal, in order to assess the reproducibility of the process. (The results of this process are described in detail below.) The independent reviewers were allocated to each paper in a random manner using an allocation schedule drawn up in advance. This used random number tables and randomisation blocks of nine. If independent reviewers were allocated a paper that they had themselves authored, a different reviewer was chosen randomly from those remaining. The independent reviewers were MM, NB and LY for the medicine-taking synthesis and RC, RP and JD for the RA synthesis.

Inclusions/exclusions

Four of the papers in the medicine-taking synthesis were the subjects of disagreement (between the first and second reviewers) about whether or not to include them. This was because they were considered borderline, either in terms of methodology (were they sufficiently qualitative?) or because the findings were somewhat slight. Disagreements were resolved through a process of negotiation (by e-mail), with the result that one paper was excluded and three were included with the proviso that if they contributed nothing to the synthesis they would ‘fall out’ later on. In the RA synthesis, a team meeting was held to decide the fate of papers over which there was disagreement. If disagreement persisted or if studies were considered borderline according to the inclusion criteria, they were included.

For the medicine-taking synthesis, five papers were excluded following appraisal; for the RA synthesis, 22 papers were excluded following appraisal. The larger number of exclusions in the RA synthesis is largely because, at the literature searching stage, GDW decided to retain any papers that might prove relevant and enter them into the appraisal process. Papers in the medicine-taking synthesis were more heavily screened for relevance by PP at the literature searching stage. Thirty-eight papers were finally included in the medicine-taking synthesis and 29 papers in the RA synthesis, although further exclusions were made during the synthesis (Box 5).

Quality

We did not exclude papers on grounds of quality. The only reasons for excluding papers were either that they were not sufficiently focused on the topic or that they did not actually report qualitative research. Sometimes this was not immediately obvious. For example, some papers described work where qualitative data had been collected, but on closer examination it became clear that the data were not analysed qualitatively. In several papers that, strictly speaking, were...
relevant and the methodology was qualitative, the analysis was considered to be very superficial or few findings were reported. In these cases we did not exclude papers, but attempted to synthesise them.

Sometimes, it was an absence of adequate reporting that could confuse matters. For example, in the RA synthesis, some of the better theoretical papers (the ‘classics’) scarcely reported their research methods. These papers were conceptually very useful to the synthesis, but if strict quality screening had been employed these papers would have been excluded. Indeed, many of the RA studies would have been so excluded. Only four papers in the RA synthesis were characterised both by good reporting of the use of qualitative methods of data collection and by analysis. (This is discussed further in relation to specific studies in Chapter 7 in which the RA synthesis is reported.) On the other hand, for both topic areas there were papers that reported on methodology in great detail, but produced very superficial findings. We remained inclusive in our approach because, although some of the papers were slight, they were synthesisable in that at least they were descriptively confirmatory of previous findings.

In the medicine-taking synthesis we did not attach greater or lesser importance to any of the papers, we simply tried to put all the findings together. Some papers contributed more in terms of findings than others, but this is not necessarily a comment on their significance. Some papers were more interesting to read than others – usually implying that they interpreted their data rather than simply offering description and content analysis – but all papers were treated in the same way. Whatever they offered, whether description or interpretation, was synthesised. Some papers contributed more to particular areas than others. For example, the asthma papers contributed the least overall, but they made a strong contribution in one area, that of identity and the psychological aspects of medicine-taking.

Several of the individual questions posed in the appraisal were difficult to interpret (e.g. some of those on sampling) and others were felt to be redundant (e.g. one on the setting in which data collection occurred). The question on theoretical perspective was felt to be of limited value, particularly as many of the later studies that we found were primarily descriptive.

**Appraising the papers in Access**

After conducting the appraisals, both PP and GDW found it easier to import the findings and concepts from the Access database into Microsoft Word and work with the data in this format. Only one of the researchers (LY) found it necessary to access the database thereafter, in order to analyse the reproducibility of the appraisals. For quantitative analyses, therefore, the
database was useful. However, the majority of the data were qualitative. Given the time that we invested in setting up the database and the small returns we obtained from doing so, we do not consider that it was a worthwhile endeavour to have conducted the appraisals in Access 2000. After conducting the appraisals, both PP and GDW found it much easier to revisit the papers themselves, rather than access the database, for any information that was subsequently needed. Those independent second reviewers who did not have access to the database conducted their appraisals in Word and reported that this was acceptable.

Reproducibility of appraisals

To evaluate the reproducibility of our quality assessment procedure, data from the Access 2000 database were exported to an SPSS (version 12; SPSS Inc., Chicago, IL, USA) save file for analysis.

Inter-rater agreement

The data were firstly examined to determine inter-rater agreement (Cohen's kappa) for each item. This calculation was based on six pairings of assessors, as there were two first assessors, each of whom was paired with three different second assessors. The number of papers rated by each of the six pairs for which complete data were available was too low (ranging from 6 to 14) to permit meaningful statistical comparison of the differences between kappa values for each pair of assessors, and so averaged kappa values for all pairs are presented in Tables 6–10.

It is evident that inter-rater agreement was typically rather low, albeit better for some classes of questions than others. There was moderate-to-good agreement on the majority of questions relating to design, sampling and recruitment (see Table 6), and modest agreement for all those relating to data analysis (see Table 8) and the value of the study (see Table 10). However, agreement was poor for questions concerning data collection (see Table 7), the interpretation and presentation of data (see Table 9) and the transferability of results (see Table 10).

Pooled ratings of papers

For all subsequent analyses, the data were pooled, such that ratings of the papers by all assessors were combined. The pooled data thus aggregated two sets of ratings for each paper: one set was

<table>
<thead>
<tr>
<th>CASP question</th>
<th>Kappa</th>
<th>% 'Yes'</th>
<th>r</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is there a clear statement of the aims of the research?</td>
<td>0.17</td>
<td>91.2</td>
<td>0.02</td>
</tr>
<tr>
<td>Is a qualitative methodology appropriate for authors' stated aims?</td>
<td>N/Aa</td>
<td>96.6</td>
<td>0.06</td>
</tr>
<tr>
<td>Is a theoretical perspective identified?</td>
<td>0.70</td>
<td>43.9</td>
<td>-0.01</td>
</tr>
<tr>
<td>Is it clear which setting(s) the sample was selected from (e.g., hospital)?</td>
<td>0.63</td>
<td>91.2</td>
<td>0.29</td>
</tr>
<tr>
<td>Is it clear why this setting was chosen?</td>
<td>0.09</td>
<td>70.1</td>
<td>0.21</td>
</tr>
<tr>
<td>Is clear and adequate information given on who was selected?</td>
<td>0.23</td>
<td>76.4</td>
<td>0.13</td>
</tr>
<tr>
<td>Is it clear why these samples were selected?</td>
<td>0.17</td>
<td>69.9</td>
<td>0.17</td>
</tr>
<tr>
<td>Is it clear how the sample was recruited?</td>
<td>0.52</td>
<td>55.1</td>
<td>0.06</td>
</tr>
<tr>
<td>Is the sample size justified by the authors?</td>
<td>0.43</td>
<td>17.6</td>
<td>0.10</td>
</tr>
<tr>
<td>Is it clear how many people accepted or refused to take part?</td>
<td>0.80</td>
<td>24.5</td>
<td>0.07</td>
</tr>
<tr>
<td>Is it clear why some participants chose not to take part?</td>
<td>0.65</td>
<td>5.8</td>
<td>-0.04</td>
</tr>
<tr>
<td>Overall, is the sampling strategy appropriate to address the aims?</td>
<td>0.18</td>
<td>74.7</td>
<td>0.32</td>
</tr>
</tbody>
</table>

N/A, not applicable.

a. Could not be computed because the first rater gave the same value ('yes') for all papers.

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TABLE 7  Critical Appraisal Skills Programme\textsuperscript{18} questions regarding data collection: percentage of ‘yes’ responses to item, inter-rater agreement (kappa) and correlation with global rating of study quality

<table>
<thead>
<tr>
<th>CASP question</th>
<th>Kappa</th>
<th>% ‘Yes’</th>
<th>(r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is it clear where the setting of the data collection was?</td>
<td>0.74</td>
<td>59.2</td>
<td>0.21</td>
</tr>
<tr>
<td>Is it clear why that setting was chosen?</td>
<td>0.17</td>
<td>27.5</td>
<td>0.18</td>
</tr>
<tr>
<td>Is it clear how the purpose of the research was explained to the participants?</td>
<td>0.23</td>
<td>14.3</td>
<td>–0.03</td>
</tr>
<tr>
<td>Is it clear how the data were collected and why (e.g. interviews, focus groups)?</td>
<td>–0.04</td>
<td>96.6</td>
<td>–0.06</td>
</tr>
<tr>
<td>Is it clear how the data were recorded (e.g. audio/video/notes, etc.)?</td>
<td>0.36</td>
<td>88.4</td>
<td>0.00</td>
</tr>
<tr>
<td>Is there evidence of flexibility/an iterative process in how research was conducted?</td>
<td>0.24</td>
<td>25.7</td>
<td>0.14</td>
</tr>
<tr>
<td>Is it clear who collected the data?</td>
<td>0.24</td>
<td>66.0</td>
<td>0.14</td>
</tr>
<tr>
<td>Overall, do you consider that data collection addresses the research aims?</td>
<td>–0.07</td>
<td>91.8</td>
<td>0.37</td>
</tr>
</tbody>
</table>

TABLE 8  Critical Appraisal Skills Programme\textsuperscript{18} questions regarding data analysis: percentage of ‘yes’ responses to item, inter-rater agreement (kappa) and correlation with global rating of study quality

<table>
<thead>
<tr>
<th>CASP question</th>
<th>Kappa</th>
<th>% ‘Yes’</th>
<th>(r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is it clear how the analysis was done?</td>
<td>0.55</td>
<td>71.7</td>
<td>0.00</td>
</tr>
<tr>
<td>It is clear how the categories/themes were derived from the data?</td>
<td>0.48</td>
<td>67.1</td>
<td>0.00</td>
</tr>
<tr>
<td>Is there adequate description of the analysis?</td>
<td>0.53</td>
<td>62.8</td>
<td>–0.03</td>
</tr>
<tr>
<td>Have attempts been made to feed results back to respondents?</td>
<td>0.58</td>
<td>14.5</td>
<td>0.01</td>
</tr>
<tr>
<td>Have different sources of data been compared where appropriate?</td>
<td>0.36</td>
<td>24.7</td>
<td>0.00</td>
</tr>
<tr>
<td>Was the analysis repeated by more than one researcher to ensure reliability?</td>
<td>0.70</td>
<td>43.1</td>
<td>–0.08</td>
</tr>
<tr>
<td>Overall, is the data analysis sufficiently rigorous to address the aims?</td>
<td>0.31</td>
<td>69.9</td>
<td>0.36</td>
</tr>
</tbody>
</table>

TABLE 9  Critical Appraisal Skills Programme\textsuperscript{18} questions regarding interpretation and presentation of data: percentage of ‘yes’ responses to item, inter-rater agreement (kappa) and correlation with global rating of study quality

<table>
<thead>
<tr>
<th>CASP question</th>
<th>Kappa</th>
<th>% ‘Yes’</th>
<th>(r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is it clear whether researchers critically examined own role, bias, influence?</td>
<td>0.32</td>
<td>19.3</td>
<td>0.14</td>
</tr>
<tr>
<td>Is the relationship between researchers and participants adequately considered?</td>
<td>0.25</td>
<td>14.6</td>
<td>0.18</td>
</tr>
<tr>
<td>Were the findings explicit and easy to understand?</td>
<td>0.20</td>
<td>93.5</td>
<td>0.22</td>
</tr>
<tr>
<td>Are sufficient data presented to support the descriptive findings?</td>
<td>0.23</td>
<td>73.6</td>
<td>0.39</td>
</tr>
<tr>
<td>Are quotes numbered/identified?</td>
<td>0.71</td>
<td>48.3</td>
<td>0.24</td>
</tr>
<tr>
<td>Do the researchers explain how the data presented were selected?</td>
<td>0.52</td>
<td>17.2</td>
<td>0.10</td>
</tr>
<tr>
<td>Do researchers say how they developed their conceptual interpretations?</td>
<td>0.23</td>
<td>27.4</td>
<td>0.44</td>
</tr>
<tr>
<td>Are negative, unusual or contradictory cases presented?</td>
<td>0.19</td>
<td>36.1</td>
<td>0.28</td>
</tr>
<tr>
<td>Is there adequate discussion of evidence for/against interpretations?</td>
<td>0.10</td>
<td>28.3</td>
<td>0.35</td>
</tr>
<tr>
<td>Overall, are you confident that all the data were taken into account?</td>
<td>0.25</td>
<td>60.0</td>
<td>0.37</td>
</tr>
</tbody>
</table>

derived from the two first assessors, and the second from the six second assessors. Consequently, it should be noted that the ratings of the two first assessors had a greater influence on the aggregated results than the ratings of the six second assessors, as each of the first assessors rated approximately three times as many papers as each of the second assessors.

Tables 6–10 show the percentage of pooled ratings that indicated that the assessors replied affirmatively to each of the CASP\textsuperscript{18} questions. The assessors answered positively to most questions about design and sampling (see Table 6); reporting of the setting of data collection and how data were recorded (see Table 7); clarity about how the analysis was conducted (see Table 8);
satisfactory presentation of data and findings (see Table 9); and the value of the studies (see Table 10). However, few studies provided details such as justification of sample size, explanation of refusals to take part, how the research was explained to the participants or how the data presented were selected. Nonetheless, most papers were given positive ratings regarding the global evaluations (see emboldened question on the bottom row of each table) of their sampling strategy, data collection and data analysis. There was, however, little inter-rater agreement concerning which papers could not be rated positively on these questions. This pattern of findings may largely reflect the publication process, whereby papers would not normally be accepted unless the basic design, collection and presentation of data and value of the study were generally considered satisfactory, but space limitations may have prevented authors from providing more detail of the research process. It is perhaps of more concern that few studies appeared to have engaged in flexible or iterative data collection, reflexive consideration of the role of the researchers and their relationship to the participants, feeding back results to participants, or consideration of contradictory evidence or alternative interpretations. However, it must be noted that not all of these procedures for increasing the credibility and rigor of the research process would be appropriate for every study or would have been expected over the period of publication.

Correlations with ratings of the overall value of the papers

Ratings of the overall value of the papers had an approximately normal distribution, and so correlations were calculated between this item (see bottom line of Table 10) and all the other CASP18 items, in order to determine which aspects of the studies assessed by the CASP18 questions were associated with positive global evaluations (see final column of Tables 6–10). All those items that were significantly correlated with the global evaluation were then entered into a stepwise multiple regression (Table 11) in order to determine which items independently predicted the global evaluation of overall value.

These results show that the assessors’ global ratings of the value of the study were, unsurprisingly, associated with their judgements of its importance and contribution to knowledge. More significantly, global evaluations were related to positive ratings of a clear description of how conceptual interpretations were developed, adequate discussion of evidence for and against the interpretations presented, and a method of data collection appropriate to the aims of the research. Papers that had clearly identified quotations and whose findings were presented in an explicit and easy to follow manner were also more likely to be positively rated.

A later date of publication was correlated with more positive global evaluations ($r = 0.24$, $p < 0.001$). Additional features of the paper weakly associated ($p < 0.05$) with more negative global evaluations were type of publication other than peer-reviewed article and a later date of appraisal (possibly indicating papers identified by means other than electronic database searching).

### Table 10

<table>
<thead>
<tr>
<th>CASP question</th>
<th>Kappa</th>
<th>% ‘Yes’</th>
<th>$r$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is there descriptive, conceptual or theoretical congruence with other work?</td>
<td>0.00</td>
<td>86.9</td>
<td>0.27</td>
</tr>
<tr>
<td>Are the findings of this study transferable to a wider population?</td>
<td>–0.02</td>
<td>71.1</td>
<td>0.38</td>
</tr>
<tr>
<td>Does the study add to knowledge or theory in the field?</td>
<td>0.38</td>
<td>69.7</td>
<td>0.63</td>
</tr>
<tr>
<td>How important are these findings to practice?</td>
<td>0.42</td>
<td>N/A</td>
<td>0.74</td>
</tr>
<tr>
<td>What is your overall view of this study?</td>
<td>0.42</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

N/A, not applicable.
Discussion

The analysis of the quantifiable data from the appraisal process indicates that inter-rater agreement was rather low. Some of this unreliability was probably due to the poor reporting of research methods in papers, which meant that reviewers were often giving a qualified ‘yes’ or ‘no’ which is not reflected in this quantitative analysis. Our findings concur with those of Dixon-Woods, who found ‘slight’ levels of agreement when comparing three approaches with appraisal: CASP, the quality framework devised by the NatCen and unprompted judgement by two experienced qualitative researchers. This research also found that using a structured approach did not result in a higher level of agreement.

As we progressed with our study, it became clear to us that, that despite having conducted very detailed appraisals, we were not excluding papers on the basis of quality. If we were to conduct another synthesis in the future, therefore, we would be unlikely to conduct such a detailed appraisal beforehand. We would not, however, recommend abandoning appraisal altogether because it has advantages in terms of encouraging the reviewers to read the papers carefully and systematically, and serves as a reminder to treat the papers as data for the synthesis.

We did not ultimately find our modified version of the CASP tool to be especially helpful as an appraisal framework prior to a synthesis. Although it did facilitate a structured reading of papers, the questions it poses place a good deal of emphasis on methodological quality. From the point of view of a synthesis, it is conceptual quality that is most important, as was highlighted in our quantitative analysis of what items contributed most to assessors’ global ratings of the value of the study. As we observed, particularly in the case of the RA synthesis, there appears almost to be an inverse correlation between methodological quality and the quality of insight and theory-building displayed in many papers. This is, in part, related to the often inadequate reporting of qualitative research methods in the past. The current requirements of funding bodies, ethics committees and publishers for greater transparency in these matters should ensure that this is less of a problem with more recent work, but it leaves the qualitative synthesiser with a dilemma about whether or not to include older ‘classic’ papers for which there may be almost no detail given about how the research was conducted. In these cases, we would suggest that the methodological integrity of the work might have to be assumed. If the work concerned turns out to be very difficult or impossible to synthesise, then that may be a strong indication that it was not well grounded empirically.

### Table 11

<table>
<thead>
<tr>
<th>Step</th>
<th>Predictor variables</th>
<th>Adjusted R²</th>
<th>Change in R²</th>
<th>F</th>
<th>F Change</th>
<th>Beta</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>How important are these findings to practice?</td>
<td>0.49</td>
<td>0.490</td>
<td>99.9</td>
<td>0.44</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Do researchers say how they developed their conceptual interpretations?</td>
<td>0.59</td>
<td>0.104</td>
<td>26.4</td>
<td>0.23</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Does the study add to knowledge or theory in the field?</td>
<td>0.63</td>
<td>0.049</td>
<td>14.1</td>
<td>0.20</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Is there adequate discussion of evidence for/against interpretations?</td>
<td>0.66</td>
<td>0.028</td>
<td>8.6</td>
<td>0.13</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Are quotes numbered/identified?</td>
<td>0.67</td>
<td>0.014</td>
<td>4.6</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Overall, do you consider that data collection addresses the research aims?</td>
<td>0.68</td>
<td>0.015</td>
<td>4.9</td>
<td>0.12</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Were the findings explicit and easy to understand?</td>
<td>0.69</td>
<td>0.013</td>
<td>4.3</td>
<td>0.12</td>
<td></td>
</tr>
</tbody>
</table>

F, F-sampling distribution.

a \( p \leq 0.001 \)

b \( p \leq 0.01 \)

c \( p \leq 0.05 \)
We would endorse the stance taken by Sandelowski et al. and Dixon-Woods et al. that where possible it is important to expose and exclude papers that are, methodologically speaking, ‘fatally flawed’. Beyond that, what is required is an appraisal tool that aids the identification of findings, interpretations and concepts. The consensus within our research team was that any future appraisal process could be much shorter, with just a few key questions. We would retain the two initial screening questions about the relevance of the paper to the topic of the synthesis, and the question about whether or not qualitative methods of data collection and analysis were employed. Beyond that, we would include questions about the clear identification of the aim and objectives of the research; whether or not the data were collected in a way that addressed the research aim; whether or not the data analysis was sufficiently rigorous to address the aims; confidence that all the data were taken into account; whether or not there were enough data presented to support the interpretations made; and whether the paper demonstrated theoretical insight, novel findings or perspectives. In addition, we would include questions that enabled the data for the synthesis to be extracted and recorded.

The exercise of judgement is crucial to a number of stages of the synthesis. This includes making decisions about what constitutes a ‘fatal flaw’ as well as decisions about which papers should and should not be entered into the synthesis on the grounds of relevance. We would suggest that teamwork facilitates the exercise of sound judgement, as does a high level of qualitative research expertise.

Ultimately, the purpose of qualitative synthesis is to achieve a greater level of understanding and insight into particular phenomena and to generate and test theory. This is very different from quantitative synthesises in health care, where the objectives are to establish whether or not an intervention does more good than harm and to calculate an effect size. In the latter case, including biased findings or giving equal weight to all study findings irrespective of their methodological quality may result in an incorrect assessment of an intervention’s effectiveness, with potentially deleterious consequences for human health. The inclusion of poorer quality qualitative research in a synthesis is unlikely to be as damaging. As can be seen in the two synthesises reported in Chapters 6 and 7, those studies whose findings were slight contributed minimally to the synthesis. When describing their meta-ethnographic method, Noblit and Hare did not advocate any form of quality appraisal prior to a synthesis. Instead they cited Hunter et al., who argued against excluding ‘methodologically deficient’ studies, and they suggested that ‘the worth of studies … is determined in the process of achieving a synthesis.’ The lack of consistency in judgements about the quality of qualitative research that we and others have identified, and the evidence garnered thus far that the inclusion of poorer quality studies is unlikely to have a very distorting impact on a qualitative synthesis, suggest that this is wise counsel.
Chapter 5
Synthesis methodologies

Introduction

In this chapter, we describe the methods used to conduct our two syntheses. Before embarking on the syntheses, each subgroup – one for the medicine-taking synthesis and the other for the RA synthesis – conducted a preliminary study to assess the reproducibility of the synthesis process. We present these preliminary syntheses first, before going on to describe, in some detail, the methods employed to conduct the two full syntheses.

Preliminary syntheses

Concerns are frequently expressed about whether or not qualitative research can be subjected to an assessment of its reproducibility. This area is heavily contested, with extremes of opinion taking the view either that reproducibility cannot be expected as different qualitative researchers will interpret the same papers differently, depending on their perspectives and background, or that reproducibility must be shown for such work to be taken seriously alongside quantitative research. There are many positions in between. We investigated the degree to which these syntheses might be comparable or reproducible.

Each of the subgroups managed the process of searching and selecting relevant articles. Four items (papers or book chapters) were identified for each preliminary synthesis, either by chronology (RA synthesis) or by disease area (medicines synthesis). Each of the two lead researchers (PP and GDW) then synthesised the four RA papers and the four medicines papers. They did this blind, i.e. without any contact between them to discuss the methods to be employed, emergent findings or methods of presentation. PP had completed a synthesis previously, but GDW had not. Each researcher submitted his or her synthesis of both sets of papers to the overall lead researcher (RC).

Medicine-taking (PP lead)

PP’s synthesis

The four papers134–137 focused on antihypertensive medications, with the aim of describing the experiences of patients taking these medications and providing reasons for not taking medicines as prescribed. For PP, the first published, by Morgan,134 was the ‘index’ paper against which the others were compared. Morgan’s134 publication was a book chapter, and the others were articles published in journals: nursing (two135,136) and clinical (one137). PP then summarised and combined the findings from each of the papers around 13 issues that she felt emerged from the papers, some of which were inter-related. PP described the combined findings on issues such as the extent of taking and not taking medications as prescribed, lay testing of medication, and characteristics of patients taking and not taking medications as prescribed. In each case, she summarised the common findings from the four papers and indicated contrary or additional findings from particular papers. She began the process of exploring ‘higher order’ concepts, for example noting that several papers indicated that patients appeared to purposefully not take medications, although this was not overtly stated.
PP produced a ‘map’ (Figure 4) that explored links between the issues identified, using labels to indicate which papers contained the categorisations, and attempted to coalesce findings into common labels. For example, in one instance, a finding that patients did not take hypertensive medication as prescribed because they wished to avoid certain side effects when engaged in a particular activity was described by the authors as ‘incidental non-adherence’.136 This was reclassified by PP as ‘purposeful non-adherence’136 as specified in another paper.

**GDW’s synthesis**

GDW indicated that the Morgan134 chapter would be the ‘index’ paper. GDW described each paper in turn in some detail, including the sample, methods of the study and the findings. He noted the main concepts emerging and issues that seemed particularly salient (e.g. ‘leaving off’ medication in Morgan’s chapter). At the end of each description of the paper, GDW listed key findings and associated interpretations. As the chapter by Morgan was the ‘index’ paper, GDW considered the findings of the other studies in the context of this first paper. These findings were compared with those from the index paper and presented in terms of those that were similar to, and those different from, the key findings that he elucidated from Morgan’s chapter. He termed this a ‘translation’ of the findings from each of the papers into those preceding. This also allowed the exploration of ideas emerging from later papers to be considered in the index paper – for example the related importance of blood pressure monitoring.

At the end of the description of the papers and their findings, GDW provided a ‘lines-of-argument’ synthesis, indicating that the four studies134–137 were essentially concerned with the same major issues about why people did or did not take the medications. He indicated that the studies did not refute each other, and that small differences between the studies related to the aims and samples in them. He presented interpretations emerging from the studies of relevance to clinical practice, as intended in most of the studies (Box 6). GDW also went on to observe what he regarded as a tension in the expression of the synthesis whereby it appeared that, on the one hand, those who adhered to their antihypertensive therapy took a passive approach, but that, on the other hand, those who experienced difficulties adhering used strategies to encourage their adherence, suggesting a proactive approach. GDW noted that, were he to have continued with the synthesis, he would have employed a grounded theory approach and searched for studies that could shed further light on this issue.

**Similarities/differences between the two medication syntheses**

The researchers undertook the syntheses in slightly different ways. Both identified Morgan’s134 chapter as the index paper, but then PP presented the work in terms of similarities and differences between the papers in relation to issues emerging from them, whereas GDW attempted to combine the findings of later papers with those preceding. Their written material therefore looks quite different, but the content, in terms of the major findings from the syntheses, is broadly similar. Both indicate that the papers are concerned with the factors affecting whether and why people do or do not take antihypertensive medications. They mention ‘active’ (GDW) or ‘purposeful’ (PP) non-compliance, which are different terms for the same concept. Both indicate that there were no great differences between the papers in terms of the key findings, although differences in samples, aims and publication format did lead to some different aspects, particularly of reasons for taking or not taking medications.

The process of synthesising was undertaken differently by the researchers and so the work is presented differently. GDW described his method in more detail and more explicitly, including taking into account the methods and samples of the papers. This difference is likely to be related, in part at least, to the fact that GDW was less familiar with undertaking a qualitative synthesis, and with this particular topic, than PP. This was GDW’s first attempt at a synthesis, and he was aware that he was not going to go on to complete this particular topic. He felt it important to
Factors influencing whether or not people take medicine as prescribed/cost–benefit analysis

- Lay evaluation, testing of drug to determine whether (1) necessary,134–137 (2) effective134,136,137 and (3) safe136,137
- Information from media, health-care providers, others138
- Worries regarding drug: current/future adverse effects,134 mix with alcohol,134 taking long term,134 fear of dependency,134 fear of addiction134
- Fear of the illness or its symptoms134,135
- Attitude towards doctors137
- Doctor–patient relationship137
- Availability of, and faith in, alternatives, e.g. herbal medicines134,135
- Culture as a filter134
- Occurrence of side effects137
- Desire to reduce drug intake,134 reduce risk of adverse effects,134,137 dependency,134 addiction134
- Fear of the illness or its symptoms134,135

No decision
Simply attempt to take drugs as prescribed. Claim never considered alternative
Could describe as ‘non-purposeful adherence’, to extend Johnson et al.’s136 categories
Less involved in care, passive, indifferent – qualities not in line with demands of informed consent and shared decision-making137

‘Purposeful adherence’136
Those who decide to take drugs as prescribed.
Includes both ‘stable’ and ‘problematic’ adherents134

But other factors affect adherence
‘Patterned adherence’ – the necessary pre-conditions of adherence136
- Physical, financial access to drugs136
- Cognitive ability to manage drugs136
- Physical ability to take and ingest drugs136
‘Incidental non-adherence’ – the incidental reasons why adherence might fail136
- Interruption of routine136
- Lack of reminders136
- Forgetting136
- Away from home136

Recommendations
Doctors should help empower patients137
Doctors should personalise treatment135

‘Purposeful non-adherence’136
Those who decide not to take drugs as prescribed
Majority of ACs but minority of whites,134 less than half Svensson et al.’s137 sample and a quarter of Johnson et al.’s136 sample
They attempt to gain control, minimise impact on daily life, take active part in decision-making. This associated with improved therapeutic outcome and in line with health-care objectives137

Patterns of medicine use can evolve over time134

Ways of not taking drug as prescribed
- Using symptomatically rather than preventively134
- Reducing doses134
- Taking drug holidays134
- To avoid side effect if doing specific activity136
*Reclassified: Johnson et al.,136 had this as ‘incidental non-adherence’, but it is clearly ‘purposeful non-adherence’

describe the papers in detail and the process he undertook, both to gain greater familiarity with this topic but also to check that he was doing this in the way that the group wanted. PP, on the other hand, had completed previous syntheses and had more confidence in her process. It is probably also the case that PP’s ‘map’ was developed in detail to summarise the findings of the papers because she knew she would need this as the basis of her longer synthesis.

As indicated above, there was similarity between the two researchers in terms of interpretation. There were, however, some differences of detail. Most commonly, these were expressed in terms of differences in categorisations when attempting to combine concepts between papers. For example, in probably the starkest difference of interpretation, PP and GDW differed over whether to combine Morgan’s134 ‘problematic adherents’ with Johnson et al.’s136 ‘adherents’ or ‘non-adherents’. Interestingly, both felt that Morgan’s134 ‘stable adherents’ could be combined with Johnson et al.’s136 ‘purposeful adherents’. This difference most likely occurred because the concept ‘problematic’ contains the intention to both comply and not comply, and thus overlaps with both of Johnson et al.’s136 categories. Such differences are small and related to the descriptive detail provided by the original authors of their concepts, which have to be reinterpreted by the synthesiser and compared with each other.

In summary, these pilot syntheses indicated some difference in approach by the two researchers. This may well be related to their experience and involvement in this work – PP had completed such work previously and was going on to complete this synthesis, whereas GDW had not done this before and was not going to continue with this topic. In terms of the accessibility of the approach, GDW’s is more explicit (which was helpful to the group), but PP’s diagram is easier to follow and a good basis from which to complete the remainder of the synthesis.
**Rheumatoid arthritis (GDW lead)**

**PP’s synthesis**

PP began with a short summary of the publications to be synthesised. She felt that they fell into two groups: two were theoretical explorations of chronic illness, using RA as an example, whereas the other two focused on the day-to-day experiences of living with RA. PP elucidated a number of key categories from the papers and then described each one, indicating how it was presented in each of the papers. PP indicated whether or not the concept was explicitly mentioned in each paper and then described how it was presented. For example, the concept 'normalising' was identified by two of the original authors explicitly and in terms of various activities in another. PP then drew several diagrams that placed the experience of RA on scales, with benefits and costs of normalising and renormalising presented as being in balance (Figure 5 and 6). There are then two tables (Tables 12 and 13), one presenting aspects of the key...

---

**FIGURE 5** Normalising and costs of normalising (pilot RA synthesis – PP).

**NORMALISING**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Covering up</td>
<td>Wiener</td>
</tr>
<tr>
<td>Keeping up</td>
<td></td>
</tr>
<tr>
<td>Pacing</td>
<td></td>
</tr>
<tr>
<td>Keeping going</td>
<td>Locker</td>
</tr>
<tr>
<td>Pacing</td>
<td></td>
</tr>
<tr>
<td>Normalising</td>
<td>Bury</td>
</tr>
</tbody>
</table>

**COSTS OF NORMALISING**

<table>
<thead>
<tr>
<th>Cost</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain, exhaustion</td>
<td>Wiener</td>
</tr>
<tr>
<td>Justifying inaction on bad days</td>
<td></td>
</tr>
<tr>
<td>Severe pain</td>
<td>Locker</td>
</tr>
<tr>
<td>Pain, loss of energy, effort</td>
<td>Bury</td>
</tr>
<tr>
<td>ordinary activities conscious and deliberate</td>
<td></td>
</tr>
</tbody>
</table>

**FIGURE 6** Renormalising and costs of renormalising (pilot RA synthesis – PP).

**RENORMALISING**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjusting to reduced activity</td>
<td>Wiener</td>
</tr>
<tr>
<td>Lowering expectations</td>
<td></td>
</tr>
<tr>
<td>Redefining limits</td>
<td></td>
</tr>
<tr>
<td>Accepting need for help</td>
<td></td>
</tr>
<tr>
<td>Repairing biographical disruption (by rethinking biography)</td>
<td>Bury</td>
</tr>
<tr>
<td>(rethinking self-concept, rethinking self-relationships)</td>
<td></td>
</tr>
<tr>
<td>Imposing meaning</td>
<td></td>
</tr>
<tr>
<td>Mobilising resources</td>
<td></td>
</tr>
<tr>
<td>Narrative reconstruction</td>
<td>Williams</td>
</tr>
</tbody>
</table>

**COST OF RENORMALISING**

<table>
<thead>
<tr>
<th>Cost</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accepting the need for help makes it less possible to appear ‘normal’</td>
<td>Wiener</td>
</tr>
<tr>
<td>Dependency</td>
<td></td>
</tr>
<tr>
<td>Loss of control over activities</td>
<td></td>
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<tr>
<td>Loss of finance</td>
<td>Locker</td>
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<tr>
<td>Loss of dignity</td>
<td></td>
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<tr>
<td>Loss of power</td>
<td></td>
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<tr>
<td>Loss of role</td>
<td></td>
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<tr>
<td>Acceptance of disabled identity</td>
<td></td>
</tr>
</tbody>
</table>

*Related to resources.*
Synthesis methodologies

TABLE 12 Uncertainty (pilot RA synthesis – PP)

<table>
<thead>
<tr>
<th>Acute flare-up</th>
<th>Author</th>
<th>Uncertainty</th>
<th>Author</th>
<th>Remission/relief</th>
<th>Author</th>
</tr>
</thead>
<tbody>
<tr>
<td>When will it strike?</td>
<td>Weiner(^{138})</td>
<td>Uncertainty about flare-ups and remission makes illness difficult to manage</td>
<td>Locker(^{139})</td>
<td>How long will it last?</td>
<td>Weiner(^{138})</td>
</tr>
<tr>
<td>How long will it last?</td>
<td></td>
<td></td>
<td></td>
<td>Timing?</td>
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<tr>
<td>How will it manifest?</td>
<td></td>
<td>Constant self-monitoring necessary</td>
<td></td>
<td>Remission mitigates against acceptance of disabled status</td>
<td></td>
</tr>
<tr>
<td>What (if anything) will control symptoms?</td>
<td>Locker(^{132})</td>
<td>Continuous testing of tolerance to activity is necessary</td>
<td></td>
<td>Hope for remission balanced against dread of disease progression</td>
<td></td>
</tr>
<tr>
<td>How much pain will there be?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>How long will pain last?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Which joints will be affected?</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>When will it happen?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Which treatments (if any) will work?</td>
<td>Bury(^{140})</td>
<td></td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

TABLE 13 Medicine and inner resources (pilot RA synthesis – PP)

<table>
<thead>
<tr>
<th>Medicine</th>
<th>Author</th>
<th>Limitations of medicine</th>
<th>Author</th>
<th>Inner resources</th>
<th>Author</th>
</tr>
</thead>
<tbody>
<tr>
<td>Is a key cultural resource</td>
<td>Bury(^{140})</td>
<td>People realise limits of medical knowledge regarding the cause and nature of RA</td>
<td>Locker(^{139}) and Bury(^{140})</td>
<td>Patients left to draw own conclusions about whether or not RA is inherited</td>
<td>Bury(^{140})</td>
</tr>
<tr>
<td>Referral to RA clinic and diagnosis gives relief and imposes order on symptoms</td>
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<tr>
<td>Diagnosis justifies people’s symptoms and behaviour</td>
<td>Locker(^{139}) and Bury(^{140})</td>
<td>Medicine cannot answer questions regarding aetiology</td>
<td></td>
<td>Patients have to construct own theories. Might search past for reasons or search family/personal biographies, or reinterpret doctor’s explanation</td>
<td>Locker(^{139}) and Bury(^{140})</td>
</tr>
<tr>
<td>People hope that medicine will provide information on how to manage RA, aetiology, fluctuating pain levels, prognosis</td>
<td>Locker(^{139})</td>
<td>Often medicine cannot provide satisfactory answers</td>
<td>Locker(^{139}) and Bury(^{140})</td>
<td>People have to impose own meaning and certainty in whatever ways possible</td>
<td>Locker(^{139}) and Bury(^{140})</td>
</tr>
<tr>
<td>Medicine provides treatment</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>People hope it will help manage pain (because pain depletes energy, disrupts personal and social life, is difficult for others to understand, and is unpredictable and intangible, so it is difficult for sufferers to secure legitimacy)</td>
<td>Bury(^{140})</td>
<td>People realise that regimens are difficult to follow, not very effective</td>
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</tbody>
</table>

concept of uncertainty and the other focusing on issues related to medication use, its limitations and other resources which patients could draw on. PP focused the synthesis around the concepts of Weiner\(^{138}\) (the first and index paper) of normalising and renormalising, and synthesised the other papers around this. This produced the key concept of ‘achieving a balance’. She attempted to combine the papers around this concept, using a grounded theory approach, which appeared to work well.
<table>
<thead>
<tr>
<th>Concepts</th>
<th>Second-order interpretations</th>
<th>Third-order interpretations</th>
<th>Fourth-order interpretations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DISEASE CONDITIONS</strong>: elusive aetiology; pain; crippling; absence of predictability; variability in progression and areas of involvement; lack of visible signs accompanying painful symptoms</td>
<td>Variability in the disease conditions of RA makes it difficult for people to maintain normal activities and be believed by others (e.g., family members, workmates)</td>
<td>(a) The essential burden in RA relates to the unpredictable nature of the disease (b) Searching for ‘personal’ aetiological explanations is a way of coping with this unpredictability</td>
<td></td>
</tr>
<tr>
<td><strong>IMPERATIVES</strong>: to identify the causes of arthritis/pain and disability; learning to live with it; avoiding embarrassment; maintaining a normal life and social roles</td>
<td>There is an evident tension between these IMPERATIVES and DISEASE CONDITIONS</td>
<td>Socially constructed norms, imperatives or expectations are a major source of the problems faced by people with RA</td>
<td>Coping with RA is as much about managing symptoms as it is about managing changing social expectations, norms and relationships</td>
</tr>
<tr>
<td><strong>BURDENS</strong>: tolerating variable uncertainty; dread of dependency or invalidity; previously taken for granted actions require conscious and deliberate physical and psychological effort; biographical disruption; problems of legitimating changed behaviour; stigma; frustration; insecurity about dangers of everyday life; costs of treatment may outweigh the benefits</td>
<td>(a) The burdens of RA centre on changes in the relationship between self, environment and others (b) These changes may be as burdensome as the symptoms of the disease itself (mainly pain)</td>
<td>In addition to the symptomatological burdens of RA, individuals face other burdens centred on changes in their relationships with their own physical bodies, other people and the medical system</td>
<td></td>
</tr>
<tr>
<td><strong>STRUCTURAL INFLUENCES</strong>: material resources; the limits of medical knowledge; emotional and social support networks; the organisation of paid employment; knowledge</td>
<td>(a) The degree to which the burdens of arthritis are onerous varies according to the social and psychological characteristics of individuals (b) The organisation of medicine and the workplace are powerful influences</td>
<td>The everyday impact of RA is exacerbated by the organisation of society</td>
<td></td>
</tr>
<tr>
<td><strong>PSYCHOLOGICAL COPING STRATEGIES</strong>: hope of remission; dread of progression; accepting it; living with it; fighting it; keeping going/Carrying on; mastering pain; withdrawal from others; reconstructing a sense of order; super-normalising (see below)</td>
<td>Psychological coping strategies involve ‘juggling’ opposites (hope/dread, accepting/fighting, carrying on/withdrawal)</td>
<td>Psychological coping strategies are limited by the unpredictable nature of the disease</td>
<td></td>
</tr>
<tr>
<td><strong>SOCIAL COPING STRATEGIES</strong>: normalising (pacing/thinking it out, covering up, keeping up); renormalising (adjusting to reduced activity; lowering expectations, developing a new set of norms for action (have to be constantly learned and released); eliciting help; balancing the options; avoiding inappropriate behaviour; planning and routing</td>
<td>(a) “Learning how to pace activities and cope with greatly diminished resources is part of … becoming disabled” (Locker, p. 40)139 (b) “The body is defined by its relationship with the world of social action not in isolation from it” (Williams, p. 182)141</td>
<td>The ways that people with RA attempt to adjust to their illness are constrained by social norms and expectations about behaviour and social action</td>
<td></td>
</tr>
<tr>
<td><strong>CONSEQUENCES</strong>: a sense of loss in relation to social roles and activities; a change from a participant in social life to an observer of it; eliciting help may give rise to feelings of dependency and decreases the possibilities of covering up and keeping up; medical knowledge is both resource and constraint; in acute phases, strategies of accepting it and keeping going break down; different aspects of biography are linked to realign the past with the present</td>
<td>The strategies used to cope with RA are undermined by the conditions of the disease itself</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
GDW’s synthesis

The synthesis was reported in several explicit stages: a description of the methods and samples in each paper, along with the major findings; tables to present the combination of findings in particular categories; a textual report of the synthesis, pointing to similarities and differences between the papers; and finally a table (Table 14) presenting the original concepts and higher order concepts derived from the synthesis. The descriptions of the papers help to set the scene for the synthesis, describing the main aims of the research, its methods and the citation of some papers by the other authors. Two of the papers are indicated to be theoretical in nature,\textsuperscript{140,141} drawing on experiences of RA as an example, whereas the others are more concerned with the lived experience of the disease.\textsuperscript{138,139} GDW noted that a major concept in one paper,\textsuperscript{139} pain, was not mentioned in another, probably because of this different approach. GDW observed that one paper was described as very difficult to synthesise as it was so theoretical\textsuperscript{140} and commented that the methods of data collection and analysis were poorly described in the theoretical papers.\textsuperscript{140,141}

In the synthesis, it was made clear that two papers\textsuperscript{138,139} were closest to the heart of the aim of the synthesis, whereas the others used the example of RA in their theoretical expositions.\textsuperscript{140,141} On the whole, it was found that concepts were similar across the studies, although one paper\textsuperscript{141} was peripheral as it was concerned with the search for a cause of the condition rather than the experience of living with it. GDW reported the various burdens and coping strategies outlined by the authors, which are basically similar in concept. In the table of concepts (see Table 14), GDW focused on ‘disease conditions’ (symptoms, unpredictability), ‘imperatives’ (need to find cause of symptoms, learning to live with them), ‘burdens’ (tolerating uncertainty), ‘structural influences’ (resources such as medical and emotional), ‘coping strategies’ (psychological, social) and ‘consequences’ (loss and change). He produced this as a framework for the continuation of the synthesis.

Similarities/differences between the two arthritis syntheses

At the heart of both syntheses is the disruption to normal life by RA and the various strategies people employ to cope with it. Both researchers grouped the papers into two sets in the same way and noted the purpose of the papers identically. Both noted that all the papers referenced each other, with the exception of Williams,\textsuperscript{141} who refers only to Bury.\textsuperscript{140} The two researchers used different terms for the core concepts. PP used Wiener’s\textsuperscript{138} thesis of normalising and renormalising, and identified that the concepts of uncertainty and balance were crucial. GDW focused more on the burdens and imperatives caused by the disease, the strategies employed and resulting consequences. PP, perhaps because of her in-depth involvement in the medicines synthesis, also focused on how those with RA became aware of the limitations of medicine and the treatments that it provides and had to draw on their own inner resources when coming to terms with the disease and its effects on them.

In summary, there are some similarities between the syntheses, but also notable differences. GDW was trying to establish a basis from which to continue the synthesis, and so displayed the information in more detail and in several ways. GDW presented a very detailed description of his attempt to synthesise these papers, probably because of his need to make this a firm foundation for the complete synthesis. He synthesised Wiener’s\textsuperscript{138} and Locker’s\textsuperscript{139} work first, as these were similar in presentation and focus, then integrated Bury’s\textsuperscript{140} findings, and attempted to include something of Williams’\textsuperscript{141} work, although this proved difficult. PP decided to focus on the key concepts from the index paper by Wiener,\textsuperscript{138} integrating the other findings into these, and this allowed the development of the higher order concept of ‘balance’. It is clear from the tables presented by both researchers that they found some aspects in common across the papers, although the terminology used was dissimilar and the syntheses were somewhat different.
The full syntheses

After the pilot syntheses had been completed, PP and GDW proceeded to conduct a full synthesis in the topic areas of medicine-taking and RA, respectively. Both synthesis subgroups reviewed and discussed the two pilot syntheses. Their experience of the pilot syntheses did not affect the way in which they conducted the full syntheses, although they used their initial syntheses as a starting point for the full synthesis. The methodologies used to conduct the full syntheses are now described in some detail.

Noblit and Hare\textsuperscript{15} suggested that there are seven stages to conducting a synthesis:

1. topic selection
2. selecting and finding the studies
3. reading the studies
4. determining how the studies are related
5. translating studies into one another
6. synthesising the translations
7. expressing the synthesis.

We described in Chapters 3 and 4 how Noblit and Hare\textsuperscript{15}' first three stages of topic selection, literature searching and reading of the studies were conducted. As noted, we appraised our studies while reading them and part of that appraisal process required us to extract and outline in some detail the main findings and concepts for each of the papers. Thus, for us, Noblit and Hare\textsuperscript{15}' third stage (reading the studies) expanded into reading, appraisal and initial data extraction. The subsequent stages varied for each of the syntheses and are described separately below. As will be seen, they did not necessarily conform to the stages outlined by Noblit and Hare.\textsuperscript{15}

Medicines synthesis

Initial extraction of data from papers

The findings and concepts that we outlined for each of the studies constituted the raw data for the initial stages of the syntheses. (The modified CASP\textsuperscript{18} appraisal tool that we used had separate questions about findings and concepts.) For the medicines synthesis, the extracted findings and concepts were compared between the two independent reviewers for each of the papers. In the rare cases in which the second reviewer had identified a finding or concept that was missed by the first, this was included to ensure that the final record of findings and concepts was comprehensive. The process of extracting the findings and concepts inevitably involves a degree of organising and summarising; thus, to some extent, an initial process of interpretation was already under way at this stage, especially when organising descriptive findings that had not been interpreted. At this stage, it also became apparent that the distinction between findings and concepts was neither simple to make nor useful, and because it was felt to be unnecessarily complicated it was abandoned.

Organising studies into medicine groups

As noted above, Noblit and Hare\textsuperscript{15} suggested that the fourth stage of the synthesis might be determining how the studies are related, followed by (5) translating studies into one another, (6) synthesising the translations and (7) expressing the synthesis. However, Noblit and Hare\textsuperscript{15} used much smaller samples in their syntheses (two or three studies at the most, although the reports were probably quite lengthy) and, with 38 studies to synthesise, the need to organise the
studies in some preliminary way became imperative. For this reason, we began by organising the studies in what seemed to us to be the most immediately obvious way, into groups according to the medicine studied, and then within medicine groups, by the date of publication (Table 15). This gave us seven groups: antihypertensive medicine, which was also used for the preliminary

<table>
<thead>
<tr>
<th>Medicine group</th>
<th>Papers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antiretroviral therapy</td>
<td>Siegel and Gorey, 1997\textsuperscript{142}</td>
</tr>
<tr>
<td></td>
<td>Stone and Clarke, 1998\textsuperscript{143}</td>
</tr>
<tr>
<td></td>
<td>Erlen and Mellors, 1999\textsuperscript{144}</td>
</tr>
<tr>
<td></td>
<td>Halkitis and Kirton, 1999\textsuperscript{145}</td>
</tr>
<tr>
<td></td>
<td>Proctor et al., 1999\textsuperscript{146}</td>
</tr>
<tr>
<td></td>
<td>Siegel et al., 1999\textsuperscript{147}</td>
</tr>
<tr>
<td></td>
<td>Siegel et al., 2000\textsuperscript{148}</td>
</tr>
<tr>
<td></td>
<td>McDonald et al., 2000\textsuperscript{149}</td>
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<tr>
<td></td>
<td>Murphy et al., 2000\textsuperscript{150}</td>
</tr>
<tr>
<td></td>
<td>Johnston Roberts and Mann, 2000\textsuperscript{151}</td>
</tr>
<tr>
<td></td>
<td>Barton Laws and Wilson, 2000\textsuperscript{152}</td>
</tr>
<tr>
<td>Antihypertensives</td>
<td>Morgan, 1996\textsuperscript{134}</td>
</tr>
<tr>
<td></td>
<td>Van Wissen et al., 1998\textsuperscript{135}</td>
</tr>
<tr>
<td></td>
<td>Johnson et al., 1999\textsuperscript{136}</td>
</tr>
<tr>
<td></td>
<td>Svensson et al., 2000\textsuperscript{137}</td>
</tr>
<tr>
<td>Psychotropic medicine</td>
<td>Keljee and Beardsley, 1992\textsuperscript{153}</td>
</tr>
<tr>
<td></td>
<td>North et al., 1995\textsuperscript{154}</td>
</tr>
<tr>
<td></td>
<td>Barter and Cormack, 1996\textsuperscript{155}</td>
</tr>
<tr>
<td></td>
<td>Rogers et al., 1998\textsuperscript{156}</td>
</tr>
<tr>
<td></td>
<td>Angermeyer et al., 2001\textsuperscript{157}</td>
</tr>
<tr>
<td></td>
<td>Usher, 2001\textsuperscript{158}</td>
</tr>
<tr>
<td>PPIs</td>
<td>Boath and Blenkinsopp, 1997\textsuperscript{159}</td>
</tr>
<tr>
<td></td>
<td>Pollock and Grime, 2000\textsuperscript{160}</td>
</tr>
<tr>
<td>Asthma medicine</td>
<td>Adams et al., 1997\textsuperscript{161}</td>
</tr>
<tr>
<td></td>
<td>Prout et al., 1999\textsuperscript{162}</td>
</tr>
<tr>
<td></td>
<td>Buston and Wood, 2000\textsuperscript{163}</td>
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<tr>
<td></td>
<td>Walsh et al., 2000\textsuperscript{164}</td>
</tr>
<tr>
<td></td>
<td>Pradel et al., 2001\textsuperscript{165}</td>
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<tr>
<td>Miscellaneous medicines</td>
<td>Donovan and Blake, 1992\textsuperscript{166}</td>
</tr>
<tr>
<td></td>
<td>Dowell et al., 1996\textsuperscript{167}</td>
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<tr>
<td></td>
<td>Ersek et al., 1999\textsuperscript{168}</td>
</tr>
<tr>
<td></td>
<td>Atkin and Ahmad, 2000\textsuperscript{169}</td>
</tr>
<tr>
<td></td>
<td>Smith et al., 2000\textsuperscript{170}</td>
</tr>
<tr>
<td>Medicines in general</td>
<td>Roberson, 1992\textsuperscript{171}</td>
</tr>
<tr>
<td></td>
<td>Britten, 1996\textsuperscript{172}</td>
</tr>
<tr>
<td></td>
<td>Dowell and Hudson, 1997\textsuperscript{173}</td>
</tr>
<tr>
<td></td>
<td>Watson and Mitchell, 1998\textsuperscript{174}</td>
</tr>
<tr>
<td></td>
<td>Lumme-Sandt et al., 2000\textsuperscript{175}</td>
</tr>
</tbody>
</table>
synthesis (four studies); human immunodeficiency virus (HIV) infection medicine (11 studies); psychotropic medicine (six studies); asthma medicine (five studies); studies on medicine in general (five studies); a group of medicines for miscellaneous illnesses (five studies); and proton pump inhibitors (PPIs) (two studies). We then proceeded to translate studies into each other within each of the medicine groups.

Translating studies into each other within medicine groups

Noblit and Hare\(^\text{15}\) suggested that the process of translating findings into each other goes something like, ‘one case is like another, except that …’ (p. 38). We followed this process in a systematic way. For example, paper 1 in the psychotropic medicine group might have findings X, Y and Z. Paper 2 in the same group might have finding w (something new that was not in paper 1), findings x and y (findings similar to findings X and Y in paper 1) and nothing like finding Z from paper 1. So this would produce a synthesis of papers 1 and 2:

- finding w (from paper 2)
- findings X and x (from papers 1 and 2)
- findings Y and y (from papers 1 and 2)
- finding Z (from paper 1).

This synthesis of papers 1 and 2 would then be compared with paper 3 in the same way. Then the synthesis of papers 1, 2 and 3 would be compared with paper 4, and so on until all the studies within, for example, the psychotropic medicine group had been translated into each other. Then the process would begin again for the next medicine group, and so on. This process of translating the findings into each other is one of ‘reciprocal translation’, as described by Noblit and Hare.\(^\text{15}\)

The process of ‘reciprocal translation’ was employed because the majority of the studies are essentially about similar issues. Another possible method of translation is ‘refutational’,\(^\text{15}\) but in this case none of the findings from the studies refuted each other.

Determining how findings relate to each other within medicine groups

The translation of findings from the studies into each other for each of the medicine groups resulted in a raw textual synthesis for each of the groups. We then needed to know how the findings related to each other within each of the groups. Noblit and Hare (p. 28)\(^\text{15}\) wrote: ‘We think it makes sense to create a list of the key metaphors, phrases, ideas and/or concepts (and their relations) used in each account and to juxtapose them’. We did this by summarising the key findings for each of the illness groups so that they fitted on one page and then drew the relationships between them. This resulted in a ‘map’ for each of the groups (Figures 7–12). These ‘medicine maps’ summarised the key findings, the way in which they translated into each other and the relations between them, for each of the groups.

Determining how studies are related across the medicine groups

We now needed to know how the findings related to each other across the medicine groups. We determined this by comparing the medicine maps with each other, and developed a model that was able to encompass the findings from all of the studies in the synthesis. Some of North et al.’s\(^\text{15}\) findings did ‘fit’ with this, but their typology of relationships to benzodiazepines did not. Our model changed slightly as we progressed with the synthesis, but the final version is shown in Figure 13. Dowell and Hudson,\(^\text{17}\) authors of one of the studies in the synthesis, had developed a model of medicine that was similar in many ways, but which was unable to ‘hold’, or account for all the findings from all the studies in our synthesis. Our model provides a useful overall map for the synthesis process. It was an invaluable organisational aid during the synthesis process, but it may also prove to be a useful model for understanding the route by which people decide whether or not and how to take their medicines. It also illustrates that seemingly disparate studies may be put together to create something new. The model, however, provides only an overview.
Factors influencing whether or not people ‘take drugs as prescribed’ (TDAP)

- Effectiveness: patients find PPIs effective for pain relief and dealing with symptoms and allow them to lead a normal life.
- Most want to continue with treatment owing to perceived efficacy, but said would follow doctors’ advice regarding stopping/switching brands if necessary.
- Worries regarding adverse effects of PPIs rare, but some worries about potential to cause harm in the long term.
- Some worry that PPIs might mask symptoms of serious illness such as cancer.

Authors’ ideas about ‘authorised self-regulation’ (ASR)

- Patients should be encouraged to actively manage their illness by finding their own level of treatment.
- Doctors need to realise that many patients already do this.
- Self-regulation of PPIs more likely to lead to restraint than profligacy.
- Patients would self-regulate if doctors told them to do so.
- If authorised and open, doctors could guide patients and make it safe.
- ASR would make care more patient centred and help patients feel in control.
- (Would this be possible for all medications?)

Purposeful non-adherence

Reasons why patients do not take drugs as prescribed

- Reductions in dose perceived as indication that condition improving.
- Desire to minimise drug intake in general.
- As and when perceived necessary, at symptom onset only or if symptoms anticipated for some reason (e.g. going out for curry), or alternatively if the patient does not anticipate a need for it, then would miss a dose.
- Trying periods without PPIs in order to test effect – naive scientists.
- Some supplemented PPIs with OTC drugs such as Gaviscon or Tagamet.

Doctor–patient relationship (doctors’ response to self-regulation)

- GPs discouraged self-management if this involved taking increased doses.
- Few patients encouraged by GPs to manage own regimen.
- GPs ambivalent about patient self-management as a strategy.

Doctors’ strategies for rationing PPIs

- Do not initiate treatment.
- Substitute different treatment or cheaper brand or reduce dose or double switch (cheaper brand and reduction in dose). Latter confusing to patients as cannot assess efficacy if unaware of reduction in dose.

Professional views of PPIs

- Clinical uncertainty regarding long-term use of PPIs.
- They allow continuation of ‘unhealthy’ lifestyle which could have long-term health implications.
- They are expensive.
## Factors influencing whether or not people take medicine as prescribed

- The nature of the regimen (HAART regimen requires strict adherence and schedule of doses affects eating, sleeping, daily routines, social and recreational activities, sex, working life, social relationships, care-giving roles, studying, weight and thereby dominates life)\(^{142–152}\)
- Type of medication (HAART has frequent doses, high number of pills per day, and off-putting taste, smell, size and shape)\(^{145,146,151,152}\)
- Gender (women more sceptical regarding safety and efficacy of HAART than men, regimen impacts more on women than on men)\(^{142,149,151}\)
- Perceived efficacy of medication\(^{142–144,148–151}\)
- Lay evaluation of medication for efficacy (with HAART subjective rather than objective indicators given more weight)\(^{143,147–150}\)
- Adverse effects (with HAART a/effects nasty, impacted on QoL and instilled distrust in medication and led to not taking drugs as prescribed as medication experienced as making you ill rather than better – again, subjective rather than objective indicators more important)\(^{142–152}\)
- Being reminded of illness status\(^{142–146,151}\)
- Quality of relationship with health-care providers\(^{143–144,147,150}\)
- Perception of health-care providers (some women very mistrustful of doctors’ motives for prescribing HAART)\(^142\)
- For significant others (staying healthy to raise family/wish to extend life for others’ sake)\(^{143,149}\)
- Illness severity (perception that if don’t take HAART will die)\(^144\)
- Understanding of how medication works\(^{143,152}\) (Not sure there is actually evidence that it affects adherence. PP)
- Informal networks/peers could influence either way\(^142,143\)
- Acceptance of HIV infection status (accepting it oneself and being able to disclose it to others)\(^{152}\)

### Purposeful or non-purposeful adherence

#### But other factors affect adherence

**Patterned adherence**
- Pharmacy running out of medicines for refills\(^{146,150}\)
- Cannot afford prescriptions\(^150\)
- Cannot afford food to take with drugs\(^148\)
- Living in institution, so no control of medicines\(^143\)

**Incidental non-adherence**
- Forgetting\(^{143,144,148,151,152}\)
- Change in routine\(^{143,150,152}\)
- Forgetting refills\(^146\)
- Stress\(^146\)
- Focus on getting fix\(^152\)
- Being busy\(^148,151\)
- Holidays, being away from home\(^151\)
- Avoid taking medicines in public for fear of disclosure\(^{143,145,148,150–152}\)

### Purposeful non-adherence (includes stable purposeful non adherents)

**Ways of not taking drugs as prescribed**
- Drug holidays to cleanse body\(^{143,144,152}\)
- Taking medicines separately to avoid a/effects\(^144\)
- Reduce doses or stopping medicines to reduce a/effects\(^{142,147–149}\)
- Altering regimen to make it more practicable.\(^{142,147–149}\) (This latter group could be ‘stable purposeful non adherents’, i.e. do not adhere and do not worry about it either. PP)

**Things that enhance adherence to regimen**
- Mechanical gadgets, e.g. pill cases, beepers, alarms\(^{143,146,150}\)
- Keeping medicines in view, written reminders\(^{143,145,150}\)
- Planning ahead, carrying medicines with you and food and water to take them with\(^{143,145,150}\)
- Reminders from significant others, social support\(^{143,145,150}\)
- Routines, schedules, habit\(^{143,145,150}\)
  - Easier, simpler regimen\(^150\)
  - Home delivery of prescription refills\(^150\)
  - Good communication with care providers\(^150\)

---

**FIGURE 8** Highly active antiretroviral therapy (HAART) synthesis. QoL, quality of life.
Factors influencing whether or not people take medicine as prescribed

- Fears regarding drug: of side effects,166,168 of becoming dependent on drug,166 of developing tolerance to drug,166,168 of addiction to drug,168 that drug will mask new pains168
- Reliance on drugs perceived as a weakness166
- Dislike of taking drugs in general/desire to minimise use of drugs166,168
- Fear of side effects could be outweighed by need for symptom relief166
- Lay evaluation to determine whether or not effective166,170 and whether or not any adverse effects170
- Would take drugs if made sense within belief system and did not conflict with lifestyle/self-image166
- Feeling that just have to live with pain168
- Whether drug regarded as effective166,170
- Whether or not drug affects function and ability to fulfill social obligations166,170
- Drug reminds self of illness166
- Availability of non-pharmacological methods/strategies for tolerating high pain without drugs166
- Drug marks out as different, threatens normality166
- Administration of drug disrupts valued social activities166
- Drug undermines cultural identity169
- Cannot see point as going to die anyway169
- Whether or not drug improves general well-being170
- Medication for mental-health problems is stigmatising170
- Perception that medicines for mental health problems used to control/make patients acceptable to society170

Knowledge and information

- Not enough information provided about prescribed drugs, their side effects and how to deal with side effects166,168,170
- Patients do not fully understand drug regimen166
- Patients do not just ask doctors for information, but consult other sources166
- Patients not sufficiently informed regarding therapeutic options166

Doctor-patient relationship

- Those who trusted doctor would accept almost any proposed prescription changes, but those who do not fiercely reject changes167
- Most had positive relationships with care providers, but some felt poor communication regarding drugs and minority felt had to ‘prove’ pain to their doctors168
- Those with poor relationship with doctor would not even discuss severe pain or treatment options with doctor168
- Desire for more involvement in decision-making regarding medication170
- Those with mental-health problems reported generally poor relationships with health professionals170

Ways of not taking drug as prescribed

- Use pump/take medicines less frequently166,169
- Disconnect needle at night169
- Hide ‘non-compliance’ from parents169
- Stop drugs completely166
- Take fewer tablets than prescribed166

Things about those who did not take

- Ntdap active not passive166
- Ntdap is a way of gaining control over illness166
- Ntdap common so hard to view as deviant166
- Ntdap not a problem for patients166

Things that enhance adherence

- Keep treatment within defined boundaries/specific times169

But other factors affect adherence

Patterned adherence

- Financial difficulties paying medical bills168
- Did not report pain as did not want to be seen as complainer so drugs not prescribed168
- Difficulties opening pill containers170

Incidental non-adherence

- Forgetfulness168

FIGURE 9 Miscellaneous medication (including for arthritis, cancer pain, mental-health problems, thalassaemia major and respiratory disease) synthesis. Ntdap, not taking drugs as prescribed.
Categories of medicine takers/rejectors

- Orthodox/active acceptors, believers/patient repertoire
  Those who gave orthodox accounts are more likely to be using medication long term, have chronic illnesses, describe taking drugs as prescribed, describe drug taking as routine, be respectful of/deferential to doctors, talk about medicines in taken for granted way, generally passive (cf. ‘non-purposeful adherents?’) Those described as passive acceptors relinquished control. Those giving the patient repertoire saw selves as patients and doctors as experts, medication taken for granted and considered neutral fact.

- Unorthodox/active users and rejectors, sceptics/self-help repertoire
  Those giving unorthodox accounts are more likely to be critical of medication, fearful of dependence on medicines, fearful of adverse effects, critical of doctors, generally active. Active users take control by modifying their regime. (cf. ‘purposeful non-adherents?’) Rejecters/rejecters reject medicine altogether (cf. rejecters/accepters in asthma synthesis?). Those using self-help repertoire preferred natural remedies, had strong negative views about medical drugs, did not obey doctors.

- Moral repertoire
  Those giving moral repertoire stressed only took a little medication, stressed that only used it responsibly and moderately, explained why needed drugs giving reasons beyond their control.

Doctor–patient relationship (challenge to medicines, occurs outside doctor’s surgery)

- Patients scolded by doctors for self-management change doctors rather than confront them
  Those giving unorthodox accounts do not reveal these to doctors. Unorthodox appear as orthodox in medical consultations so agenda is not addressed, but they regain control outside surgery by accepting or rejecting medicines.

Knowledge/information

- Scant knowledge regarding illness and medication
- Understanding illness and medicines linked to acceptance of medicines
- People seek information from informal networks, alternative healers, various sources, not only doctors

Factors influencing whether or not people take medicine

- Does regimen fit with personal goals
- Availability of effective alternatives
- Worry about medication
- Experimentation
- General dislike of taking medication
- Non-acceptance of illness
- Information

Purposeful non-adherence

Ways of not taking drugs as prescribed

- Reduced doses
- Stopped completely
- Took as felt necessary
- People set self-imposed limits

‘Purposeful adherence’ and ‘non-purposeful adherence’

But other factors affect adherence

Patterned adherence

- Some too poor to visit doctor, so stopped when ran out
- Some had no transport to get to doctor for refills
- Medication too expensive for some

Incidental non-adherence

- Some simply forgot to take drugs

Model of medicine-taking behaviour

Patients test treatment to see whether it works or not (some bypass testing process as already have firm views on medication) Testing results in rejecters, passive accepters, active users/modifiers Active users and passive users both take medication, but in different ways (cf. ‘stable and problematic adherents?’) Nevertheless, practical limitations still apply (cf. incidental non-adherence and patterned adherence) Problem may be reviewed and people may recenter process.

FIGURE 10 General medication synthesis.
Factors influencing whether or not people take medicine as prescribed

Acceptance of, or distancing from, asthma identity

Perceived effectiveness of need for medicine

Whether or not medication supports ordinariness or stigmatises

Worries regarding steroids, side effects of steroids, dependency on steroids

Lay evaluation/experimentation regarding necessity of taking drugs

Nature of the medicine itself: difficulty or ease of use of inhalers, convenient or inconvenient to use, speed with which it works, taste, confusion between preventers and relievers (among children)

Negative experiences such as hospitalisation

Benefits of asthma medication

Inhalers allow targeting of precise quantities of medication to lungs, are small, unobtrusive, easy to use, allow rapid onset of drug action and reduce side effects.

Inhalers not seen as strong medicine as inhaled (rather than pills/liquid) and not seen as systemic? (PP)

Asthma medicines seen as preferable to avoidance/(non-medical) preventive strategies

But because asthma medicines seen as so effective, hospital admission delayed even when serious

So possibly inhaler/reliever medicines over used, but preventer medicines underused (PP)

Purposeful adherence (or possibly non-purposeful adherence – not adherence)

Acceptors

Incorporate asthma into lives

Accept chronicity of asthma

Accept need to take both preventive and reliever medication

All carry reliever medication

Do not like preventers because fears regarding steroids, side effects, dependency and have tried to reduce use of preventers, but unsuccessful so have accepted and routinised their use

Medication seen as aid to normalisation

Operate policy of full disclosure; to deny asthma would be to deny important part of selves. Asthma identity not in conflict with other social identities

Asthmatics viewed positively

Model of asthma is nearer medical understanding

Purposeful non-adherence (of preventers – and over use of relievers)

Deniers/distancers

Viewed asthma as acute, not chronic

Rejected identity of asthmatic, e.g. said had ‘bad chest’

Downplayed severity – ‘only asthma’/slight asthma

Denied, doubted they had asthma

Did not take preventers because: entailed acceptance of chronicity and asthma identity, fears regarding steroids, side effects, dependency

Used reliever medicine, but more than optimum dose, because: did not fear dependency as only taken for symptom relief, seen to control condition

Concealed use of inhalers because seen as stigmatising and obstacle to normalisation

Operated policy of partial disclosure re their condition, preferring to present selves as normal

Asthmatics viewed as weak and wimpy

Have mechanical model of asthma as blocked tubes

But other factors affect adherence

Things that help people take drugs as prescribed, i.e. that discourage ‘incidental non-adherence’:

- Relatives and friends reminding person to take medicines
- Establishing routines for taking medicines

Children and asthma medication

Older children described using brand names, younger using colours, shapes, general terms like pill, puffer

The older children got, the more autonomous they became in controlling own medicines

Children told caretakers when sick, but latter decided whether or not to seek medical help

FIGURE 11 Asthma medication synthesis.
Factors influencing whether or not people take medicine as prescribed

- Cost versus benefit calculation:
  - Actual and hoped for benefits of medicine: improved QoL, reduced symptoms, ability to deal with symptoms, reduced risk of relapse/hospitalisation, normality/sameness\(^{104,106-108}\)
  - Actual and feared physical and psychological adverse effects of medicine\(^{154,156-158}\)
  - Fear of dependency and addiction\(^{154,157}\)
- Lay evaluation to determine difference between effects of illness and effects of medication\(^{158}\)
- Availability of alternatives\(^{103}\)
- Acceptance of illness (schizophrenia)\(^{156,157}\)
- Imposed compliance\(^{158}\):
  - People on neuroleptics more likely to tdap due to threat of social disapproval, hospitalisation\(^{156}\)
  - Perception of social contract: tdap if to be tolerated by community\(^{156}\)
  - Some had experienced coercion from CPNs when had ntdap\(^{156}\)
  - Some had experienced surveillance and coercion from significant others\(^{156,158}\)
  - Some had drugs injected into them to improve compliance\(^{153}\)
- Specific attributes of medication (psychotropic):
  - Taking psychotropic medicine is stigmatising and can lead to social alienation and discrimination\(^{156,158}\)
  - Adverse effects of psychotropic medicine can mark out as different, be stigmatising\(^{158}\)
  - Neuroleptic medicine implies you are a lesser person and not whole without it\(^{158}\)
  - Experienced by some as form of external control\(^{158}\)
- Specific attributes of medication (benzodiazepines)
  - Withdrawal is difficult\(^{154,155}\)
  - Attitudes: shameful, embarrassing, evil, stigmatising, addictive, immoral, powerful, destructive, okay because doctor legitimises, good because relieves symptoms/improves QoL, ambivalence\(^{153,154}\)
  - Relationships: acquaintance, workmate, old friend, keeper, master – scale of dependency on benzodiazepines, feeling of controlling/being controlled by benzodiazepine use\(^{154}\)

Purposeful non-adherence: ways of ntdap

- Taking drugs symptomatically rather than curatively/preventively\(^{153}\), e.g. clozapine\(^{157}\)
- Self-regulation: taking more/less than prescribed or stopping completely to gain sense of control (easier for outpatients and with PRN)\(^{153,156}\)
- When competing priorities, i.e. to avoid side effect if going out drinking\(^{156}\)

Doctor–patient relationship (benzodiazepines)

- Some doctors discouraged patients’ use of benzodiazepines\(^{154,155}\)
- Some doctors continued unseen repeat prescribing which interpreted by patients as legitimising benzodiazepine use\(^{154,155}\)
- Some doctors and patients enjoyed mutual trust\(^{154}\)

Doctor–patient relationship (other psychotropics)

- Patients on neuroleptics who were ntdap did not let health professionals know as climate of coercion/enforced compliance with psychotropic drugs, perception (correct) that others have power over their lives\(^{156}\)
- Patients’ use of alternatives to drugs (e.g. prayer) can lead health workers to label them as ‘uncompliant’\(^{153}\)
- Patients’ attempts at regulating medication could lead to either conflict or negotiation with health professionals\(^{153}\)

But other factors affect adherence

- Incidental non-adherence
  - Running out of tablets\(^{153}\)
  - Missing appointments\(^{153}\)

FIGURE 12 Psychotropic medication synthesis. CPN, community psychiatric nurse; ntdap, not taking drugs as prescribed; PRN, prescribed as needed; QoL, quality of life.
As an abstract representation of the findings and the relationships between them, it lacks necessary detail.

**Synthesising translations across the illness groups**

The next stage was to return to the detail, to the textual translations that were conducted for each of the medicine groups, and synthesise these across the medicine groups. So whereas the medicine maps were ‘brought together’ to provide our overall model of medicine-taking, now the textual syntheses for each of the medicine groups were ‘brought together’, or synthesised, to provide an overall synthesis of medicine-taking. This initially involved reading and rereading the textual syntheses for each of the medicine groups (referring back to the original papers where clarification was necessary), then bringing together the data thematically. So, for example, if ‘worries about medicine-taking’ were identified in the asthma, HIV infection and hypertension textual syntheses, these data were brought together beneath this heading in the overall synthesis. Thus, the data for the overall synthesis were analysed thematically rather than according to medicine group. This overall synthesis is a ‘lines-of-argument’ synthesis, in that it says something about the whole based on studies of the parts.

Noblit and Hare (p. 64) explained the relationship between the translating of cases into each other and the subsequent expression of the synthesis as ‘lines-of-argument’: ‘In short, the translation of cases into one another sets the stage for a second-level inference about the relationship between the studies. It is the second-level inference that assigns interpretive significance to each study to be synthesized’. The ‘lines-of-argument’ synthesis, therefore, essentially involves a process of interpretation and conceptual advancement and is at the same level as Britten et al.’s ‘third-order interpretations’. The entire process of the medicine-taking synthesis is summarised in Box 7.
Exclusions during synthesis

At the end of the synthesis, it became apparent that one of the papers in the medicines synthesis had contributed nothing at all to the final synthesis even though it had been included in the asthma medications map. The paper had been borderline because much of it was about children’s response to asthma as well as asthma medicine. On further reading, it became clear that its emphasis precluded it from contributing to the synthesis. The final number of papers that contributed to the medicines synthesis was therefore 37 (Box 8).

Rheumatoid arthritis synthesis

The papers were read and synthesised chronologically, according to publication date. The earliest paper in the series was used as an index paper, i.e. subsequent papers were compared with this one. The synthesis proceeded as ‘reciprocal translations of each case (i.e. paper) into each of the other cases’. There was a conceptual coherence among the majority of the studies that eased the synthesis process. The synthesis was conducted in a cyclical fashion. The papers were read carefully three times by the first author: once for appraisal and data extraction; once during the reciprocal translation phase; and once to fill in any holes in the emerging theory. As noted above, the papers were also read once independently by second assessors. Thus, each paper was read in detail on four different occasions by two different people.

Reciprocal translation involves comparing the findings and concepts from each included paper with those of the others. The coherence of the papers in the RA synthesis meant that it was possible to do this simply by means of a table, with columns representing all of the included studies, and rows representing differently themed groups of findings or concepts. The whole table is too large to be included here, although an extract of some of the studies and theme groups has been given (Table 16) by way of example. Some theme groups were renamed, added to, merged or deleted during the reciprocal translation process. In order to preserve the authors’
### TABLE 16 Reciprocal translations of four classic qualitative studies of the lived experience of RA

<table>
<thead>
<tr>
<th>Disease conditions</th>
<th>Bury, 1982</th>
<th>Stenström et al., 1993</th>
<th>Shaul, 1995</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signs and symptoms (p. 170)</td>
<td>‘The respondents were uncertain about themselves and their own interpretations of their symptoms, and whether they would be able to get adequate help to master the disease and manage their everyday lives’ (p. 237)</td>
<td>‘Most common symptoms involved the hands, wrists, shoulders, and feet. In some cases, pain and swelling occurred simultaneously, in others, pain occurred alone, followed by swelling. In either case, there was usually weakness in the affected limb and general fatigue’ (p. 292)</td>
<td>‘Fatigue and pain were cited as the most distressing of the symptoms and contributory to an increase in depressive symptoms’ (p. 293)</td>
</tr>
<tr>
<td>‘Implies premature ageing (p. 171, after Singer, 1974)’</td>
<td></td>
<td>‘Weakness (p. 293)’</td>
<td>‘Of all the physical symptoms that the women described, none were as pervasive as fatigue. Fatigue was often the first sign of an impending flare, and continued long after the flare had subsided. Some women commented on being “always tired”’ (p. 294)</td>
</tr>
<tr>
<td>‘An outside force … [invading] … all aspects of life’ (p. 173)</td>
<td></td>
<td></td>
<td>‘Of all the physical symptoms that the women described, none were as pervasive as fatigue. Fatigue was often the first sign of an impending flare, and continued long after the flare had subsided. Some women commented on being “always tired”’ (p. 294)</td>
</tr>
<tr>
<td>‘A crippling disease’ (p. 173)</td>
<td></td>
<td></td>
<td>‘Of all the physical symptoms that the women described, none were as pervasive as fatigue. Fatigue was often the first sign of an impending flare, and continued long after the flare had subsided. Some women commented on being “always tired”’ (p. 294)</td>
</tr>
<tr>
<td>‘Nature of the disease remains elusive’ (p. 173)</td>
<td></td>
<td></td>
<td>‘Of all the physical symptoms that the women described, none were as pervasive as fatigue. Fatigue was often the first sign of an impending flare, and continued long after the flare had subsided. Some women commented on being “always tired”’ (p. 294)</td>
</tr>
<tr>
<td>Burdens or needs (added while synthesising Stenström et al.)</td>
<td>‘Many saw a future of growing dependency and invalidity’ (p. 173)</td>
<td>Needs accentuated after disease onset:</td>
<td>‘Denial (p. 292)’</td>
</tr>
<tr>
<td>‘Functional limitations’ (p. 175)</td>
<td></td>
<td>‘Maintenance of one’s identity, personal integrity, mental balance, practical help, support from significant others, social support, communication about the disease, social contacts’ (p. 237)</td>
<td>‘Learning to live with it’ (p. 293)</td>
</tr>
<tr>
<td>‘The erstwhile taken-for-granted world of everyday life becomes a burden of conscious and deliberate action’ (p. 176)</td>
<td></td>
<td></td>
<td>‘They incorporated the illness into their self-image, and although they could not predict the course of the illness or when another flare would occur, they were better prepared to cope with it when it came’ (p. 293)</td>
</tr>
<tr>
<td>Chronic illness as a disruptive experience (biography, relationships)</td>
<td></td>
<td></td>
<td>‘The thought of death seemed to bring some form of relief, even as a fantasy’ (p. 294)</td>
</tr>
<tr>
<td>Problems of legitimating changed behaviour; stigma</td>
<td></td>
<td></td>
<td>‘The thought of death seemed to bring some form of relief, even as a fantasy’ (p. 294)</td>
</tr>
<tr>
<td>Psychological strategies</td>
<td>‘The search for a more comprehensive level of explanation, a more certain basis of coping with the illness is often a long and profound one’ (p. 174)</td>
<td>Strategies/attitudes used to reduce uncertainty, regain control and achieve security in everyday life: ignore; mental and physical struggle; solve problems; distract; enjoy; hope; adjust/accept; reappraise; escape/deny/suppress; be resigned (p. 239)</td>
<td>‘Denial (p. 292)’</td>
</tr>
<tr>
<td></td>
<td></td>
<td>‘Sometimes it is possible to confront the disease, whereas at other times the only thing to do is hope for better times’ (p. 240)</td>
<td>‘Learning to live with it’ (p. 293)</td>
</tr>
<tr>
<td>Social strategies</td>
<td>‘The attempt to normalise in the face of disruption’ (p. 177)</td>
<td></td>
<td>‘They incorporated the illness into their self-image, and although they could not predict the course of the illness or when another flare would occur, they were better prepared to cope with it when it came’ (p. 293)</td>
</tr>
<tr>
<td></td>
<td>‘Departing from behaviour which is deemed appropriate carries it’s own specific disadvantages and thus is avoided as far as possible’ (p. 177)</td>
<td></td>
<td>‘The thought of death seemed to bring some form of relief, even as a fantasy’ (p. 294)</td>
</tr>
</tbody>
</table>
own findings and concepts, verbatim quotations from the papers were retained at this stage of the synthesis, and the page numbers in the cells in the table refer to the pages from which the quotations were taken. The concepts and findings in the table derived primarily from those listed by the first and second assessors when the papers were read and appraised. The initial framework extracted information from the studies into the following headings: sample, data collection setting, data collection method, data analysis method, disease conditions, imperatives, burdens, structural factors, psychological ('coping') strategies and social ('coping') strategies. However, it was necessary in some cases to return to the original papers at this stage, both for clarification, and for further elucidation of findings or concepts which achieved added importance only during the reciprocal translation stage itself. The initial framework was adapted to fit the reciprocal translations as the synthesis progressed.

The synthesis was concerned with perceptions of aetiology, experience and treatment. However, the substance of the studies, and their main theoretical significance, concerned the 'experience' of daily life with RA following the changes brought by the condition. Although issues around aetiology and treatment did feature, the papers that focused on these issues felt marginal to the wider collective voice of the papers. In expressing the synthesis here, the three areas of aetiology, experience and treatment are separated out, although far more attention is devoted to 'experience' as it was the focus of most of the studies. This process resulted in two sociological models, one concerned with gender relations and one concerned with adaptation to RA, which were constructed after the reciprocal translations, perhaps in a manner akin to moving to a 'higher' level of conceptual development (although not in the 'by theme' manner exemplified by Britten et al.18).
**Exclusions during synthesis**

Four papers were excluded at the synthesis stage, leaving a total of 25 papers in the RA synthesis, describing 22 studies (Box 9). One was excluded because the focus was doctor–patient communication, rather than the lived experience of RA. Another was excluded because it focused on conditions other than RA, whereas the remaining two were marginal in terms of whether or not the research methods employed meant that the study fell within our definition of qualitative research.

**BOX 9 Exclusions during synthesis (RA)**

| Number of papers initially included in synthesis | 29 |
| Number of papers excluded during synthesis | 4 |
| Number of papers in final synthesis | 25 (describing 22 studies) |

**Discussion**

**Preliminary studies of reproducibility**

Reproducibility of qualitative research is difficult to assess as it is not possible to collapse the detail and richness of the data into simple quantified components. Assessing the reproducibility of qualitative research is, then, a matter of judgement. There is considerable debate in the methodological literature about whether or not ‘reproducibility’ is a reasonable concept with which to assess qualitative research. It is always likely that different researchers with different backgrounds and different levels of experience will interpret qualitative data in different ways. It is likely, therefore, that the reproducibility of syntheses of qualitative research is also likely to be variable and related to the particular researchers involved. This can be seen in the comparison above, particularly in relation to researchers’ different levels of experience of synthesising: PP had undertaken this before and so did not feel the need to describe the process in as much detail as GDW who was doing this for the first time. Similarly, as the two researchers were going on to complete only one of the syntheses each, there were differences in their preparation of the material for continuing to synthesise beyond the first four papers.

Many of the concepts and findings in these two syntheses were nevertheless combined similarly by the researchers. There were differences of detail, but often these were related to the way in which the original authors presented the concepts, which presented the opportunity for terminology to be interpreted differently. The approaches to the syntheses were somewhat different. GDW, for example, preferred to place the findings in tables to facilitate comparison, whereas PP preferred to describe the process of combining findings more textually. Such differences are reflective of individuals’ particular modes of working, and it is unlikely that a single approach could be deduced and applied universally. Indeed, this is also true of primary qualitative research. The key to synthesising qualitative research lies in becoming immersed in the findings and concepts presented by the original authors, and then attempting to combine these to reach summaries of findings and higher level concepts.

Overall, this preliminary and exploratory attempt to investigate the reproducibility of the first stage of a qualitative synthesis has shown that the combination of concepts and findings is fairly similar, even when the approach to synthesising is quite different. It is possible to deduce reasons for differences related to approach and background of the researchers and a lack of explicitness on the part of the original authors. This investigation involved experienced qualitative
researchers. The complexity of the task suggests to us that it should always be undertaken by experienced researchers. It is our view that less experienced researchers would find the tasks of translating concepts more difficult, with the likely result of less coherent syntheses – although, this remains to be tested. This investigation examined reproducibility only at an early stage with only four papers, and it is not clear whether the syntheses would have become more or less similar with the inclusion of further papers. Only two researchers independently synthesising the same full set of papers would show this. We would suggest that the evidence from this investigation of reproducibility should assure researchers and policy-makers that the process of qualitative synthesis is reasonably robust when undertaken by experienced researchers.

**Full syntheses**

As with the preliminary syntheses, the two full syntheses were conducted differently from each other, primarily reflecting the differing amounts of literature to be synthesised. Because there was more literature for the medicines topic, additional work had to be done to organise the literature and make the synthesis process manageable. Another research group might simply have synthesised the medicine papers chronologically, but it is likely that an analysis by medicine or disease would still have been conducted at some later stage. More detailed reflections on the full syntheses will be offered at the conclusion of the synthesis findings (see Chapter 6).
Chapter 6
Resisting medicines: a synthesis of qualitative studies of medicine-taking

Introduction

The year 2003 saw the publication of three high-profile reports on medicine-taking: one from the World Health Organization (WHO), one from the government-funded Medicines Partnership and the last from the King’s Fund. Meanwhile, an initiative called ‘Ask about Medicines Week’ (www.askaboutmedicines.org) was launched to ‘help promote partnership in medicine-taking between medicine users, carers and health professionals’. During ‘Ask About Medicines Week’, the British Medical Journal dedicated an issue to the theme of ‘concordance’, as did the official journal of the Royal Pharmaceutical Society, The Pharmaceutical Journal. Additionally, the NHS Service Delivery and Organisation Research and Development programme commissioned a scoping exercise to review ‘Concordance, adherence and concordance in medicine taking’ which reported in 2005. What prompted this attention?

The WHO report aimed to improve worldwide rates of ‘adherence’ to long-term treatments for chronic conditions, which the authors of the report currently estimated to be 50%. They argued that the consequences of low adherence to long-term treatments were firstly poor health and secondly increased costs of health care, claiming: ‘Poor adherence is the primary reason for sub-optimal clinical benefit. It causes medical and psychosocial complications of disease, reduces patients’ quality of life and wastes health care resources’ (p. 25). Likewise, the Medicines Partnership cited ‘a rising drugs bill and the key role of medicines in promoting health’ as the key reasons for improving compliance. The editor of The Pharmaceutical Journal explained that the themed issue aimed to clarify the concept of concordance because, if concordance can be achieved, ‘… people are more likely to take the medicines prescribed for them and stick to the regimen’ (p. 480). Although noting that the result may also be a refusal of the medicine, the editor went on to remind readers that many medicines are taken by less than half the patients to whom they are prescribed and that ‘… the waste alone should make them think again’ (p. 480).

The reports by the WHO and by the Medicines Partnership were wide-ranging and reviewed the literature to establish what was known about medicine-taking, the factors affecting it and interventions that could enhance it. The Medicines Partnership report noted that several factors can affect medicine-taking, including demographic factors, issues surrounding the medicine itself (including adverse effects and the complexity of the regimen), ‘beliefs’ about medicines, concerns about the value or appropriateness of medicines, psychosocial issues, confusion and physical difficulties. The WHO report suggested that the main influences on medicine-taking were a wide range of social and economic factors (including poverty, lack of education and ‘culture’), health-care team and system-related factors (such as the patient–provider relationship and the quality of the health services offered), condition-related factors (for example, the severity of the illness and its symptoms), therapy-related factors (i.e. the efficacy of the treatment itself and whether or not it has adverse effects) and, finally, patient-related factors (such as anxieties about the treatment, stress, perceptions that the medicine is not needed or not effective, and non-acceptance of illness).
The independent King’s Fund report was quite different.\textsuperscript{185} It was written in response to a ‘growing recognition’ that the interests of those who provide health care do not necessarily coincide with the needs of those who use it. Harrison\textsuperscript{185} suggested that at present an implicit public–private partnership exists between the NHS and the pharmaceutical industry, but that the pharmaceutical industry is largely free to provide products it considers will be profitable, whereas the NHS is a passive purchaser. Harrison\textsuperscript{185} argued that this has led to an emphasis on pharmaceutical products at the expense of non-pharmaceutical approaches to health care such as behavioural therapies, illness prevention, public-health interventions and ‘alternative therapies’. Furthermore, because research priorities are not determined by consumers, the needs of major groups, including women, children and older people, have been ignored, and important therapeutic areas have been neglected. “These circumstances have produced health care systems that are highly drug dependent and that in general cannot see themselves being any different.”\textsuperscript{185}

The reports by the WHO\textsuperscript{183} and the Medicines Partnership are, arguably, products of this type of health-care system and culture, where health is seen to depend upon pharmaceutical medicine. Both reports assumed that adherence to medicine will lead to better health. Yet, a systematic review of interventions to increase adherence to medicines found that successful interventions (of which there were 19 out of 39) did not lead to large improvements in adherence or treatment outcomes.\textsuperscript{189}

The bulk of the research upon which both reports relied is quantitative and much of it has been conducted from the professional or systems perspective, taking little account of lay perspectives. Vermeire \textit{et al.}\textsuperscript{190} observed that during three decades of quantitative research into non-compliance, > 200 variables have been studied, but none consistently predicts compliance. The authors suggested that despite continuing research, there have been few insights since the 1980s. Most of the research has been on the extent and determinants of non-compliance, but Vermeire \textit{et al.}\textsuperscript{190} argued that this research has been fragmented, of variable methodological quality and lacking an integrating model or theory. The authors attributed the lack of success in this field to neglect of patients’ perspectives and beliefs, as well as to an absence of qualitative research.

In 1996, Blaxter and Britten\textsuperscript{191} reviewed lay beliefs in the context of medicine-taking and concluded, usefully, that lay people do not view medicine as something to be taken ‘as prescribed’, but rather as a resource for use as they themselves see fit. The theory that medicines are used as a lay resource provides an alternative perspective on the behaviour that professionals find bewildering. If lay people use medicines as a resource then it is understandable that they might modify their regimens to suit their daily routines, take medicine only when they experience symptoms, cease taking it if they experience adverse effects or draw upon other resources, such as ‘natural remedies’ alongside it. Blaxter and Britten\textsuperscript{191} suggested that more research, especially qualitative research, is needed to understand lay theories of bodily systems, the actions of medicines, the origins of lay beliefs about medicines and the influence of the media on public perceptions of medicines.

Our synthesis brings together 10 years of qualitative research into lay experiences of medicine-taking and we hope that the emphasis on lay perspectives will shed new light on this field. Before outlining the various perspectives of the studies, however, a brief discussion of the terminology of medicine-taking is necessary because, as can be seen, this varies. Medicine-taking has been characterised in three different ways during the last few decades: compliance, adherence and now concordance. Compliance is traditionally assumed to refer to doctors’ desire for patients to comply with their instructions about taking medicine. Adherence was subsequently offered as a more neutral expression, in the hope of avoiding the paternalism for which compliance came to be criticised. The tone remained prescriptive, however, and a new model, that of concordance, was proposed.\textsuperscript{195} Concordance refers to the anticipated outcome of the consultation between
doctors and patients about medicine-taking, if both parties can be encouraged to work together as partners. In practice, the three terms are often used interchangeably, despite the subtle differences in meaning and perspective.

The studies

Key features of the studies in this synthesis are summarised in Table 17 and the citation tracking of papers is recoded in Table 18. As can be seen, most studies are concerned with medicines taken on a long-term basis for chronic illnesses.

**TABLE 17 Features of the 37 studies synthesised**

<table>
<thead>
<tr>
<th>Source paper (n=38)</th>
<th>Country setting</th>
<th>Participants (female)</th>
<th>Sample origins</th>
<th>Medicines/condition</th>
<th>Data collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Donovan and Blake, 1992&lt;sup&gt;168&lt;/sup&gt;</td>
<td>UK</td>
<td>54 (39) people</td>
<td>Hospitals</td>
<td>Miscellaneous (inflammatory arthropathy/osteoarthritis/minor rheumatology complaints)</td>
<td>Semi-structured interviews, before and after recorded consultation with doctor</td>
</tr>
<tr>
<td>Kaljee and Beardsley, 1992&lt;sup&gt;153&lt;/sup&gt;</td>
<td>USA</td>
<td>70 people</td>
<td>Rural mental-health clinic</td>
<td>Psychotropic medicine</td>
<td>Observation of meetings, group sessions, in-depth interviews</td>
</tr>
<tr>
<td>Roberson, 1992&lt;sup&gt;173&lt;/sup&gt;</td>
<td>USA</td>
<td>23 (14) black adults aged 19–77 years, varied education; 18 hypertension, 8 arthritis, 5 diabetes, 1 asthma, 1 hepatitis, 1 erythema multiforme</td>
<td>Southern rural African American community population</td>
<td>Medicines in general</td>
<td>Unstructured interviews</td>
</tr>
<tr>
<td>North et al., 1995&lt;sup&gt;154&lt;/sup&gt;</td>
<td>New Zealand</td>
<td>22 Europeans (11) aged from 34 to 82 years</td>
<td>One group from the community and one from self-help group</td>
<td>Psychotropic medicine</td>
<td>In-depth interviews</td>
</tr>
<tr>
<td>Barter and Cormack, 1996&lt;sup&gt;155&lt;/sup&gt;</td>
<td>UK</td>
<td>11 (10) aged 60–90 years. Also sample of 20 (12) elderly people ‘randomly’ sampled on street, not on benzodiazepines</td>
<td>Community (those in receipt of benzodiazepines for at least 1 year)</td>
<td>Psychotropic medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Britten et al., 1996&lt;sup&gt;156&lt;/sup&gt;</td>
<td>UK</td>
<td>30 (11) aged 20–70 years, 27 white, 2 black and 1 Middle Eastern</td>
<td>Two London GP practices: poor area (9 patients), affluent (21)</td>
<td>Medicines in general</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Dowell et al., 1996&lt;sup&gt;157&lt;/sup&gt;</td>
<td>UK</td>
<td>17 people</td>
<td>Urban Scottish general practice</td>
<td>Miscellaneous (rapid prescribing changes)</td>
<td>In-depth interviews, twice over 6 months</td>
</tr>
<tr>
<td>Morgan, 1996&lt;sup&gt;158&lt;/sup&gt;</td>
<td>UK</td>
<td>60 (30): 30 white, 30 African Caribbean (30) all aged at least 35 years</td>
<td>15 general practices in London Borough of Lambeth</td>
<td>Antihypertensive medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Adams et al., 1997&lt;sup&gt;159&lt;/sup&gt;</td>
<td>UK</td>
<td>30: 14 women (10 working class, 4 middle class) and 16 men (8 working class and 8 middle class). Aged 19–57 years</td>
<td>Single general practice in south Wales (former mining) town</td>
<td>Asthma medicine</td>
<td>In-depth interviews</td>
</tr>
<tr>
<td>Boath and Blenkinsopp, 1997&lt;sup&gt;160&lt;/sup&gt;</td>
<td>UK</td>
<td>20 (11) aged 28–68 years, 15 married, 10 working, 1 unemployed, 1 housewife, 4 unable to work and 4 retired</td>
<td>One fund-holding group medical practice</td>
<td>PPIs</td>
<td>Semi-structured interviews</td>
</tr>
</tbody>
</table>

**continued**
<table>
<thead>
<tr>
<th>Source paper</th>
<th>Country setting</th>
<th>Participants (female)</th>
<th>Sample origins</th>
<th>Medicines/condition</th>
<th>Data collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dowell and Hudson, 1997</td>
<td>UK</td>
<td>44 (24), 23 aged &gt;65 years, 12 living alone</td>
<td>Two samples, both from GP registers</td>
<td>Medicines in general</td>
<td>In-depth interviews</td>
</tr>
<tr>
<td>Siegel and Gorey, 1997</td>
<td>USA</td>
<td>71 women, 42% black, 17% white, 40% Puerto Rican. Mean age 34.9 years. 37% working, 82% were parents. Median household income low (US$12,500 per annum)</td>
<td>HIV organisations, including hospitals and community organisations</td>
<td>Antiretroviral medicine</td>
<td>Unstructured interviews</td>
</tr>
<tr>
<td>Rogers et al., 1998</td>
<td>UK</td>
<td>34 (12) aged 18–56 years, varied social class</td>
<td>Voluntary groups, MIND centres/outpatient centres, inpatients</td>
<td>Psychotropic medicine</td>
<td>In-depth interviews</td>
</tr>
<tr>
<td>Stone and Clarke, 1998</td>
<td>USA</td>
<td>56 (28), 50% white, 29% black, 21% Latino, mean age 37 years, varied education</td>
<td>Five hospital and community centres in Boston, MA, USA and Providence, RI, USA</td>
<td>Antiretroviral medicine</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Van Wissen et al., 1998</td>
<td>New Zealand</td>
<td>19 (15), 2 Maori, 17 European. Age range 41–67 years, mean age 54 years</td>
<td>Register of people previously involved in hypertension research</td>
<td>Antihypertensive medicine</td>
<td>In-depth interviews</td>
</tr>
<tr>
<td>Watson and Mitchell, 1998</td>
<td>Australia</td>
<td>37 women aged 59–83 years living independently in the community</td>
<td>Two community pharmacies</td>
<td>Medicines in general</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Erlen and Mellors, 1999</td>
<td>USA</td>
<td>Six (two), five ‘on disability’, one employed, three white, three black, all high school educated</td>
<td>University trial unit/ community AIDS ministry/private practice</td>
<td>Antiretroviral medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Ersek et al., 1999</td>
<td>USA</td>
<td>21 (18), mean age 60 (range 42–79) years, mostly white, educated, married</td>
<td>Larger longitudinal study (not known where patients from)</td>
<td>Miscellaneous (cancer)</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Halkitis and Kirton, 1999</td>
<td>USA</td>
<td>37, 22% female, mean age 42 years. Men: 24% heterosexual, 76% bisexual or gay. Women: majority heterosexual, 38% black, 24% Latino, 35% white</td>
<td>Attendants of large city AIDS service organisation</td>
<td>Antiretroviral medicine</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Johnson et al., 1999</td>
<td>USA</td>
<td>21 (17), aged 65–92 years, Caucasian, retired</td>
<td>Local emergency centres and physicians’ offices</td>
<td>Antihypertensive medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Proctor et al., 1999</td>
<td>USA</td>
<td>39: 27 men age 30–69 years, 12 women age 29–60 years. 9 white, 16 black, 4 Hispanic. 10 gay men, 9 heterosexual men, 9 heterosexual women, 11 injecting drug users</td>
<td>Five New York institutions providing AIDS services: three state, one federal and one private</td>
<td>Antiretroviral medicine</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Prout et al., 1999</td>
<td>UK</td>
<td>Nine families: five middle class, four working class, including four girls, five boys aged 10–12 years</td>
<td>Larger European Union Biomed project. Asthma clinics in two general practices in towns in Midlands</td>
<td>Asthma medicine</td>
<td>Repeated interviews, some open-ended some using check list</td>
</tr>
<tr>
<td>Siegel et al., 1999</td>
<td>USA</td>
<td>78 (20), age 50–68 years, 41% black, 19% Puerto Ricans, 40% non-Hispanic white. Majority single, most living alone. 51% heterosexual, 42% gay. Varied education, low income (only 18% on &gt; US$20,000 per annum)</td>
<td>Community-based health and social organisations, support groups, advocacy groups and drug treatment centres in New York, NY</td>
<td>Antiretroviral medicine</td>
<td>Questionnaires and semi-structured interviews</td>
</tr>
<tr>
<td>Source paper</td>
<td>Country setting</td>
<td>Participants (female)</td>
<td>Sample origins</td>
<td>Medicines/condition</td>
<td>Data collection</td>
</tr>
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</tr>
<tr>
<td>Atkin and Ahmad, 2000[@]</td>
<td>UK</td>
<td>25 children: 12 boys, 13 girls, average age 13.9 years. 22 Pakistani Muslim, 1 Bangladeshi Muslim, 1 Iranian Muslim, 1 Indian Hindu. 15 school, 7 college, 2 unemployed, 1 university</td>
<td>Health professionals’ records, e.g. paediatricians/specialist haemoglobinopathy workers in six localities in Midlands/northern England</td>
<td>Miscellaneous (thalassaemia major)</td>
<td>In-depth interviews, twice over 6 months</td>
</tr>
<tr>
<td>Barton Laws and Wilson, 2000[@]</td>
<td>USA</td>
<td>25 (8) aged 27–57 years, 9 white, 3 black, 12 Latino, 1 Portuguese</td>
<td>AIDS groups in Boston and other parts of Massachussets, word of mouth</td>
<td>Antiretroviral medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Buston and Wood, 2000[@]</td>
<td>UK</td>
<td>49 (29), mean age 15.6 years. Mean age of diagnosis 4.9 years. 35 at school, 7 in further education, 5 unemployed and 2 employed</td>
<td>Hospital asthma clinics in Greater Glasgow</td>
<td>Asthma medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Lumme-Sandt et al., 2000[@]</td>
<td>Finland</td>
<td>No information, except originally 448 people aged &gt; 90 years. 250 interviewed. Data from 151 interviews referring to medication.</td>
<td>All people &gt; 90 years living in the city of Tampere in southern Finland</td>
<td>Medicines in general</td>
<td>Narrative/biographical interviews</td>
</tr>
<tr>
<td>McDonald et al., 2000[@]</td>
<td>Australia</td>
<td>76 (13) aged 25–62 years. 52 gay men, 2 women and 6 men bisexual, 11 women and 5 men heterosexual. 84% of interviewees currently using antiretroviral medicine</td>
<td>Larger HIV Futures Study of people with AIDS/HIV in Australia. AIDS organisations/ mailing lists, hospitals, doctors’ surgeries, adverts</td>
<td>Antiretroviral medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Murphy et al., 2000[@]</td>
<td>USA</td>
<td>39, aged 33–54 years, 69% male. 44% black, 39% white, 6% other/mixed race, 6% Latino (16% graduate college degrees, 24% undergraduate college degrees, 32% some college, 19% high school, 8% less than a high school education)</td>
<td>HIV infection clinic and advertisement in an AIDS publication</td>
<td>Antiretroviral medicine</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Johnston Roberts and Mann, 2000[@]</td>
<td>USA</td>
<td>20 women aged 25–54 years. 50% Hispanic, 35% black, 15% white. Varied education. Two-thirds with more than one child</td>
<td>Los Angeles, CA, USA, HIV infection/AIDS clinic</td>
<td>Antiretroviral medicine</td>
<td>Participants kept a journal</td>
</tr>
<tr>
<td>Siegal et al., 2000[@]</td>
<td>USA</td>
<td>49 (9) HIV-positive adults aged 50–67 years, in New York, NY metropolitan area. 45% black, 51% white, 4% Latino, 51% heterosexual, 18% bisexual, 31% gay/lesbian. Education varied, 86% unemployed. 63% live alone</td>
<td>Community-based health and social organisations, HIV infection support groups and HIV infection advocacy organisations in New York, NY</td>
<td>Antiretroviral medicine</td>
<td>Semi-structured interviews</td>
</tr>
<tr>
<td>Smith et al., 2000[@]</td>
<td>UK</td>
<td>No information on participants</td>
<td>Relevant national and local voluntary organisations</td>
<td>Miscellaneous (arthritis/respiratory disease/mental health)</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Svensson et al., 2008[@]</td>
<td>Sweden</td>
<td>33 (15), mean age 58 (range 35–83) years, mean duration of hypertension 10 (range 1–30) years</td>
<td>Rural GP centre and specialist hypertension unit in major hospital, both in southern Sweden</td>
<td>Antihypertensive medicine</td>
<td>Semi-structured interviews</td>
</tr>
</tbody>
</table>
Sociological and anthropological studies challenging compliance

The earlier sociological and anthropological studies in this synthesis challenged decades of quantitative research that had worked within the paradigm of compliance. The first three studies noted that previous research rested on an ideology of compliance which implied medical dominance. Donovan and Blake, in the context of a study into lay experiences of treatment for RA, argued that the concept of ‘compliance’ lays the responsibility squarely on the patient, and questioned the assumption that patients should comply with their medical treatment, observing that ‘non-compliance’ is a problem for doctors, not patients. Roberson, whose study is about general experiences of taking medicine, also highlighted the assumptions upon which earlier research was based: that people’s health will improve if they follow their doctors’ orders, that biomedicine is the only option and that those who do not follow medical advice are irrational, deviant or uncooperative. Kaljee and Beardsley, writing about psychotropic medicines, noted that the term ‘compliance’ assumes the authority of doctors over patients, the latter being represented as ignorant and unable to understand the doctors’ orders. Like Roberson, Kaljee and Beardsley noted that the biomedical approach is only one among many other ways of achieving health (a point that is seldom made after these two early papers). Both Roberson and Kaljee and Beardsley concluded that if health professionals want to understand compliance it is important to understand the patients’ perspective, and Donovan and Blake recommended the ‘development of active, co-operative relationships between patients and doctors’ (p. 512).

Later studies by sociologists Adams et al. of treatment for asthma, Rogers et al. of neuroleptic medicine, Lumme-Sandt et al. of older people’s experiences of medicine-taking and Atkin and Ahmad of the ways in which young people respond to chelation therapy for thalassaemia major also gave primacy to lay perspectives and tried to illustrate that patients’ behaviour could be understood as rational and strategic, rather than passive and powerless. Like the earlier studies, they suggested that qualitative investigations of lay explanations and experiences would provide clues as to how people can be better supported in taking their medicine. Thus, although they challenged the ideology of compliance for its paternalism, these studies nevertheless displayed varying levels of attachment to the biomedical paradigm.
### TABLE 18 Citation tracking in the medicines synthesis

<table>
<thead>
<tr>
<th>Medicine</th>
<th>Author citing</th>
<th>Date</th>
<th>Author cited</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIV infection</td>
<td>Siegel and Gorey¹⁴²</td>
<td>1997</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Stone and Clarke¹⁴³</td>
<td>1998</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Erlen and Mellors¹⁴⁴</td>
<td>1999</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Halkitis and Kirton¹⁴⁵</td>
<td>1999</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Proctor et al.¹⁴⁶</td>
<td>1999</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Siegel et al.¹⁴⁷</td>
<td>1999</td>
<td>Siegel and Gorey, 1997¹⁴²</td>
</tr>
<tr>
<td></td>
<td>Siegel et al.¹⁴⁸</td>
<td>2000</td>
<td>Siegel and Gorey, 1997;¹⁴² Siegel et al., 1999¹⁴⁷</td>
</tr>
<tr>
<td></td>
<td>McDonald et al.¹⁴⁹</td>
<td>2000</td>
<td>None</td>
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<td>Murphy et al.¹⁵⁰</td>
<td>2000</td>
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<td></td>
<td>Johnston Roberts and Mann¹⁵¹</td>
<td>2000</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Barton Laws and Wilson¹⁵²</td>
<td>2000</td>
<td>Stone and Clarke, 1998¹⁴³</td>
</tr>
<tr>
<td>Antihypertensives</td>
<td>Morgan¹³⁴</td>
<td>1996</td>
<td>Donovan and Blake, 1992¹³⁴</td>
</tr>
<tr>
<td></td>
<td>Van Wissen et al.¹¹³⁵</td>
<td>1998</td>
<td>Roberson, 1992¹³⁷</td>
</tr>
<tr>
<td></td>
<td>Johnson et al.¹¹³⁶</td>
<td>1999</td>
<td>Dowell and Hudson, 1997;¹¹³ Van Wissen et al., 1998¹¹³⁵</td>
</tr>
<tr>
<td></td>
<td>Svensson et al.¹¹³⁷</td>
<td>2000</td>
<td>None</td>
</tr>
<tr>
<td>Psychotropic medicines</td>
<td>Kaljee and Beardsley¹¹³</td>
<td>1992</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>North et al.¹¹⁴</td>
<td>1995</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Barter and Cormack¹¹²⁴</td>
<td>1996</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Rogers et al.¹¹²³</td>
<td>1998</td>
<td>None</td>
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<td></td>
<td>Angermeyer et al.¹¹²⁷</td>
<td>2001</td>
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<td>Usher¹¹²⁸</td>
<td>2001</td>
<td>None</td>
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<td>PPIs</td>
<td>Boath and Blenkinsopp¹¹⁵⁹</td>
<td>1997</td>
<td>Dowell et al., 1996¹¹⁵²</td>
</tr>
<tr>
<td></td>
<td>Pollock and Grime¹¹⁶⁰</td>
<td>2000</td>
<td>Boath and Blenkinsopp, 1997;¹¹⁶⁰ Donovan and Blake, 1992¹¹⁶⁰</td>
</tr>
<tr>
<td>Asthma medicine</td>
<td>Adams et al.¹¹³¹</td>
<td>1997</td>
<td>Donovan and Blake, 1992;¹¹³⁴ Morgan, 1996¹¹³⁴</td>
</tr>
<tr>
<td></td>
<td>Prout et al.¹¹⁶²</td>
<td>1999</td>
<td>Adams et al., 1997¹¹²⁵</td>
</tr>
<tr>
<td></td>
<td>Buston and Wood¹¹²³</td>
<td>2000</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>Walsh et al.¹¹⁶⁴</td>
<td>2000</td>
<td>Adams et al., 1997;¹¹³¹ Donovan and Blake, 1992¹¹³¹</td>
</tr>
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<td>Miscellaneous medicines</td>
<td>Donovan and Blake¹¹⁶⁶</td>
<td>1992</td>
<td>None</td>
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<td>Dowell et al.¹¹⁶⁷</td>
<td>1996</td>
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<td>Ersek et al.¹¹⁶⁸</td>
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<td>Atkin and Ahmad¹¹⁶⁹</td>
<td>2000</td>
<td>Prout et al., 1999¹¹²³</td>
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<td>Smith et al.¹¹²⁰</td>
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<td>Medicines in general</td>
<td>Roberson¹¹²¹</td>
<td>1992</td>
<td>None</td>
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<td>Britten¹¹²²</td>
<td>1996</td>
<td>Donovan and Blake, 1992¹¹²²</td>
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<td>Dowell and Hudson¹¹²³</td>
<td>1997</td>
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<td>Lumme-Sandt et al.¹¹²⁵</td>
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<td>Donovan and Blake, 1992;¹¹²² Britten 1996;¹¹²² Adams et al., 1997¹¹²⁵</td>
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Sociological studies not focusing on compliance

A small number of sociological studies investigated medicine-taking, but without a focus on compliance. North et al. developed a typology of relationships to benzodiazepines in order to understand how people use them, whereas Britten took Cornwell's work on 'private' and 'public' accounts as a starting point for her analysis of 'orthodox' and 'unorthodox' lay accounts of medicine-taking. Morgan was primarily concerned with how ethnicity influences the way people manage their antihypertensive medicines. She noted that reducing blood pressure through antihypertensive medicines was regarded as a major preventive strategy for decreasing stroke and heart disease, but she did not ally herself with attempting to resolve what the medical profession regarded as a problem of 'non-compliance'. Rather, she suggested that reliance on medicine as a response to ill health can be regarded as culturally rooted. Prout investigated the role of medicine in the lives of families with asthma and concluded that people much prefer it to the 'non-medical' preventive strategies that are advocated alongside the use of inhalers.

Nursing study not focusing on compliance

Usher, writing in a nursing journal, explored the experience of taking neuroleptic medicine. She considered the issue of compliance, but her focus was on understanding what it is like to take the medicine.

Economically motivated studies

Four studies set out to answer specific and largely economically motivated questions about medicines. All but the first of these were published in the journal Social Science and Medicine. Aiming to reduce benzodiazepine use, Barter and Cormack investigated why people continue to take this medicine despite a reduction in prescribing. Dowell et al. explored patients' experiences of having their prescriptions changed in order to determine whether or not expensive prescriptions can be switched for cheaper ones without upsetting too many people. Boath and Blenkinsopp and Pollock and Grime set out to discover, respectively, an explanation for the dramatic increase in prescribing for PPIs and ways of decreasing the costs of PPI prescribing.

Professional agendas

Watson and Mitchell, writing in a nursing journal, investigated how older women use medicines, claiming a dearth of information on women's health. Their discussion, however, was more suggestive of an interest in expanding the role of community nurses in the care of older women. Similarly, Smith et al., writing in a pharmacy journal, investigated long-term users' experiences of medicine use, but they seemed most interested in whether or not there was an opportunity for pharmacists to extend their information-giving role.

Increasing adherence

Just over half of the studies in the synthesis were primarily concerned with finding out how to increase people's use of medicines. These papers were published mainly in medical or nursing journals and varied in the extent to which they questioned the concept of compliance.

Dowell and Hudson writing in Family Practice about medicine-taking in primary care, noted that compliance is based upon problematic assumptions and acknowledged that many people have good reasons for not taking their medicine as prescribed. Nevertheless, they aimed to increase compliance as long as this is effective, safe and acceptable to patients and they suggested that doctors could help by supervising the changes patients make to their regimens. Ersek and Miller, writing in Cancer Practice, aimed to find ways of overcoming cancer patients' non-adherence to analgesic therapy. They concluded, somewhat unusually, that although patient education may help some, assisting people to use non-pharmacologic pain strategies may be more successful. Angermeyer et al.'s study published in a psychology journal investigated...
people's perceptions of neuroleptic medicine in order to find 'hints' for increasing compliance with treatment.

In addition to Morgan's study, three other studies investigated the experience of taking antihypertensive medicine. The first two were published in nursing journals and the third in a cardiology journal. All encouraged adherence and none questioned the assumption that medicines are the only way of managing high blood pressure and preventing strokes and heart disease. Van Wissen et al. took the view that 'Compliance with treatment is a fundamental prerequisite for therapeutic benefit' (p. 568) whereas Johnson et al. suggested, 'Poor adherence to prescribed hypertensive pharmacotherapy is responsible for unnecessary complications, leading to increased health care spending, premature progression of complications, and early death' (p. 319). Interestingly, Johnson et al. seemed to be attributing the morbidity and mortality to non-adherence rather than to the hypertension itself, illustrating the extent to which health is seen to be allied to taking medicine. Svensson et al., on the other hand, although promoting adherence, suggested that modifying the regimen was 'sound behaviour' and should be encouraged if it represented an attempt to gain control of the condition.

In addition to the studies by Adams et al. and Prout et al. (above), two further studies dealt with taking medicine for asthma. Buston and Wood, writing in *Family Practice*, were concerned with surmounting the barriers to compliance in the case of adolescents with asthma, and Walsh et al. were interested in how the doctor–patient relationship influenced compliance. Although the psychotherapeutic framework they adopted was unique in the synthesis, Walsh et al. shared with Buston and Wood the assumption that compliance is desirable. However, Walsh et al.'s emphasis that patients' behaviour is rational and 'sense making' recalled the earlier sociological studies that stressed the importance of lay meanings.

Eleven studies (10 of which were American) focused on adherence to antiretroviral therapy for HIV infection. Most of the authors of these studies took a fairly strong position, repeatedly citing the benefits of taking the medicine (increased survival, delayed disease progression, improved immune functioning, reduced incidence of opportunistic infections) and the problems associated with not taking it (vertical transmission to babies, viral resistance, limited future treatment options). Siegel and Gorey investigated women's experiences of, and views about, taking azidothymidine (AZT; zidovudine) in the hope of increasing their uptake of the drug. Stone and Clarke recommended simplifying the antiretroviral regimen to improve adherence but suggested that only patients who committed themselves to the regimen should be treated. Erlen and Mellors suggested that health professionals need to know what patients' experiences are so that they can individualise regimens to suit the patient. Both Halkitis and Kirton and Proctor et al. focused on the obstacles that prevent adherence to antiretroviral therapies and the strategies people use to enhance adherence.

Siegel et al. took the position that patients were misinterpreting the symptoms of HIV infection as the adverse effects of antiretroviral therapy, either through ignorance or because they could not accept that the HIV infection was progressing. They recommended that health professionals help people distinguish between symptoms and adverse effects, provide strategies for alleviating symptoms and educate patients about the harmful consequences of non-adherence. Siegel et al. categorised those who did not adhere to antiretroviral therapy as offering 'excuses' or 'justifications' for their behaviour and recommended that misconceptions be debunked and the importance of adherence promoted. They noted that ‘… a set of beliefs may be growing among patients that may support non-adherence’ (p. 39). Similarly, Barton Laws and Wilson explored people's experience of taking antiretroviral therapy and concluded that people come to believe that whatever standard of adherence they can achieve is good enough. Murphy et al. investigated factors that facilitate and prevent adherence, and McDonald et al. investigated the
differences between men’s and women’s experiences of treatment. Noting that women with HIV are less likely than their male counterparts to use antiretroviral therapy, the authors concluded that this is because women are less likely to believe in the treatment’s efficacy. Johnston Roberts and Mann set out to explore the unique barriers to adherence to antiretroviral therapy faced by women. They wrote from the standpoint that adherence should be encouraged, but were sensitive to the social factors that impinge uniquely on women, such as motherhood.

In summary, the earlier studies in the synthesis can be understood as reacting to the concept of compliance that had held sway in the preceding decades. They advocated a qualitative methodology for the purpose of providing an alternative, lay perspective and emphasised the rationality of lay behaviour. A small number of sociological studies investigated medicine-taking in its own right, but the absence of compliance as a starting point was rare in the studies. A few studies answered specific economic questions and a couple seemed to be influenced by specific professional agendas. Among the later studies, few were sociological and more were conducted from the perspective of increasing adherence. Taken as a whole, the studies in this synthesis illustrated that the employment of a qualitative methodology, or a sociological perspective, does not necessarily imply abandonment of the medical agenda. The majority of the studies focused on the reasons why people do not take their medicine as prescribed. Few considered the experience of those who reject medicine outright or those who accept it uncritically. Additionally, the vast majority of studies were concerned with chronic illness. Because the protracted nature of chronic illness presents more challenges to Western medicine than acute illness, this is probably a reflection of medical concerns. The ideology that people should take their medicines as prescribed remained dominant in the studies.

Findings

Our findings fall into four parts. We begin with the ways in which people evaluate their medicines: trying out the medicine and weighing up its costs and benefits, stopping the medicine and observing the effect, observing others and obtaining information. We also detail the lay use of subjective and objective indicators, the way in which gender influences evaluations and the several difficulties lay people face in the process of evaluating their medicines. Secondly, we describe the concerns that cannot be resolved through lay evaluation, including concerns about dependence, tolerance and addiction, the potential harm from using medicine on a long-term basis and the possibility of medicine masking other symptoms. The ways in which medicine and identity are linked is the third aspect. Here we considered non-acceptance of illness and consequently of the medicine, medicine as a reminder of being ill, medicine presenting problems of disclosure and also the stigmatising potential of some medicines. The last section presents the ways in which some authors have categorised medicine-taking before going on to detail the various methods that people use to modify their regimens. Attempts to minimise medicine intake were common, and this was reflected in many of the reasons why people modified their regimens: to decrease unwanted effects, to make the regimen more acceptable and for financial reasons. People were also found to take their medicines symptomatically and, in some cases, strategically. Others replaced or supplemented their regimens with non-pharmacological treatments. Additionally, there were incidental reasons for regimen breaks. Finally, we consider the communication between doctors and patients about regimen modifications.

Lay evaluation of medicines and its difficulties and limits

Trying out the medicine and weighing up the costs and benefits

People weighed up the benefits of taking the medicine against the costs of doing so. The costs included adverse effects and the difficulties faced when trying to incorporate the regimen into daily life. In general, the studies focused less on the benefits of medicines than their costs.
Nevertheless, it was clear that people did place hope in their medicines and that these hopes varied according to the medicine in question. The most common reason for taking medicine was to relieve symptoms, whether physical, as in the case of digestive disorders\(^{159,160}\) and RA,\(^{166}\) or mental.\(^{154-156,158,170}\) However, in the case of neuroleptic medicine, people with schizophrenia hoped to avoid relapse and hospitalisation in addition to obtaining symptom relief.\(^{156}\) With treatment for HIV infection, people hoped to slow down or halt the progression of the disease;\(^{142-144}\) asthma medicine was taken both to relieve physical symptoms\(^{163,164}\) and in the hope of sustaining normality.\(^{161,162}\) Those with hypertension hoped that antihypertensive medicine would prevent or reduce the risk of a future illness such as heart attack or stroke\(^{134}\) and control blood pressure and symptoms.\(^{157}\)

The costs, or undesired outcomes, of taking medicines were reported more frequently. Adverse effects were a significant issue for people taking medicine and a key criterion in the evaluation of treatment. Worries about adverse effects were found in the context of RA,\(^{166}\) cancer,\(^{168}\) asthma,\(^{161}\) digestive disorders,\(^{159,160}\) high blood pressure\(^{136,137}\) and schizophrenia.\(^{156,158}\) In the last case, the considerable physical and psychological effects of neuroleptic medicine were highlighted by participants in both studies: effects that could alter thoughts, inhibit social interaction and identify the person as having schizophrenia. However, it was the studies on HIV infection that stood out because of their participants’ emphasis on the undesirable effects of antiretroviral therapy.

All of the 11\(^{142-152}\) studies on antiretroviral therapy reported people’s experiences of adverse events while having treatment, which included nausea, vomiting, gastrointestinal distress, kidney stones, insomnia, headaches, rashes, dry skin, diarrhoea, dizziness, numbness, feeling generally lousy, a bad taste in mouth, neuropathy, anaemia, breathing difficulties, fatigue, stiffness, mood swings, visual problems, leg pain, hair loss, liver damage, cancers, blackening of fingertips and nails, loss of appetite, general ill health and sweating. These reactions were experienced as unpleasant and challenging in their own right, but because they could be so frequent, severe and unpredictable they were also described as instilling fear and distrust of the medicine. Furthermore, they could have the effect of restricting social activities, affecting friendships, relationships and work, making it difficult to look after families and sometimes necessitating disclosure of the illness.\(^{141,143-145,148-150}\) One man described the effect of his regimen:\(^{146}\)

> The first regimen that I was on was Norvir [ritonavir]. I found out that it was almost next to impossible to stay on it. I literally told my doctor, ‘I’m not taking this stuff any more. I don’t care if I die.’ It made me feel 10 times sicker when I took it than when I didn’t take it.

With such overwhelming undesirable effects, it is not surprising that many people viewed their medicine negatively. The following are excerpts from diaries kept by HIV-positive women in Johnston Roberts and Mann’s study:\(^{151}\)

> … I choke down the Viracept [nelfinavir] and it’s stuck in my throat again, chalky and grainy, so it drops into my stomach … like a ton of bricks … The poison that’s supposed to save my life makes me miserable.

(p. 380)

I actually feel as though I am damaging my body with each pill that I take. I believe these medicines are too toxic and powerful and I do not feel comfortable taking them.

(p. 381)
Many of the women in Siegel and Gorey’s study, especially the African American women, also had very negative attitudes to their treatment (AZT), claiming that at best the drug did more harm than good and at worst it did serious harm. They feared that the drug’s high toxicity was damaging to the internal organs and that the drug could not discriminate between HIV cells and healthy cells.

**Acceptability of regimen**

In addition to the occurrence of adverse effects, people evaluated their regimen in terms of whether it was feasible or acceptable given their daily schedules. All of the studies on HIV infection, except two, described the impact of the regimen in strikingly similar ways. Stone and Clarke’s participants found the regimen highly demanding because of the need to take the protease inhibitors exactly as prescribed. The regimen required people to radically alter their daily routines of sleeping and eating and was experienced as interrupting life’s normal flow. Participants described it as becoming ‘the central organising principle’ and it was experienced as being so overwhelming that they were no longer in control of their lives:

> Everything is amplified. I’m more aware of, ‘Oh I ate that cheeseburger. Now I’ve got to wait another two hours before it digests to take my pills’. I feel very out of control with the regime that I have to follow. My life is focusing around that, rather than the other way round.

McDonald et al.’s sample found that the drug regimen had a long-term impact on social relationships, employment and studying, not just the daily routines of sleeping and eating. The authors found that the regimen had a different impact according to gender: for men the social world was made to fit round their medical regimen, whereas for women the social world took priority and impinged on the regimen. The authors also suggested that men were more concerned with the impact of the regimen on themselves, whereas women were more concerned about its impact on relationships. The women in their study were found to be more resistant to the demands of the regimen than men and less worried about not taking the drugs as prescribed. Johnston Roberts and Mann also reported that their women-only sample found the regimen exceptionally difficult to follow. In addition to reporting that it took away their freedom regarding daily routines and activities, the women found that it competed with other goals such as losing or gaining weight, forming relationships and fulfilling their care-giving roles. As one woman wrote in her study journal:

> I am a single parent of three children. So being a mother and father put into one and living with this illness, well, I’m sure people must have easier lives. I take eight different medicines three times a day. Some on an empty stomach and some with food. Some 8 hours apart. It’s extremely hard to stay focused on my medicine being a full-time mother and, on the other hand, it’s hard to stay a full-time mother while fighting for my life.

The frequency of doses and the sheer number of pills (sometimes between 25 and 35 per day) were also found to be problematic in the case of antiretroviral therapy, as was the nature of the pills themselves – their taste, smell, size and shape.

None of the other regimens seemed to have as great an impact as antiretroviral therapy, although the one study on young people with thalassaemia major suggested that chelation therapy was found to be profoundly disruptive. As Atkin and Ahmad described, an infusion pump delivers a drug that helps excrete the excess iron that accumulates as a result of the frequent blood transfusions that people with thalassaemia major need. The pump has to be used five to seven nights a week, for 8–12 hours each time. Nearly all of the young people in the sample cited the
use of the pump as the most difficult and hated aspect of their illness. Older children especially hated it because it disrupted their social life and marked them out as different from their peers. For this reason, some decided not to pursue the treatment. One boy said, ‘To be honest I couldn’t be arsed [sic] to use it. I just got fed up. I wanted to be like everyone else.’ In such cases, the regimen is assessed as being more problematic than the condition it is meant to treat. Roberson found that some of her participants similarly concluded that their regimens (for a variety of illnesses) were too disruptive.

The process of weighing and balancing

As noted above, when assessing whether or not to persevere with a regimen, people had to weigh up the costs and benefits. Most of the studies that referred to this weighing-up process were those that had outlined severe costs as a result of taking the medicine, i.e. those relating to treatment for HIV infection and mental illnesses. In the case of antiretroviral therapy, the undesirable effects of the treatment could be so bad that some people had to seriously question whether or not it was worth continuing with the medicine. Five of the HIV studies noted this and outlined the stark choices involved. McDonald et al. noted that the overwhelming impact of adverse effects affected people’s evaluations of antiretroviral therapy because it was difficult to appreciate the potential benefits when experiencing such unpleasant symptoms. Some of the women in Siegel and Gorey’s study felt that, despite the possible benefits of AZT, the effects were too debilitating to make it worthwhile; others felt that ‘alternative’ treatments were better for quality of life. In Siegel et al.’s study, some of the participants described the medicine as more threatening to their well-being than the disease for which they took it. Similarly, Proctor et al. found that some of their participants chose quality of life with HIV infection in preference to taking antiretroviral therapy.

People had to balance the costs and benefits in the case of psychotropic medicine too. North et al. reported that some of their participants weighed up the costs of taking benzodiazepines (adverse effects, dependence and social alienation) against the benefits (improved quality of life). Both studies of neuroleptic medicine for schizophrenia found high gains associated with taking the medicine in terms of reduced symptoms, improved ability to deal with symptoms and reduced risk of relapse, but also high costs in terms of physical and psychological adverse effects, the stigma of taking neuroleptic medicine and the consequent discrimination. Similarly, Angermeyer et al.’s participants had to weigh up the benefits of clozapine (improved sleep, feeling calmer, less depressed, less anxious, reduced risk of relapse, perceived protection against mental illness) against its undesirable effects which included feelings of sedation, fatigue and lack of motivation, difficulty getting up in the morning, lethargy, passivity, hypersalivation, constipation, weight gain, sexual ‘dysfunction’, increased perspiration and restlessness.

Although the losses and gains experienced by those taking other medicines may appear less extreme than in the cases of HIV infection and mental illness, the process of weighing up the costs and benefits still occurred. Donovan and Blake found, in the context of RA, that the costs included the unpleasantness or stigma of having to take drugs or wear supports, the need to attend clinics for tests and the risks of adverse effects and dependence upon drugs.

I had Naprosyn [naproxen] and they made [the arthritis] a bit easier, but not much, so I stopped them. To me, the odds, the results from the tablets didn’t balance the risks of taking them. The risk was higher than the result.

The benefits of taking drugs in the case of RA were the immediate relief of symptoms or the anticipation of relief in the long term. Donovan and Blake noted that some people decided whether or not to accept the medicine without even trying it, whereas others made a decision based on the results of tests.
Resisting medicines: a synthesis of qualitative studies of medicine-taking

**Stopping the medicine and observing the effect**

Siegel *et al.*, referring to patients as ‘naive scientists’, suggested that people formulate hypotheses about medicines causing effects and then test these hypotheses by altering the dose or stopping the medicine in order to observe the effects. This seems to be a popular method with several different types of medicine: antiretroviral therapy, PPIs, antihypertensive medicine, inhalers and non-steroidal anti-inflammatory drugs. Although several of these studies implicitly referred to this type of test, none outlined it in much detail. Dowell and Hudson suggested that the lay testing of medicines may be either an explicit or a subconscious act and that people are more likely to conduct tests if the medicine is to be taken in the long term. They gave an example of a woman with systemic lupus erythematosus who was trying to manage without her steroid therapy:

> Prednisolone, I felt that I could be off them, I thought you know, I was feeling well at the time, I didn’t need to be taking this lot (…) and, err, I mean I did it, they knew I was doing it but it didnae work and I mean I had to go back on them.

This woman suffered rapid weight loss following the cessation of her medicine, which convinced her that she needed to commit herself to the regimen. Another woman, quoted by Buston and Wood, reached the same conclusion after experimenting with her asthma medicine:

> I stopped taking my inhalers for a while [laughs] cos I didnae need them so I just stopped taking them. I never told anyone but I just stopped taking them. I'd been all right for a wee while but then I was in hospital. I was in hospital at the end of the year and I think it's because I stopped taking them for a wee while and then it just built up.

Donovan and Blake found that many of the people with RA in their sample conducted a process of testing or evaluating their medicines before deciding whether or not to take the medicine as prescribed. The authors found that 4 of 41 people taking non-steroidal anti-inflammatory drugs eventually reduced their doses and six gave up altogether. One woman said:

> Feldene [piroxicam] worked OK but they make me feel sick and dizzy. I just stopped taking them because I couldn't do anything when I was feeling giddy … I'd rather just cope with the pain than the pain and the side effects as well.

The authors gave another example of a person who gave up the drug having experimented with it:

> [The doctor] gave me anti-inflammatory tablets. I tried them for a week and they didn't seem to help at all and I don’t particularly like taking tablets anyway, so I stopped taking them a week after I started.

Donovan and Blake noted that some people did not give the drugs sufficient time to work, but were unaware of this.

**Observing others**

One study of women’s experiences of, and attitudes towards, taking AZT found that some women relied on observations of how others fared on AZT in order to come to a decision about whether or not to take it. Siegel and Gorey's participants felt that AZT was experimental and consequently they placed more faith in their observations than in their doctors’ advice, as the following two quotes illustrate.
The doctors try to tell me, ‘Oh it depends on the person,’ or ‘They took it too late’. Well, I’ve made my own research, and my research has told me that those who don’t take it live longer than those who take it. I would not take it … They are experimenting. They are not sure themselves, but they don’t want to say it.

I don’t want to take it [AZT] because from what I, from what I’ve seen, friends of mine that have taken it, they just die quicker, or all of a sudden start getting sick.

Observation of others as a form of evaluation was found only in this one case.

Obtaining information about medicines
Both Dowell and Hudson and Donovan and Blake noted that before accepting their medicines people consult a variety of sources and do not rely solely on their doctors’ advice. Some of Roberson’s sample used ‘root’ doctors as well as informal networks, while Watson and Mitchell’s participants sought information about medicine from pharmacists, friends, peers, relatives and the media, as well as from their GPs. Erlen and Mellors found that information was obtained from support groups, peers, books and the internet when participants were deciding whether or not to take the medicine.

Objective and subjective indicators used in lay evaluation
Johnson et al. found that people used blood pressure monitoring to help them evaluate their antihypertensive medicine. They noted that in order to establish the efficacy of the medicine, ‘Participants needed to sense a reduction in blood pressure and, preferably, controlled blood pressure readings’ (p. 327). Roberson’s study was of medicine in general, but most of her participants were on antihypertensive medication for high blood pressure. She noted that people would use their own or their doctor’s blood pressure readings to help them decide whether or not the medicine was working. This finding was confirmed by Morgan and Van Wissen et al. Blood pressure readings were, therefore, a key tool for monitoring the effectiveness of the antihypertensive medicine.

However, in the case of antiretroviral therapy for HIV infection, there was evident tension between objective and subjective indicators of health, the objective indicators in this case being laboratory results such as T-cell counts. A 46-year-old woman in McDonald et al.’s study illustrates the problem:

… the incentive, I suppose, for somebody to go on to tablets is if they are going to make you feel better, and my experience is that whilst they are physically making me better, and my viral load has dropped and my T-cell count has gone up, so they are obviously doing my immune system good, I don’t feel better on them, and that’s the difficult thing to weigh up.

Siegel et al. and Stone and Clarke found likewise, with a participant in Stone and Clarke’s study making a similar point:

You know, it might serve the purpose, okay, make your T-cells go up or whatever, or make your viral load get to where it’s supposed to, but what about the way you’re feelin’ overall?

People that were affected with HIV were more influenced by subjective indicators, such as symptom alleviation and a sense of health and well-being, than by laboratory results. Siegel et al. noted ‘… participants seemed to rely heavily on their subjective sense of how they were
feeling and their perception of how symptomatic they were in order to assess the efficacy, benefits, and risks of the antiviral medicines they had been prescribed’ (p. 251).

**Gender differences in evaluating medicines**

A few studies\(^ {141,142,149,151} \) found that women were more sceptical than men about antiretroviral therapy. Only a few (mainly white) women believed that the drug could be beneficial under some circumstances. Additionally, women were more sceptical about the scientific basis of drugs for treating HIV infection, arguing that trials had not been conducted with women, so its effectiveness in women was unknown.\(^ {142,149,151} \) Both Siegel and Gorey\(^ {142} \) and Johnston Roberts and Mann\(^ {151} \) found that distrust of drugs was associated with not taking the drugs as prescribed. Some of the women in Siegel and Gorey’s\(^ {142} \) study suggested that doctors were ill informed about the drug, yet failed to acknowledge the limits of their knowledge. Stone and Clarke\(^ {143} \) pointed out that, unlike gay men or injecting drug users, women do not belong to the social networks that people infected with HIV form. They argued that informal social networks are important in promoting confidence in drugs and for circulating up-to-date information about treatments, but women’s lack of access to these networks may account for women’s greater negativity towards the medicine.

**Difficulties with evaluating medicines**

A small number of studies noted that, in the course of evaluating their medicine, a few people had difficulty distinguishing the effects of their medicine from the effects of their illness. Morgan\(^ {134} \) found this in the case of antihypertensive medicine, but Usher\(^ {158} \) found, in the case of neuroleptic medicine, that some people with schizophrenia learned to monitor their bodies in order to discern the difference between the symptoms of their illness and the effects of the medicine. One person explained: ‘…lots of times you get confused; what’s the side effects and what’s the illness? You have to focus on your body and try to work it out, but it’s not easy … the side effects would also get confused with my thinking and get linked to my illness.’\(^ {158} \) In the case of antiretroviral therapy, Siegel *et al.*\(^ {147} \) suggested that patients wrongly interpreted the symptoms of the disease as the undesirable effects of their treatment and, as a result, rejected the treatment mistakenly. Whether or not this is the case is hard to say, but it does highlight the difficulty patients faced when attempting to evaluate their medicine. Particularly with HIV infection, people found it confusing if objective indicators indicated improvement after taking the medicine, but they felt no better or felt worse. In such cases, there might also be disparity between professional and lay assessments of the efficacy of treatment, in which cases patients might feel compelled to ignore their doctors’ advice to continue with treatment.

Dowell and Hudson\(^ {173} \) noted that the method of testing depends on the person’s understanding of the medicine’s function; thus, analgesics with a short-term, symptomatic effect may be easily assessed, but other medicines which cannot be assessed using the presence or absence of symptoms may still be evaluated by conferring with doctors or obtaining information from friends or the media. With preventive medicine, the evaluative process is difficult for patients because immediate symptoms cannot be used as indicators of efficacy. Although health professionals can evaluate medicine in terms of its long-term and preventive effect, lay people can evaluate it only in terms of its immediate impact on their lives. For example, Angermeyer *et al.*\(^ {157} \) found that health professionals prescribed clozapine to reduce psychotic symptoms, but also to prevent relapse, whereas patients primarily evaluated it in terms of its effect on symptoms: the fact that it was calming, relaxing and helped them sleep. Again, it is difficult for lay people to evaluate the effectiveness of antihypertensive medicine given that the anticipated benefits are long term. Because they cannot assess its impact, some people may also be uncertain about whether or not the medicine is necessary, in other words whether or not they actually have hypertension; this uncertainty was an issue in all four studies of hypertension.\(^ {134-137} \)
Occasionally, people might have difficulty evaluating their medicine because they lack information. In one case, where doctors were replacing patients’ usual brand of PPIs for a cheaper brand, they ‘double-switched’ by reducing the dose at the same time as changing the brand. The authors noted that this confused patients because, being unaware of the dose reduction, they could not accurately assess the efficacy of the new drug, so many rejected it as being less effective than their original brand. There was also a lack of desired information on adverse effects. Smith _et al._ found that people with respiratory diseases had to educate themselves about the adverse effects of asthma medicine and lacked knowledge about the appropriate doses of inhalers. They also found that people with arthritis were given inadequate and superficial information on the adverse effects of arthritis drugs and what to do about them. Donovan and Blake _et al._ noted the same in the context of arthritis, and Ersek _et al._ noted that few people with cancer knew how to deal with adverse effects, that some confused tolerance, dependence and addiction and others were unaware that analgesics did not always provide immediate pain relief. The absence of such basic information on medicines and their adverse effects cannot have helped people with their evaluations.

**Worries about medicine that lay testing and evaluation cannot resolve**

Although many of the concerns that people have can be resolved through lay evaluation of medicines, some are less easily resolved and may linger, affecting decisions about medicine-taking.

**Dependence, tolerance and addiction**

Dislike of dependence upon medicines was noted in the context of medicines in general, asthma, cancer, hypertension, mental health and arthritis. Both Donovan and Blake _et al._ and Ersek _et al._ noted that fear of dependence was a reason why some people did not take their medicines. These two studies also reported that participants were worried that they would become accustomed to the medicine and that it would thereby lose its effectiveness; again this resulted in people using less than the prescribed doses. As one woman with RA said:

> I have two tablets at night and I put two by the bed if I need them. But I don’t take them unless it is absolutely necessary. It’s no good taking tablets when you’re not in pain because when you are, your body will be used to them.

Some of Morgan’s participants feared that if they took their antihypertensive medicine as prescribed, they might become unable to manage without the drugs or become addicted to them. As one African Caribbean respondent said:

> Sometimes I remember and sometimes I do not because I don’t want to build my hopes on tablets. I don’t want to become an addict.

Similarly, Angermeyer _et al._ found that a few people expressed fear about becoming addicted to clozapine. Benzodiazepines are well known for their addictive potential, and North _et al._ referred to a spectrum of dependence in their sample, with people who felt that they could control their benzodiazepine use at one end and those who were completely reliant on the medicine at the other end. There also appeared to be a spectrum of attitudes to dependence, with participants at one end believing that constant low doses represented only ‘partial dependence’ as compared with dependence on escalating doses. Barter and Cormack found that only a small number of participants in their sample worried about dependence on benzodiazepines.
Potential harm from long-term use

Three studies\textsuperscript{134,159,160} reported people’s fears about taking medicines on a long-term basis. Morgan\textsuperscript{134} found this to be a major concern, expressed by both men and women, both white and African Caribbean, as the following quotes illustrate:\textsuperscript{134}

I have thought about adverse effects. If you keep on with these tablets, what is it going to do eventually? That’s what bothers me.

I used to say to my friend at work that these tablets are going to affect my insides. She said don’t be silly … taking so much of them each day and for so long, they might do something to me.

Both of the studies on PPIs\textsuperscript{159,160} found that people’s generally positive opinions of the medicine were tempered by their concern about the potential for long-term harm. Pollock and Grime\textsuperscript{160} noted that the people in their study who were worried about the long-term effects of taking PPIs were more likely to experiment with their regimen to achieve the lowest possible dose. The authors noted that this is desirable ‘in view of the clinical uncertainty about possible hazards of long term acid suppression’ (p. 1837).\textsuperscript{160}

The possibility of medicine masking other symptoms

Two of Boath and Blenkinsopp’s\textsuperscript{159} participants were concerned about the potential of PPIs to mask symptoms that they should know about, symptoms that might indicate a more serious condition such as cancer. A woman with cancer in Ersek \textit{et al.}\textsuperscript{168} study was also worried about this. She was reluctant to take analgesics in case it masked new pains that might indicate an infection that she should be aware of.

Medicine and identity

Not accepting the illness

Dowell and Hudson\textsuperscript{173} found that acceptance of medicines was linked to acceptance of the illness in question. They suggested that taking medicine was equated with having an illness, so if people did not accept that they had the illness then they were unlikely to continue with or start the treatment. Acceptance was a strong theme in the asthma studies. Adams \textit{et al.}\textsuperscript{161} reported that some people with asthma denied that they had asthma, or distanced themselves from their asthma. Although these people – termed ‘deniers and distancers’ by the authors – had asthma that was just as severe as it was among those who accepted their asthma, the deniers/distancers tended to downplay its significance, claiming that they did not have asthma at all or that they had only slight or ‘not real’ asthma. They viewed their asthma as acute rather than chronic and did not take the preventive asthma medicine as this entailed acceptance of the asthma identity and the chronicity of the condition. However, they did take the reliever medicine ‘just in case they demonstrated symptoms in social/public situations’.\textsuperscript{161} As such, they were found to use relievers and avoid preventers for the same purpose: maintenance of normality.

Proust \textit{et al.}\textsuperscript{162} also found a tendency among the people in their study to downplay the severity of asthma, as did Busto and Wood.\textsuperscript{163} In Proust \textit{et al.}’s\textsuperscript{162} study, families did not define childhood asthma as a serious illness, preferring to stress the ordinariness of children instead. As with some of Adams \textit{et al.}’s\textsuperscript{161} sample, Proust \textit{et al.}\textsuperscript{162} found that some of their families doubted that their children actually had asthma. The authors argued that inhalers were enrolled into the task of maintaining this sense of ordinariness. Walsh \textit{et al.}\textsuperscript{164} identified three categories of people who did not take their asthma medicine as prescribed: those in denial, those in avoidance and those in depression. Those in denial did not want to see themselves as having a chronic illness and felt that they could manage the asthma on their own; they were therefore motivated to reduce their
medicine. Those in avoidance felt that nothing helped resolve their breathing difficulties, so they did not attend clinic appointments and neglected their regimens. These two categories resemble the people in Adams et al.\textsuperscript{161} study who denied or distanced themselves from the asthma, but those in Walsh et al.\textsuperscript{162} third category, depression, are unlike any described in the other asthma studies. These people accepted that they had asthma and, in as far as this goes, they are similar to those in Adams et al.\textsuperscript{161} study who accepted their asthma, but here the similarity ends because Walsh's et al.\textsuperscript{162} 'depressed' group keenly felt the loss of their previous abilities and also a vulnerability that led them to restrict their lives in order to avoid asthma attacks. They became more and more withdrawn and depressed until they neglected their regimen and became ill. By contrast, Adams et al.\textsuperscript{163} 'accepters' appeared to be well balanced and had come to terms with the asthma and the need for medicine.

Only one\textsuperscript{152} of the HIV studies and one\textsuperscript{158} of the studies on psychotropic medicine dealt with the issue of illness acceptance. Barton Laws and Wilson\textsuperscript{152} suggested that the ability to confront the fact of being HIV positive is a key factor determining whether or not people take their drugs as prescribed, whereas Usher\textsuperscript{158} noted that the acceptance of schizophrenia is a necessary precursor to the acceptance of neuroleptic medicine. Usher\textsuperscript{158} observed that this acceptance occurs over time and is not easy as it entails taking on a stigmatised label, the unlikelihood of recovery and the need to take medicine long term.

**Being reminded of the illness**

Medicine can also be a constant and unwelcome reminder of illness, whether or not that illness is accepted. Four of the HIV studies found this.\textsuperscript{144–146,151} One of the men in Halkitis and Kirton's\textsuperscript{145} study said:

\[
\ldots\text{it's hard every time you take that pill, to recognize what you're taking it for, and to know that this is going to be probably the course of my life…}
\]

One woman wrote in her journal for Johnston Roberts and Mann's\textsuperscript{151} study:

\[
\text{Each time I take a pill it is a reminder that I am sick. I do not want to focus on my sickness or let it control my life. I want so badly to be 'normal' and healthy again.}
\]

Only one other study reported that taking medicine reminded people of their illness, in this case cancer.\textsuperscript{168}

**Disclosing the illness to others**

A separate but related issue in the case of HIV infection (and a significant reason why some people did not always take their medicine as prescribed) was fear of disclosing their illness to others.\textsuperscript{143,145,148,150,151} Consequently, rather than take the medicine in front of people and risk disclosing their HIV status, people might postpone or forgo their medicine.\textsuperscript{151} A participant in Stone and Clarke's\textsuperscript{143} study said:

\[
\text{If you don't want everybody to know, you know, you don't take 'em in front of people. It raises questions.}
\]

This could include public social situations such as parties and the work place, but also private, family life. One woman in Johnston Roberts and Mann's\textsuperscript{151} study who had not disclosed her HIV status to her children found it hard to take all her pills because she had to do so out of sight of her children.
The stigmatising effects of medicine

Some of the people in Rogers et al.’s study reported that neuroleptic medicine was stigmatising because it could mark out a person as having schizophrenia. Similarly, Smith et al. found that those with mental-health problems reported feeling stigmatised and labelled by their medicine, as did Usher, again in the case of neuroleptic medicine. Her respondents reported that the adverse effects of the medicine drew attention to their illness, as one person illustrated:

When you go out it’s like advertising you have a mental illness, so the side effects draw attention to the fact that you have a mental illness. And even though you might be quite well mentally, the side effects stigmatize you … you cannot even go over to your sister’s place and go out into the yard without the neighbours thinking she’s got someone there who is mentally ill … you know your legs are going up and down all the time and they think you’re a lunatic. It’s like wearing a sign on your forehead.

Other studies in the psychotropic medicine group did not cover this issue, although North et al. noted that some people reported feeling ashamed of using benzodiazepines. In the context of HIV infection, Barton Laws and Wilson found that some people did not initiate treatment because they worried that the regimen would identify them as having HIV. Finally, Atkin and Ahmad found that for children with thalassaemia major, one of the most hated aspects of their chelation therapy was the fact that it marked them out as different from their peers.

Ways of taking medicine

A small number of studies categorised or characterised the various ways in which people take their medicines. This section begins with these studies, before going on to examine the range of modifications people make to their regimens, the reasons behind them and how far these are communicated to doctors.

Those who take their medicine as prescribed (Figure 14)

Passive accepters

Dowell and Hudson described people who accepted the regimen that they had been prescribed, who assumed a more passive role and who relinquished control to their doctors as ‘passive accepters’. Similarly, in the context of hypertension, Svensson et al. reported that those who accepted the prescribed regimen were passive, indifferent and less involved in their care. The authors noted that many had ‘rather obscure motives for continuing with treatment, and often claimed not to have considered the possibility of changing or omitting medication’ (p. 162).

Active accepters

In the context of hypertension, Johnson et al. developed a category of medicine-taking which they termed ‘purposeful adherence’. ‘Purposeful adherence’ followed a conscious decision to pursue the prescribed regimen and was dependent upon having first tested the medicine to ensure its effectiveness. Those Morgan described as ‘stable adherents’ and ‘problematic adherents’ – also in the context of hypertension – could be seen as practising ‘purposeful adherence’. These people took their medicine as prescribed but the ‘problematic adherents’ expressed concern about doing so and had worries about adverse effects and dependence, whereas the ‘stable adherents’ seemed unconcerned, even by the prospect of remaining on the tablets for all of their lives. Although ‘stable adherents’ seem similar to Dowell and Hudson’s ‘passive accepters’, they were not necessarily passive and may have reached their position after a period of testing their medicines.
Accepting accounts

Both active and passive accepters, but especially the latter, might give a more ‘orthodox’ account of their medicine-taking. In Britten’s study of lay accounts of medicine-taking, she developed an analysis based on ‘orthodox’ and ‘unorthodox’ accounts. Those giving ‘orthodox accounts’ were more likely to describe the ‘correct behaviour’ for taking medicines, such as cashing prescriptions, taking the medicine as prescribed, following the doctor’s advice even if they had doubts about it and not wasting the doctor’s time. These people appeared to be more deferential to doctors, talked about medicine in an unquestioning manner and were generally passive.

Similarly, those offering what Lumme-Sandt et al. described as the ‘patient repertoire’ offered ‘very traditional representations of themselves as patients’ (p. 1847) and regarded doctors as experts who should be obeyed. Medicine was simply taken for granted.

Those who do not take their medicine as prescribed (Figure 15)

Rejecters/sceptics

Dowell and Hudson reported that some people rejected their prescribed medicine outright, without testing it first. These people, whom the authors describe as ‘rejecters’ or ‘sceptics’, preferred to maintain control by using ‘alternative’ therapies or by tolerating their symptoms. They completely rejected Western medicine and its emphasis on drug taking. In the context of hypertension, Johnson et al. noted that some people, whom they termed ‘purposeful non-adherers’, made a conscious decision not to take the medicine, but in this case possibly following a period of testing.

Rejecting accounts

Both the ‘active modifiers’ (see below) and the ‘rejecters’ (but more likely the ‘rejecters’) might give ‘unorthodox accounts’ or draw on the ‘self-help repertoire’, both of which imply a rejection of modern medicine. Britten found that those giving ‘unorthodox accounts’ were ‘almost entirely negative’ (p. 60) about medicines in general. These people preferred not to take medicine, describing it as unnatural, damaging, dealing with symptoms rather than causes and not being tailored to individuals. They were critical of doctors, mainly because of overprescribing,
Resisting medicines: a synthesis of qualitative studies of medicine-taking

and their criticisms of doctors and medicines went hand in hand. These people were generally active in their approach. Similarly, those drawing on the ‘self-help repertoire’ described by Lumme-Sandt et al.\textsuperscript{175} had strong negative views about medicine, and some explicitly reported that they did not follow their doctors’ advice. They reported using both medical and home remedies side by side, but had greater confidence in the latter.

**Those who modify their regimen – the ‘active modifiers’**

Donovan and Blake’s\textsuperscript{166} study of RA is the earliest in the synthesis to describe the characteristics of those who do not take their medicine as prescribed. Most of Donovan and Blake’s\textsuperscript{166} sample stated that they disliked taking drugs, most chose to ignore advice or alter doses and few saw this as an issue. The authors suggested that modification of the regimen represented a means of gaining some control over the illness. Similarly, Dowell and Hudson\textsuperscript{173} found that a number of people in their sample took control by making a conscious decision to modify the regimen following a period of deliberation and testing. These people, whom the authors describe as ‘active users’, consciously decide to modify the regimen, following period of deliberation and testing. Active users take control by modifying their regimen. May be critical of doctors and medicine in general

Dowell and Hudson, 1997\textsuperscript{173}

‘Active users’ consciously decide to modify the regimen, following period of deliberation and testing. Active users take control by modifying their regimen. May be critical of doctors and medicine in general

Svensson et al., 2000\textsuperscript{137}

Characteristics of those who do not take drugs as prescribed: attempting to gain control; minimise impact of drug on lives; and taking active part in decision-making. Qualities associated with improved outcome and in line with health-care objectives

Siegel et al., 2000\textsuperscript{148}

‘Justifiers’ had chosen not to take drugs as prescribed and justified modifications to regimen by arguing that flexibility won’t hurt and strict adherence not necessary, not possible

‘Excusers’ argued that although they wanted to adhere this was not possible due to side effects, or circumstances

**FIGURE 15** Categories of those who do take their medicine as prescribed.
regimen on the basis that some flexibility would not hurt and that strict adherence was neither necessary nor possible, and ‘excusers’ claimed that they wanted to adhere to the regimen, but that various circumstances and adverse effects prevented them from doing so.

Before going on to look in detail at the various ways in which people modify their regimens, it should be noted that there is another category of medicine-taking that none of the above studies considered, but which came out strongly in the studies on mental illnesses: those who are forced or strongly coerced into taking their medicines.

‘Imposed compliance’

Usher\textsuperscript{158} used the term ‘imposed compliance’ to describe the result of the pressure exerted by relatives or health professionals to take medicine. Writing in the context of neuroleptic medicine for schizophrenia, she suggested that some people take their medicines only because they feel powerless to do otherwise. Some of her participants reported that friends and relatives surveyed them for signs of illness and, if they felt it necessary, exerted pressure on them to take their medicine. Rogers \textit{et al.}\textsuperscript{156} also found this to be true in the case of neuroleptic medicines. Some of their participants had experienced relatives or friends strongly encouraging or forcing them to take the medicine, as one of their participants illustrated:\textsuperscript{156}

> My husband is supposed to sort of see that I take it, I don't see that it's up to your husband to see that you take your medication, it's up to you … but my husband gets blamed if I don't take my medication.

A different type of pressure was found in the context of HIV infection. McDonald \textit{et al.}\textsuperscript{149} noted that some relatives or partners might encourage the HIV-positive person to take the medicine because they wanted their loved one’s lifespan to be extended. As one of their participants said:\textsuperscript{149}

> … I mean, when you’re in a relationship you have to take into consideration your partner’s thoughts and feelings on things and what she sees in tablets is ten or more years of being together … And that’s what she wants of course and to some extent of course I want it too but I don’t want it if I’m going to be sick all the time taking them.

At the other end of the scale, Kaljee and Beardsley\textsuperscript{153} noted that some patients with mental illness have no control at all over their medicine because it is administered by injection. This is the most extreme form of ‘imposed compliance’. Somewhere in between the two ends of the scale is the pressure to take medicine that is imposed by communities. Rogers \textit{et al.}\textsuperscript{156} suggested that people on neuroleptic medicine are more likely to take their drugs as prescribed because of the threat of social sanctions (for example social disapproval, withdrawal of acceptance, hospitalisation) if they do not. Their participants perceived the existence of an unwritten social contract: take the medicine in order to be tolerated by the community. Furthermore, some had experienced coercion from health professionals when in the past they had not taken their medicine, as a woman in Rogers \textit{et al.}'s\textsuperscript{156} study illustrated:

> You’re supposed to do as they [doctors] tell you, you know, otherwise you get sectioned and I have been sectioned so er it's not very nice, police bringing you down to [local psychiatric hospital], so, you take them to stop yourself from being sectioned.

Similarly, one of Usher’s\textsuperscript{158} participants said:

> The community nurse comes to see if I’m taking my medicines … I call them the thought police … I felt like I didn’t have any choice.
Some of the people with mental illness in Smith et al.'s study also felt that medicine was used to control them and make them acceptable to society. As the authors wrote, ‘Medicines for mental health problems were seen as a tool used by professionals to control individuals within a social context. The aim of treatment was viewed as a means to achieve behaviour acceptable to society’ (p. 93). As can be seen then, imposed compliance was almost exclusively an issue for people with mental illnesses.

**A general desire to minimise medicine intake**

Dowell and Hudson suggested that people take medicine within limits, either those established by the doctor or self-imposed. They noted that people ‘… have a powerful drive to minimize treatment use. Indeed, the more powerful the drug, the more its importance is emphasized and the stronger may be the desire to reduce it’ (p. 375). Few of the rural African Americans in Roberson’s study took their drugs as prescribed, and most talked openly about this. They regularly reduced the number of doses that they were supposed to have, for example by taking them once rather than twice daily. A general impulse to minimise the intake of tablets was also found in the context of treatment for arthritis. Donovan and Blake found that more than four-fifths of all the people in their sample spontaneously expressed their dislike at having to take drugs. Watson and Mitchell found that the older women in their sample avoided taking medicine unless they felt it to be essential. People taking PPIs also preferred to minimise their intake, despite PPIs being regarded by doctors as a medicine that is overconsumed by patients. Similarly, North et al. found that many people taking benzodiazepines, which are also regarded by doctors as being overused, set self-imposed limits on their consumption of this medicine.

Although the above authors referred to a general desire to minimise the intake of medicine, many of the following strategies reflect a similar desire.

**Adjusting doses to decrease adverse effects and addiction**

As noted earlier, many of the HIV studies reported people’s concerns with the adverse effects of the antiretroviral regimen. The following studies illustrate the ways in which people adjusted their regimens to try and realise some therapeutic gain at the same time as reducing adverse effects. Siegel and Gorey found that many of the women who took or had taken AZT reduced the dosage, for example by taking it three or four times daily rather than five times, or by skipping a day here and there. Some of Siegel et al.’s older respondents reported that if altering the dose reduced the adverse effects and improved their well-being, then they felt that such alterations were acceptable. Some would stop taking the medicine completely to reduce the debilitating adverse effects, whereas others would interrupt their regimen if the adverse effects were too severe, then continue once the effects had abated. One man, when asked if he took his medicine as prescribed, answered:

> No, it’s too strong … You’re not supposed to stop taking the medicine. It is supposed to be taken consecutively so that it can have effect. But since I start taking it and I see that it is attacking parts of my body, that is more dangerous. I will stop taking the medicine … if my liver begins to hurt, or my prostate, or this or that, I stop taking it for up to two days.

Some of Erlen and Mellors’ participants who were infected with HIV reported taking their tablets separately instead of all together, to avoid adverse effects. Several of Siegel et al.’s sample reported reducing their doses, or taking their medicine at different times, to reduce adverse effects. One man explained:

> Yes, I decided that I would double up [doses] at the end of the day because it [medication] was giving me diarrhoea so bad. And I figured like I didn’t want to be out
in public and have to go, and not be able to have any place to go. So what I would do, I would take one at mid-day when it wouldn't bother me too bad. But the morning dose, I would not take, because if I did and had to go out, then I would be in trouble.

Some people specifically referred to ‘drug holidays’ as a means of stopping the build-up of toxicity and of cleansing their bodies.144,152 Several of Stone and Clarke’s143 participants reported going without medicine for periods ranging from 1 week to 1 month or more. One person said:143

Every once in a while l’d have drug holidays, and you do feel like a toxicity’s building up, and so on and so forth. That something just tells you to stop. And you stop for, like, a week.

One of Barton Laws and Wilson’s152 respondents reported taking a 6-week break during which his viral load had ‘shot up’, but he did not believe that the practice was harmful and was convinced that almost every person infected with HIV took drug holidays. Morgan134 found that people also took drug holidays in the context of antihypertensive medicine, in the hope of reducing the total drug intake and reducing the risk of adverse effects, dependence and addiction. She suggested that ‘leaving off’ the drugs took two forms: one method was to stop taking the medicine for a few days each week regularly and the other was to take a break of several weeks or months if the person was feeling well. Donovan and Blake166 suggested that people modified their regimen to reduce their fears about adverse effects. ‘They reasoned that there would be fewer side effects with fewer tablets, and so kept their doses to the minimum they could. In some cases, this meant them putting up with considerable amounts of pain and discomfort . . .’ (p. 509).166 North et al.154 reported that many people tried to restrict their use of benzodiazepines in order to avoid addiction. The authors noted that doctors gave people a high degree of autonomy in the management of their benzodiazepines. Most people were found to set self-determined controls on their doses and many were trying alternative ways of managing their symptoms owing to their ambivalence about using benzodiazepines.

Adjusting the regimen to make it more acceptable

Some of Siegel et al.’s147 participants amended their antiretroviral regimen to fit in with their daily schedule, arguing that complete adherence was not necessary for therapeutic gain. Similarly, some of those in Siegel et al.’s148 study were not worried about straying from their regimen, feeling that some flexibility in the regimen was acceptable and harmless, as the optimum regimen was not known anyway. They argued that flexibility with the regimen allowed life to continue without too much disruption and that strict adherence to the regimen was neither an attainable nor realistic goal. In the same way, Barton Laws and Wilson152 found several people adjusted their doses or the timing of their doses to fit in with their daily routine rather than modifying their daily routine to accommodate the regimen. One person, who took his medicine whenever he ate rather than every 8 hours, said:152

They’re not spaced like they’re supposed to, but I know enough about the medication where I know that they still overlap … These medications don’t flush out of your system in 8 hours like they make … people believe.

Another intrusive regimen was the nightly infusion pump for administering chelation therapy to children with thalassaemia major. Atkin and Ahmad169 reported that some children would use the infusion pump less frequently than prescribed or would disconnect the needle at night. The authors suggested that children did this because the therapy dominated their lives and marked them out as different from their peers, thus making them feel physically and symbolically tied to the pump and the regimen.
Reducing doses for financial reasons

Johnson et al.\textsuperscript{136} noted that in order to take medicines, people need insurance or sufficient personal income to purchase or access their treatment. Some of Roberson\textsuperscript{8,171} sample reduced doses because they could not afford the prescribed amounts. One of her respondents with hypertension, diabetes and arthritis was on a low income and had no health insurance. Her medicines cost between US$100 and US$125 per month. She said, ‘If I see I’m about to run out and I don’t have no money to get no more, I just cut down my pressure pills to once a day’ (p. 16).\textsuperscript{171} Financial difficulties that impacted on medicine-taking were also reported by Ersek et al.,\textsuperscript{168} Barton Laws and Wilson\textsuperscript{152} and Siegel et al.\textsuperscript{148}

Using medicine symptomatically

Morgan\textsuperscript{134} is the only person to have considered the influence of ethnicity on medicine-taking. She found that some of her African Caribbean respondents used their antihypertensive drugs in the same way as they used their herbal remedies, in other words, symptomatically. She noted that although high blood pressure is regarded as an asymptomatic condition, some of the people in her study described feelings of weakness or tiredness, eye problems, dizziness or hotness and perceived these as indications that their blood pressure was elevated. If this was the case, they might be prompted to take the medicine. One respondent said:\textsuperscript{134}

Please don’t tell him [Dr] this, but I’m a person that if I find it’s stable and I’m not getting any funny feelings then I don’t take them. If I feel peaky again, then I go back on them.

Similarly, one of Johnson et al.’s\textsuperscript{136} respondents reported taking his or her antihypertensive medicine sporadically, when his or her blood pressure readings was elevated. Roberson\textsuperscript{171} found likewise, that some people would check their blood pressure and decide on dosage accordingly. Some of the people with RA in Donovan and Blake’s\textsuperscript{166} study altered their doses according to the symptoms they experienced. As one woman put it:\textsuperscript{166}

When [the arthritis] is bad, I take two [Feldene, i.e. piroxicam] because that’s how it’s directed, but otherwise, if I get a twinge, I only take the one. There is days I don’t take any. I’d rather not take them if I can.

Kaljee and Beardsley\textsuperscript{153} suggested that some people on psychotropic medicine manipulate their drug doses to control symptoms. One person diagnosed as having manic depression felt that his drugs provided immediate relief from his symptoms: ‘… a couple of sinking feelings, take a pill, (the feelings) would go away\textsuperscript{2,155} A few of Rogers et al.’s\textsuperscript{156} participants adjusted their doses of neuroleptic medicine to cope with distress when it arose, and Angermeyer et al.\textsuperscript{157} also found that some people with schizophrenia used clozapine symptomatically.

Using medicine strategically

Some of Morgan’s\textsuperscript{134} respondents reported that they would not take their antihypertensive medicine if they intended to drink alcohol, for fear that the interaction would have severe consequences:\textsuperscript{134}

I am a man that goes out to parties sometimes and has a nice drink. When you are taking drugs you have to limit your drinks. If I know that I’m going out on a Saturday night or Friday or Sunday I won’t take the tablets as you do not want to mix the drugs with the alcohol.

Interviewer: Why is that?

It is not something you should do, you can die if you mix them.
Similarly, Rogers et al.\textsuperscript{156} found that several people adjusted their doses of neuroleptic medicine when they wanted to go out for a drink, because they knew that alcohol interacted poorly with their medicine. Most stopped taking their medicine on the day they drank alcohol; thus the need for social activity was weighed against the need for medicine. The authors of the studies on PPIs reported that some people took their medicine strategically rather than continuously.\textsuperscript{159,160} Boath and Blenkinsopp\textsuperscript{159} found that some took PPIs as and when they thought it necessary, relating the dosage to their proposed diet. Thus, some might take a pill in advance of a planned indulgence, whereas others might miss a dose if they did not anticipate a need for it. Pollock and Grime\textsuperscript{160} also found that several of their participants took PPIs only at symptom onset, or if symptoms were anticipated.

**Replacing or supplementing medicines with non-pharmacological treatments**

Most of Roberson’s\textsuperscript{171} sample undertook ‘self-care activities’ in addition to their prescribed regimen. Vinegar, Epsom salts, lemon and garlic were taken for hypertension, BENGAY\textsuperscript{8} (Johnson & Johnson, New Brunswick, NJ, USA) and copper wire bracelets were used for arthritis, horseradish tea or buttermilk for diabetes and rabbit tobacco tea and pine top for asthma. A few people sought advice from ‘root doctors’, who are described as having an extensive knowledge of herbs and roots. However, although Roberson\textsuperscript{171} noted that people took traditional remedies in addition to their prescribed medicine, it is not clear whether or how this affected people’s drug regimens. Lumme-Sandt et al.\textsuperscript{175} noted that their respondents who spontaneously talked about traditional remedies seemed to use them as their primary medicine. As one man said, ‘Yes, I have slight blood pressure and I eat garlic for that. I didn’t take any medicine when the doctor told me to. I cure it with garlic’ (p. 1847).\textsuperscript{175} Others used both medical drugs and traditional remedies alongside each other.

Several of Ersek et al.’s\textsuperscript{168} participants used non-pharmacological methods to reduce pain and also to lower their use of analgesics, and about half of the people in Donovan and Blake’s\textsuperscript{166} study used some sort of ‘alternative’ remedy either instead of or as well as their medicine. Kelp, cod liver oil, feverfew, dietary changes and homeopathy were mentioned. Morgan\textsuperscript{134} found that over half of the African Caribbean people in her study took herbal remedies. Respondents reported taking cerasee, which was described as good for reducing high blood pressure, and Constitution Bitters, a drink containing a blend of seven herbs. The African Caribbean women boiled grapefruit skins with garlic and drank the water to reduce blood pressure. Morgan\textsuperscript{134} noted that these herbal remedies formed an additional resource for many people and that they were taken either in addition to or instead of their prescribed medicines. Furthermore, she observed that people were familiar with these remedies and regarded them as potentially less harmful and powerful than prescribed drugs because they were seen as ‘natural’ substances. If people were worried about the potentially harmful effects of prescribed drugs they might take a break from them for a period and use natural remedies instead.

**Temporary breaks in regimen for incidental reasons**

Several studies reported forgetfulness to be a reason why some people occasionally did not take their medicine as prescribed,\textsuperscript{136,143,144,148,151,152,168,171} and three studies\textsuperscript{156,146,153} reported that occasionally people unintentionally ran out of tablets. Interruption in routine or an irregular daily schedule could lead to a break in the regimen.\textsuperscript{136,143,148,150–152} Strategies for minimising what Johnson et al.\textsuperscript{136} referred to as ‘incidental non-adherence’ were developed mainly by those infected with HIV. Stone and Clarke,\textsuperscript{143} Halkitis and Kirton\textsuperscript{145} and Murphy et al.\textsuperscript{152} reported that people used mechanical reminders to take their pills (including beeper pill cases, weekly or monthly pill boxes, stopwatches or alarm clocks) or visible cues. Reminders from significant others were found to help in the context of asthma\textsuperscript{163} and HIV infection.\textsuperscript{143,145,150} The routinisation of medicine-taking was a key strategy for those infected with HIV,\textsuperscript{143,145,150} hypertension\textsuperscript{136} or asthma.\textsuperscript{163}
Doctor–patient communication about regimen modifications

A small number of studies suggested that a better relationship with one's doctor meant that patients were more likely to take their medicines as prescribed\textsuperscript{137,143,144,173} or accept prescription changes,\textsuperscript{157} whereas a poor relationship with the doctor could lead to not taking medicines as prescribed.\textsuperscript{150} More specifically, with regard to regimen changes, some of Roberson's\textsuperscript{171} respondents reported that they were scolded by their doctors for making their own decisions about their health care. However, instead of confronting them, patients would change doctors whom they were dissatisfied with. Britten\textsuperscript{172} also suggested that people who gave 'unorthodox accounts' tended not to mention their beliefs or criticisms to their doctors because they did not perceive their criticisms to have medical legitimacy. Thus, people giving 'unorthodox' accounts appeared 'orthodox' while in the surgery. Once outside the surgery, however, they regained control by modifying or rejecting their prescription, offering an unseen challenge to the medical system.

Most of Ersek et al.'s\textsuperscript{168} sample reported positive relationships with care providers, but a minority felt that doctors did not communicate well about medicine. Smith et al.,\textsuperscript{170} whose study included people with arthritis, respiratory disease and mental illness, found that people with arthritis varied in their attitude to involvement in decision-making about medicine, and those with respiratory disease wanted more involvement. People with mental-health problems reported having generally poor relationships with health professionals and participation in decision-making was rare for this group, particularly during the acute phases of their illnesses. Also in the context of mental illness, Rogers et al.\textsuperscript{156} observed that people with schizophrenia who did not take their neuroleptic drugs as prescribed were unlikely to reveal this to health professionals because of their awareness or experience of coercion and their knowledge of the power that health professionals held over their lives. One person put it like this:\textsuperscript{156}

He's [psychiatrist] bawling and shouting at me in my own house, and quite a few other people have had this experience, he's got a very poor personal style. I feel intimidated by him and he's over-bearing and he's got the power to section me, he's got the power to do just whatever he wants to me so I'm very careful what I say to him.

Another of Rogers et al.'s\textsuperscript{156} participants felt unable to discuss her use of 'alternative' remedies with the psychiatrist because she felt that he would consider this as further evidence of mental illness. Similarly, Kaljee and Beardsley\textsuperscript{153} noted that patients' use of alternatives (e.g. prayer) to drugs for mental illness could lead care providers to label them as 'uncompliant'.

Evidently then, although people are indirectly challenging medicine outside the surgery, in the day-to-day practice of using alternative remedies, discarding their prescriptions and modifying regimens to suit themselves, no resolution occurs between doctors and patients on this issue. Some suggestions about resolving the issue have been made. In the context of PPIs, Pollock and Grime\textsuperscript{160} argued that the best solution is to let people find their own dose and manage it themselves, given that their tendency is to under- rather than overmedicate. They recommended that patients be encouraged to actively manage their illness by finding their own level of treatment and suggested that doctors need to recognise that many patients do this already. They argued that more patients would self-regulate if doctors told them to do so and that if the practice were authorised and open, with doctors guiding patients, it would be safer than it is at present. They suggested that self-regulation would involve people in their own care and help them feel in control. Similarly, Dowell and Hudson\textsuperscript{173} suggested that as there is a powerful drive to minimise the use of medicines and as patients will continue to test and modify their medicines, doctors may as well assist them in doing so.
Svensson et al. reached similar conclusions with regard to antihypertensive medicine. They suggested that 'non-adherence' should be encouraged because it is a way for patients to gain control and take an active part in decision-making. They suggested that it ‘must be considered sound behaviour, the danger of which is only apparent when it occurs frequently and is not communicated to health professionals’ (p. 162). The authors further claimed that taking an active part in care and gaining control over illness are practices associated with improved therapeutic outcomes and are also in line with health-care objectives. Conversely, they noted that indifference and passivity may co-exist with adherence but are not qualities associated with informed consent or shared decision-making, which are assumed to be characteristics of good clinical practice.

### Summary of main findings

A small number of sociological studies investigated medicine-taking in its own right, but the absence of compliance as a starting point was rare in the studies and the ideology that people should take their medicines as prescribed held sway. The majority of the studies focused on the reasons why people do not take their medicine as prescribed, with few considering the experience of those who reject medicine outright or those who accept it uncritically. Additionally, the vast majority of studies were concerned with chronic illness.

The synthesis revealed widespread caution about taking medicines and highlighted the lay practice of testing and evaluating medicines prior to accepting or rejecting them, principally owing to worries about adverse effects and other unwanted consequences of taking medicine. Although some concerns about medicines may be resolved through lay evaluation, many cannot be dealt with in this way. These include worries about dependence, tolerance and addiction, the potential harm from taking medicines in the long term and the possibility of medicines masking important symptoms. Additionally, it is clear that in some cases medicine has a significant impact on identity, presenting problems of disclosure and stigma, among others.

People were found to accept their medicine either passively or actively, or to reject it. Additionally, some were strongly coerced into taking medicines. Most of the studies focused on those who did not take their medicines as prescribed. Regimens might be modified symptomatically or strategically, adjusted in order to decrease adverse effects and other unwanted consequences, or adapted to fit more acceptably into daily life. Many of the modifications reflected a general desire to minimise the intake of medicines and this desire was echoed in some people's use of non-pharmacological treatments to either supplant or supplement their prescribed regimen. Few discussed regimen modifications with their doctors.

The lines of argument presented in this synthesis, and summarised textually in this section, are displayed in the form of a model of medicine-taking in Figure 16.

### Discussion

It is important to remember that our sample of studies represented only a small range of medicines, taken mainly for chronic illnesses. Furthermore, the illness most frequently represented in the studies was HIV infection, which provided relatively extreme experiences of medicine-taking. A person's experience of medicines is likely to differ according to the medicine in question and the nature of the illness for which it is taken. For example, it is possible that those classified in some of the studies as 'rejecters' could become 'accepters' if they were given
a different medicine or vice versa. Equally, it is important to remember that in some cases the consequences of not taking medicines as prescribed can be serious or fatal.

**Lay evaluation**

At least two studies preceding our synthesis considered the topic of lay testing, but we found only one study that specifically set out to investigate this issue. In common with our findings, the participants in the study by Arluke, which also predated our synthesis, tested their medicines in a number of ways including observing the effect of stopping the drugs or reducing the dose, and observing the reactions of others on the same medicine. However, although some lay people conduct their own tests of medicines this does not imply that they are ‘experts’ in the field. Prior criticised the sociological inclination to treat lay knowledge as ‘every bit as valuable’ as professional knowledge, arguing instead that lay knowledge is often partial and restricted. An example from the synthesis that may support this is the gender difference evident in lay evaluations of treatment for HIV infection. McDonald et al. argued that gay men collectively engaged with the AIDS (acquired immunodeficiency syndrome) epidemic and achieved an unusual level of biomedical expertise on the subjects of HIV infection and trial methodologies, as well as credibility within medical circles (see also Epstein). On the other hand, women's decisions tended to be based on ‘personal and idiosyncratic health histories or current or very recent health status. Women were simply unused to making judgements about the efficacy of HIV treatments and their frame of reference was not historically and communally situated, as it was for the men.

However, we believe that the significance of the lay evaluation of medicines lies not simply in the fact that it occurs, but in the reasons why it does. The urge to evaluate suggests widespread caution with regard to taking medicine as well as the possibility that people do not completely trust what they have been told about their medicines. It may also reflect an (instinctive) understanding of the principle of individual treatment response and a desire to ensure that the medicine is suitable in their particular case. As Johannessen explained: 'The controlled single subject trial is concerned with determining the efficacy of a drug compared with placebo or another drug in one particular patient. Such trials can also be employed to assess the optimal dose of a drug, to identify adverse effects, or to stop established, unnecessary treatment by convincing the patient that there is no benefit beyond the placebo effect. In N of 1 trials each subject serves as his or her own control.'

The principle of the 'N of 1' trial methodology is often compared with the strategy of 'trial and error' in clinical practice, where perhaps the effect of a therapy or the optimal dose is uncertain. Lay people, in the way in which they test their medicines, are attempting to determine whether or not there are any adverse effects and whether or not the medicine works for them and, if so, to establish the most suitable dose for themselves. This suggests that lay people may have an intuitive understanding that population set doses may not be appropriate for individuals. Recently, the worldwide vice president of GlaxoSmithKline acknowledged that most prescription medicines do not work on most people who take them. Steve Connor, the Science Editor for the Independent, wrote ‘It is an open secret within the drugs industry that most of its products are ineffective in most patients’ (see also Smith). Although this may be an open secret within the drugs industry, it is well concealed from those who take the medicines. Fortunately, as this synthesis suggests, many lay people work this out for themselves and modify their regimens accordingly. In support of a flexible approach to medicine-taking, a study on the use of non-steroidal anti-inflammatory drugs found that an adaptable approach that involved varying the dose according to the presence of symptoms was less likely to result in hospital admission for upper gastrointestinal bleeding than was rigid compliance to prescribed doses.
Adverse drug reactions

The literature on adverse drug reactions (ADRs) suggests that people are right to be cautious about taking medicines. A meta-analysis of studies in the USA found the incidence of fatal ADRs (excluding those because of prescribing errors) to be between the fourth and sixth leading cause of death, whereas in England and Wales the Audit Commission estimated that just under 11% of hospitalised patients suffer an ADR, with mortality following ADR showing a marked upward trend. Yet the very real risks involved in taking medicine are either ignored or treated dismissively in the literature. For example, the WHO report stated, ‘Concerns about medication typically arise from beliefs about side effects and disruption of lifestyle and from more abstract worries about the long term effects and dependence. They are related to negative views about medications as a whole and suspicions that doctors over-prescribe medicines …’ (p. 44, italics added).

It is worrying that the WHO is so dismissive of people's concerns about medicines. The qualitative studies suggest that these concerns are real and sensible; they are not ‘beliefs’, ‘abstract worries’ or ‘suspicions’ and the statistics on ADRs bear testimony to this. Nevertheless, lay reports of the adverse effects of medicines are consistently dismissed, despite the fact that the current Yellow Card Scheme, which relies on doctors voluntarily reporting ADRs, has been estimated to capture only 1% of actual ADRs. Heath pointed out that more people are admitted to hospital for ADRs than for problems relating to ‘non-compliance’. ‘Medication errors’ caused by poor prescribing, poor dispensing and poor labelling are also a significant problem for patients and the NHS, with the potential to cause death, morbidity, increased or extended hospital admissions, litigation and financial burden to the NHS.

Concordance

Pollock and Grime, Dowell and Hudson and Svensson et al. suggested that doctors need to accept that patients modify their regimens and that the way forward is to encourage ‘authorised’ self-regulation. This is similar to Donovan and Blake's earlier call for ‘active and co-operative relationships between patients and doctors’ and to the newer concept of concordance, which ‘… describes the process whereby the patient and doctor reach an agreement on how a drug will

![Model of medicine-taking.](https://example.com/figure16.png)
be used, if at all.207 The aim of ‘concordance’ is to involve patients in making decisions about their medicines, to ensure that they have enough information for doing this and to support them with any problems that they might have. Thus, doctors might help patients with their testing and their modifications, providing feedback and guidance. ‘Crucially, concordance advocates a sharing of power in the professional–patient interaction.’

Concordance and related approaches would appear to make sense in cases where doctors can help patients determine appropriate individualised doses, and provide information on adverse effects and how to deal with them as well as the information necessary to conduct their evaluations safely and effectively. However, concordance depends on doctors sharing information honestly with patients. Unfortunately, Cox et al.209 have found that doctors emphasise the benefits of treatment rather than discussing its possible harms or risks, despite patients regarding these topics as essential. Equally, the concordance model depends on patients sharing information with doctors, but a few of the studies in the synthesis found that people were unlikely to raise issues with their doctors that they felt would be frowned upon.156,171,172 Furthermore, Rogers et al.156 noted that people with mental illness were very cautious about what they revealed to doctors because they were aware of the power that health professionals held over their lives. This was confirmed more recently by Britten et al.16 Socially excluded patient groups, such as those with poor mental health or HIV infection, may be less likely to achieve ‘concordance’ with their doctors. This could be because they are less normative in their attitudes or because they feel disempowered owing to their membership of marginalised groups. On the other hand, they might feel unable to reveal how they actually take their medicine; for example, people infected with HIV might feel unable to ‘admit’ modifying their regimen because of public health fears about viral mutation. This is not to suggest that health professionals should give up trying to acknowledge and deal with the concerns of marginalised groups, but to make the point that the power imbalance inherent in the doctor–patient relationship will not easily be resolved by the model of concordance.

It is has also been suggested that concordance is simply another way of encouraging people to take their medicine, except that, unlike the ‘compliance’ model, this time the coercion is concealed.206 This is an important consideration. Nevertheless, although doctors hold the balance of power during the consultation, the medical profession is oddly powerless once the patient has left the surgery. Whether or not there is coercion and whether this is concealed or explicit, lay people have always exercised their power to reject prescriptions or modify their regimens. Undoubtedly they will continue to exercise this power, as this is ‘normal’ behaviour for them. Unless health professionals learn to acknowledge their lack of power over how people take their medicines, ‘concordance’ is unlikely to be achieved.

**Relationship between health and medicine-taking**

Arguably, the ideologies of compliance, adherence and concordance, because they revolve around the axis of pharmaceutical medicine, distract attention away from non-pharmaceutical approaches to health. Heath206 made the point that the rhetoric of both concordance and compliance uncritically endorses medicine-taking. This relates to the most profound and hidden assumptions in the medicine-taking debate: firstly, that taking medicine is good for you and, secondly, that taking medicine is the only way of achieving desired health outcomes. In fairness, the original conceptualisation of concordance, as set out in the Royal Pharmaceutical Society of Great Britain report,210 acknowledged that taking medicine was not always the best thing to do and that ‘Almost all medicines have the capacity to harm as well as to do good.’ However, the WHO182 report made unsubstantiated claims about the health effects of not taking medicines as prescribed: ‘Poor adherence to long term therapies severely compromises the effectiveness of treatment making this a critical issue in population health …'183
Both the WHO\textsuperscript{183} report and the Medicines Partnership report\textsuperscript{184} assumed that taking medicines as prescribed would lead to good health outcomes. The introduction to the latter report, for example, stated, ‘In the context of a rising drugs bill and the key role of medicines in promoting health, the review underlines the vital importance of maintaining a clear focus on improving compliance and making better use of medicines’ (p. 2).\textsuperscript{184} Yet in the body of the report, the authors noted that the Cochrane Review of interventions to improve compliance (also published as a journal article)\textsuperscript{189} found little evidence that such interventions would lead to large improvements in treatment outcomes.\textsuperscript{189} Similarly, the review of the literature contained in the report indicated that the evidence relating ‘adherence’ to improved biomedical outcomes varies according to the illness.\textsuperscript{184} At the very least this suggests caution against generalising about the benefits of taking medicines, as clearly the evidence for the benefits of pharmacological treatments is stronger in some areas than in others.

Although the bulk of the medicine-taking literature uncritically assumes that pharmaceutical drugs and Western medicine are the only response to ill health, those people who reject their prescriptions outright clearly think otherwise. Presumably so too did the one in three Americans in 1990 who reported using at least one ‘unconventional therapy’ in the previous year, and who spent almost US$13.7B on alternative health treatments in the same year.\textsuperscript{211} It is estimated that approximately half the general population in developed countries uses complementary and alternative medicine.\textsuperscript{184} Harrison\textsuperscript{185} argued that user support for complementary and alternative medicines ‘…indicates in the clearest possible way that a significant number of people do not like conventional medicine or that it has failed them in particular ways’ (p. 36).

**Sociology, qualitative methodologies and the adoption of the medical agenda**

The sociological studies in this synthesis barely touched the assumptions underlying the compliance debate: firstly, that taking medicine leads to good health outcomes and, secondly, that it is the only possible response to ill health. As such, sociologists have to a large extent adopted the medical agenda. Only a few studies considered those who rejected medicine,\textsuperscript{172,173,175} despite the fact that it is this group that probably has the most to tell us about the reasons for the widespread reluctance to take medicine. Nevertheless, the early sociological and anthropological studies in this synthesis\textsuperscript{134,153,156,161,166,171} demonstrated that people have understandable and logical reasons for not taking their medicine as prescribed and challenged the assumptions inherent in the ideology of compliance. As such they undoubtedly influenced medical culture to the extent that, 10 years later, the term ‘compliance’ is no longer considered appropriate and there is a greater readiness to accept that some people will disregard their doctor’s advice and that they may be perfectly sensible for doing so.

There was a discernible tendency in some of the studies to regard qualitative methods as a means of obtaining information on the patient’s perspective that might ultimately be used to encourage people to take their medicine as prescribed. Broadly speaking, the sociological studies advocated the use of qualitative methods as a means of understanding why patients do not take their medicine as prescribed, whereas the medical and nursing studies advocated the methodology as a means of finding out why people did not comply. Angermeyer et al.\textsuperscript{157} put it bluntly, ‘Our findings point to the necessity for psychiatrists to explore patients’ subjective views and motivations with regard to their medicine in order to improve treatment adherence’.\textsuperscript{157} Given that qualitative methods are the preferred means of exploring ‘patients’ subjective views and motivations’ it is possible to argue that in some cases qualitative methods may have been used to advance the medical agenda.
Bringing medicine into the medicine-taking debate

In our view, one of the key conclusions produced by this synthesis is that the main reason why people do not take their medicines as prescribed is because of concerns about the medicines themselves. On the whole, the findings suggest that there is considerable reluctance to take medicine and a preference to take as little medicine as possible. This is in contrast to earlier theories that attributed 'non-compliance' to failings in patients, or current theories that attribute it to failures of the 'system' or the doctor–patient consultation. Although medicines are obviously at the centre of the medicine-taking debate and lay people have repeatedly demonstrated their wariness of medicines, their worries have tended to be marginalised or, as noted earlier, treated as 'beliefs about medicines', despite the well-documented existence of ADR and instances where concerns about the safety of medicines, for example over addiction to benzodiazepines, have been highlighted by patient groups based on their lay expertise, well before they were acknowledged by professionals (p. 145).²¹²

It is difficult to imagine how or why this most obvious factor has been overlooked, but it may be because of the dominance of the cultural belief in the benefit of pharmaceutical medicines. Or perhaps it was dismissed because of the view that patients, doctors and systems are easier to modify than medicines themselves. It may be because most of the research has ignored the lay perspective and has failed to involve consumers, resulting in a mismatch between the priorities of researchers and those of patients.²¹³,²¹⁴ However, sociologists are also responsible here because we have tended to focus on 'perceptions' of medicines or the 'meanings' people attach to medicines. This tendency has significant consequences. It makes the person, rather than the medicine, the focus of attention, in much the same way as the WHO¹⁸³ report (as noted in the introduction to this synthesis) considered anxieties about medicines to be 'patient-related' rather than 'therapy-related' factors. The result of this is that attempts are made to modify patient behaviour or the doctor–patient consultation rather than question the appropriateness of the medicine. Thus, because people's accounts are not taken at face value, the more mundane issues about the physical reality of medicines and the effects that they have on people's bodies and minds have become obscured, with the result that patients' priorities and concerns are neglected.

Resisting medicines

We feel that many lay people's response to medicine is best captured by the concept of resistance. In part this is because the term encapsulates the ways in which people take medicines at the same time as attempting to minimise their intake. However, the majority of people represented in this synthesis were not just unwilling or reluctant to take their medicines, nor simply cautious. The strategies they adopted to manage their medicine-taking indicate varying degrees of resistance to the prescription which they were given and, if the term resistance sounds strong, it should be remembered that the huge literature on 'non-compliance' exists only because so many people have continued to resist taking medicines in the face of sustained advice, interventions and admonitions. The term resistance also captures lay people's active engagement with their medicines, as well as the ingenuity and energy that they bring to dealing with their medicines. Additionally, it carries the suggestion of a clandestine operation, which it is, as most people do not tell their doctors about the modifications they make to their regimens.

The term resistance is usually employed in relation to the exercise of power or coercion. As such, it entails acknowledging that the traditional approach to medicine-taking has been, and in varying degrees continues to be, coercive. In this context it should be noted that the noun 'medicine' refers both to the pharmacological substance as well as to the profession that prescribes it. Therefore, we could also consider the possibility that any resistance is not only to the pharmacological substance, but also to Western biomedicine. From this perspective, the evaluation and modification of drug regimens that people engage in could also be understood as ways of reasserting control over their bodies and the decisions about it.
Policy and practice implications

There is a need to accept that people are unlikely to stop resisting their medicines. Doctors could assist people in their lay evaluations of medicines by providing the necessary information, feedback and support and by prescribing safely. However, doctors will need training and support to do this effectively. The policy emphasis needs to be less on attempting to modify people’s behaviour and more on developing safer medicines. This huge undertaking involves questioning the present methods used to develop and test medicines and ensuring that more attention is paid to safety at the licensing stage, possibly by having a probationary period. Additionally, safer ways need to be found of administering medicines, and of monitoring their effectiveness and acceptability to individual patients. Effective ways of identifying ADR in all patients need to be developed and implemented. Furthermore, in recognition of the fact that many people prefer not to take medicines, funds should be allocated, firstly, to determine what sort of treatments patients prefer and, secondly, to evaluate the safety, efficacy and cost-effectiveness of those preferred treatments.
Chapter 7

A synthesis of qualitative studies about lay experiences and perceptions of rheumatoid arthritis

Introduction

This chapter presents a synthesis of published reports of qualitative studies concerning lay perceptions and experiences related to the aetiology, treatment, management and lived experience of RA. RA was chosen as a topic because it was known to be a field where published qualitative studies had impacted widely on medical sociology.\textsuperscript{140,141} Furthermore, RA was selected as a narrow, disease-specific field in order to investigate what value qualitative synthesis might have for clinical practice for a particular condition.

RA, unlike the other main cause of arthritis, osteoarthritis, is not primarily a disease of ageing; the most common age of onset is between 30 and 50 years (www.arc.org.uk). RA is an inflammatory disease of the joints, which is experienced as a spectrum from minor twinges to destruction of joints, often with an uncertain pattern of flares and remissions. It is the potentially chronic and disabling nature of the condition and its uncertain course that has brought it to the attention of medical sociologists and others. RA affects women disproportionately (ratio 3:2 to men).

The central preoccupation of the studies, and their main theoretical significance, concerned the ‘experience’ of daily life with RA following the changes brought by the condition. Although issues of aetiology and treatment did feature, the papers that focused on these issues were marginal to the wider collective voice of the papers. Salient features of the included studies are detailed in Table 19. In expressing the synthesis here, the three areas of aetiology, experience and treatment are separated out, although far more attention is devoted to ‘experience’ as it was the focus of most of the studies.

Methodological features of included papers

As noted in the methods chapter, of the studies initially included, only four were characterised by both good reporting of the use of qualitative methods of data collection \textit{and} good analysis.\textsuperscript{138,177,229,233} One further paper contained good reporting of qualitative methods of data collection and analysis,\textsuperscript{223} but in this case the reviewers were concerned that the methods described were not actually reflected in the way in which data were analysed or presented in the paper. Studies by Bury,\textsuperscript{140,216} Williams,\textsuperscript{141} Williams and Wood\textsuperscript{218} and Pinder\textsuperscript{221} were conceptually very useful to the synthesis, but were characterised by poor reporting of methods of data collection and/or analysis. These papers described various forms of case study approaches that sought to build lines of sociological argument. It might not even be obvious to the casual reader that these studies employed interpretive or qualitative research methods, as this was not always stated explicitly.

The synthesis was principally constructed on the seven following studies. The studies by Bury,\textsuperscript{140,216} Williams\textsuperscript{141} and Williams and Wood\textsuperscript{218} stood out for their theoretical contribution.
alone, whereas that by Stephens and Yoshida stood out for good reporting of methods. The studies by Wiener, Shaul and McPherson et al. managed to combine conceptual coherence with good reporting and/or use of methods. The last two papers were both also grounded in the results of earlier ‘classic’ studies. The main value of Locker’s approach was that the book format allowed for a higher degree of ‘thick description’ and contextualisation of findings than that is found in the shorter journal articles.

The main contribution of three later studies to the synthesis was primarily as descriptive confirmation or elucidation of previous findings. The value of the studies reported by Williams and Barlow and Brown and Williams became apparent only during the reciprocal translation stage. This is as an important point because, following initial ‘assessment by checklist’, the papers might have been rejected on grounds of quality or relevance. The studies by Rao et al. and Archenholtz et al. did not look as if they had employed an interpretive approach, although their findings did not conflict with other papers in the synthesis and they were not

### TABLE 19 Salient features of 22 studies (reported in 25 papers) included in the synthesis following appraisal

<table>
<thead>
<tr>
<th>Source paper(s) (n = 25)</th>
<th>Country setting</th>
<th>Participants, male (female)</th>
<th>Sample origins</th>
<th>Type of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wiener, 1975</td>
<td>USA</td>
<td>21 (16)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Bury, 1982; Bury, 1988</td>
<td>UK</td>
<td>30 (25)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Locker, 1983</td>
<td>UK</td>
<td>24 (16)</td>
<td>‘Severely disabled by RA’</td>
<td>Interviews</td>
</tr>
<tr>
<td>Williams, 1984</td>
<td>UK</td>
<td>30 (19)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Donovan et al., 1989</td>
<td>UK</td>
<td>1989: 32 (ns)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Donovan, 1991</td>
<td>Denmark</td>
<td>8 + 3 sexual partners (ns)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Bjørner and Hansen, 1993</td>
<td>Sweden</td>
<td>9 women</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Rice and Young, 1994</td>
<td>USA</td>
<td>50 (38)</td>
<td>Senior activity centre</td>
<td>Semi-structured questionnaire</td>
</tr>
<tr>
<td>Pinder, 1995</td>
<td>UK</td>
<td>25 (18)</td>
<td>Snowballed, mainly voluntary group and hospital patients</td>
<td>Case studies developed from interviews</td>
</tr>
<tr>
<td>Brown and Williams, 1995</td>
<td>UK</td>
<td>7 women</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Shaul, 1995</td>
<td>USA</td>
<td>30 women</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Dildy, 1996</td>
<td>USA</td>
<td>14 (9)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Ryan, 1996</td>
<td>UK</td>
<td>7 (ns)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Moss, 1997</td>
<td>Canada</td>
<td>25 women</td>
<td>Hospital patients</td>
<td>Case studies developed from interviews</td>
</tr>
<tr>
<td>Rao et al., 1998</td>
<td>USA</td>
<td>33 (ns)</td>
<td>Hospital patients</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Williams and Barlow, 1998</td>
<td>UK</td>
<td>14 (9)</td>
<td>From arthritis charities/research centre</td>
<td>Interviews</td>
</tr>
<tr>
<td>Archenholtz et al., 1999</td>
<td>USA</td>
<td>50 women with SLE and 50 women with RA</td>
<td>Hospital patients</td>
<td>Structured telephone interviews</td>
</tr>
<tr>
<td>Stephens and Yoshida, 1999</td>
<td>Canada</td>
<td>46 (32)</td>
<td>Arthritis charity</td>
<td>Interviews</td>
</tr>
<tr>
<td>Lambert and Butlin, 2000</td>
<td>USA</td>
<td>12 (?)</td>
<td>Hospital patients</td>
<td>Focus groups</td>
</tr>
<tr>
<td>Gilworth et al., 2001</td>
<td>UK</td>
<td>47 (29)</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
<tr>
<td>Grant, 2001</td>
<td>UK</td>
<td>4 mothers</td>
<td>Hospital patients</td>
<td>Case studies based on observation of OT sessions, interviews, OT case note analysis</td>
</tr>
<tr>
<td>McPherson et al., 2001</td>
<td>New Zealand</td>
<td>10 women</td>
<td>Hospital patients</td>
<td>Interviews</td>
</tr>
</tbody>
</table>

*ns, not stated; OT, occupational therapy; SLE, systemic lupus erythematosus.*
excluded, although they could have been on these grounds. In addition, neither paper was focused on RA, the latter also including patients with systemic lupus erythematosus, although both clearly incorporated the views of patients with inflammatory arthropathy. Three papers were characterised by poor reporting and/or poor methods, and also added little conceptually to the synthesis, although again they were not excluded on these grounds. However, and in a similar manner to the Williams and Barlow paper mentioned above, the worth of the paper by Stenström et al. became apparent only during the reciprocal translation stage. The importance of the paper by Bjørner and Hansen was that its topic – sexuality – was ignored in most of the other papers.

Findings

Lay perceptions of the causes of arthritis

Three of the five factors mentioned by Rice and Young's respondents (damp and/or cold weather, ageing and injury) were also found in the two patient focus groups conducted by Lambert and Butin. Rice and Young's participants also mentioned heredity and diet. In the Lambert and Butin focus groups, diet was not mentioned as a cause of arthritis, but rather specific foods (red meat, coffee, sugar and junk food) were identified as exacerbating arthritis pain. Other causes mentioned only in the focus groups were viruses, bacteria, bursitis, close-fitting shoes and air pollution. However, in comparing the two studies, the main problem was that Rice and Young's participants included people who did not have arthritis and these responses were not separated out from those of the other respondents.

Rice and Young noted that lay accounts of the aetiology of arthritis fell broadly into two types of explanation: firstly, just those that suggested internal, 'mechanical breakdown' as responsible, where ageing and injury were identified as factors; and secondly, those that saw arthritis as resulting from an outside agent that infected the body. It was reported both by Shaul, and Brown and Williams that people had attributed initial symptoms to 'commonsense' explanations like minor trauma or overexertion, which also pointed towards 'mechanical' explanations.

For people with RA, the search for an understanding of disease aetiology became a more personal matter, as part of the broader 'attempt to establish points of reference between body, self and society, and to reconstruct a sense of order from the fragmentation produced by chronic illness'. In the perceived absence of biomedical explanations for the disease, 'the actual nature of the disease remains elusive'. Without adequate scientific–medical explanations for such devastating 'physical and social breakdown', people turned to lay explanations reflecting that 'the body is defined by its relationship to the world of social action not in isolation from it' (p. 182). In the account to follow, it will be shown that particular features external to the body become significant precisely because of characteristics within the individual, his or her place in society or his or her social milieu.

Living with the symptoms of rheumatoid arthritis

Pain dominated patients' accounts of the illness experience in RA to such an extent that one could wonder whether pain was the issue at the heart of the matter, rather than RA itself. Many studies contained detailed accounts of the nature of pain in arthritis, although this concept was noticeably absent in Bury's study. However, the abundance of work had fundamentally little to add to Wiener's study of the sociological aspects of pain management in chronic illness. Wiener was also the earliest published study considered in the synthesis. Whether or not later studies referenced Wiener's work, all those that contained findings about 'pain' demonstrated conceptual coherence with her framework and concepts.
Wiener noted four features of pain in RA. Firstly, there is an ‘absence of predictability’ to RA’s symptoms, including pain, which is characterised by ‘flare-ups’ that come without signs or warning. Secondly, there is a ‘variability of progression, severity and areas of involvement’.

In other words, people with RA do not know from one day to the next whether they will be feeling better, worse or the same as they do today, or whether or not the disease will affect the same body part(s) as last time. However, the one ‘certainty’ is that sooner or later there will be pain. Thirdly, Wiener also found that pain in RA had a ‘circuits’ relationship with fatigue, such that ‘pain drains energy and fatigue produces more pain’ (p. 98). In later studies, ‘fatigue’ was sometimes mentioned as a separate symptom in its own right, rather than a concept in relationship with pain. Fourthly, Wiener noted that these features of pain in RA led to a ‘reduction in personal resources’ for the individual.

Wiener’s findings led her to two key interpretations. Firstly, the key concept in living with arthritis relates to ‘tolerating variable uncertainty’. Secondly, the disease conditions engender a ‘dread of [future] dependency [on others]’. Wiener’s twin theories were confirmed and rediscovered by most of the later published studies, but no single study has captured these concepts with such clarity and groundedness. Wiener’s findings were consistently validated by the findings of the other studies included in the synthesis. Furthermore, her reporting of research methods and methods of data analysis compared favourably with that found in other included studies.

Other work added further findings to Wiener’s basic schema. Locker completed detailed case histories and interviewed people twice, 1 year apart. He noted a further feature of pain in RA: that it ‘is often not accompanied by any signs which would help to verify the sufferer’s complaints’ (p. 132). This is similar to Williams and Wood’s assertion that the symptoms of RA are ‘often vague’ although, for Brown and Williams, ‘vague’ was rather used to describe earlier symptoms, which were also characterised as ‘mild’ and ‘non-disabling’. Similarly, Pinder noted that both pain and fatigue provide considerable ambiguity, as they are not visible forms of disability.

Interestingly, apart from pain and fatigue, the only other typification of the condition of RA in the earlier literature reviewed was as a ‘crippling disease’ by Bury (p. 173), who also identified RA as ‘an outside force’ (see Lay perceptions of the causes of arthritis, above). The focus of earlier work was much more on coping strategies and the effects on social relations, which will be considered later. However, by 1989, Donovan et al., although still describing pain as the major symptom, also included ‘stiffness’ and ‘burning’ (p. 59). Bjørner and Hansen noted that flare-ups of arthritis were associated with reduced libido for some patients. Rice and Young’s folk model of arthritis identified ‘painful movement’ as a core concept, and also included ‘inflamed’ and ‘deformed’ joints as conditions, although their model was not necessarily applicable to specific types of arthritis.

From the mid-1990s, the practitioners of qualitative research into RA changed, and there was a discernible move towards descriptive studies highlighting the symptoms and conditions of the disease and their impact on physical and social functioning. This probably reflected the fact that these studies were being undertaken by people also involved in clinical practice, whereas earlier work had focused more on RA as an example of a painful, disrupting, disabling, chronic illness. Wiener was interested in the sociological aspects of pain management; Bury in the disruption to biography and relationships by chronic illness; Locker in RA as a form of social disadvantage; Williams in the application of the idea that people reconstruct their personal biographies in light of subsequent events; and Donovan in RA as a clinical encounter.

In later qualitative studies undertaken by nurses and others (although we should note that Wiener also came from a nursing background), there was a definite move towards a
'disease-specific' approach. At their best, these studies confirmed the findings of previous work while presenting results to clinical audiences. However, the reviewers characterised many of these studies as being of 'poor' methodological quality. One exception was the article by Shaul,176 which stood out as a far better study. Hers was the first paper to discuss areas of the body involved in RA symptomatology in any depth. For the women in Shaul's study, 'Most common symptoms involved the hands, wrists, shoulders and feet. In some cases pain and swelling occurred simultaneously, in others, pain occurred alone, followed by swelling. In either case, there was usually weakness in the affected limb and general fatigue' (p. 292).177 Shaul identified fatigue as the most 'pervasive' symptom, although both fatigue and pain were described as the most 'distressing' symptoms. These findings were further developed in a later study, where the root dissatisfaction with RA for female informants concerned the 'visible swollen and disfigured appearance of these painful body parts'.227 As one respondent to this study put it, 'The swellings are like a physical effect of pain. You don't want people to see the pain you are suffering' (p. 131).227 Although these findings seem partly to conflict with those of other studies which noted that the primary symptoms of RA are not obvious, they most probably reflect that in each case findings derived from people in different stages of the disease course (but see Discussion). The primary symptoms of arthritis lead to secondary symptoms, largely related to pain, and identified by: exhaustion, anger, depression, despair, self-pity and [perception of] loss of control.223 Bjørner and Hansen similarly identified 'low self confidence', reduced libido and difficulties reaching orgasm.

Consequences for identity and problems maintaining taken-for-granted activities

This section is largely about the problems of maintaining ‘taken-for-granted’ actions and behaviours and the concomitant challenges to norms of behaviour (as perceived by self or others). The best term to capture these issues in the context of RA is probably Bury’s140 ‘biographical disruption’. In a later reflection on these data, he noted: 'The onset and development of arthritis is simultaneously an assault on the body and a disruption of social life'.216 This ‘biographical disruption’ centred on the loss of previous life and work identities, or in the metaphor employed by McPherson et al.,233 loss of the individual’s ‘life definition’. Another metaphor related to these issues is Brown and Williams’222 ‘role incompetence’ (p. 699).

The experience of people with RA was mediated by certain sociological and physiological needs. The metaphors relating to these needs are detailed in Table 20 and are grouped according to the focus on symptoms, personal biography, presentation of self and social roles, and further categorised into intrinsic and interpersonal needs. The studies showed how RA brought previously taken-for-granted actions and behaviours into sharp focus. As Pinder221 put it, 'The conditions under which people take themselves for granted may be compromised' (p. 610). As important in this context were the conditions under which other people take the person with arthritis for granted. Some of these conditions were 'expectations governing behaviour in the wider culture'.216 As Williams and Wood179 noted, such social or cultural 'norms' may be unspoken, which seems to be a large part of the problem for the person with RA: 'Informal structures like families contain unstated norms of reciprocity or give-and-take, and for many people the experience of disablement is one of having these norms upset' (p. 130). It is evident that there are as many expectations governing physical functioning or the ability to do things as there are in relation to fulfilling social obligations. Table 21 shows that many of the factors causing disruption in the social lives of people are related to expectations governing social and cultural actions whether at home or in the workplace.

So far as 'the material world' and physical activities were concerned, the most detailed exposition of the effects of RA 'as a disablement' was that provided by Locker.139 Usefully, he followed Wiener,138 and noted that the uncertain course of symptoms in RA called for 'continuous monitoring and self-management since little can be taken for granted'.139 'Activity, however
TABLE 20 Needs of people with RA

<table>
<thead>
<tr>
<th>Types of need</th>
<th>Study</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intrinsic</strong></td>
<td></td>
</tr>
<tr>
<td>‘Relief from pain’</td>
<td></td>
</tr>
<tr>
<td>‘The feeling of being in control of the symptoms is vital’</td>
<td>Gilworth et al., 2001, p. 345235</td>
</tr>
<tr>
<td>Biographically focused (Bury, 1982)</td>
<td>Williams, 1984, p. 192241</td>
</tr>
<tr>
<td>‘To reaffirm telos and to reconstruct order in the presence of profound disruption in the biographical processes of daily life’</td>
<td>Williams, 1984, p. 192241</td>
</tr>
<tr>
<td>‘To prevent the onset of disability and deformity’</td>
<td>Donovan et al., 1989, p. 60217</td>
</tr>
<tr>
<td>‘Maintenance of one’s identity’</td>
<td>Stenström et al., 1993, p. 237228</td>
</tr>
<tr>
<td><strong>Attitude focused</strong></td>
<td></td>
</tr>
<tr>
<td>‘Learning to live with it’</td>
<td>Bury, 1982, p. 173240</td>
</tr>
<tr>
<td>‘Learned to live with pain’</td>
<td>Locker, 1983, p. 167239</td>
</tr>
<tr>
<td>Does not ‘give in’</td>
<td>Williams and Wood, 1988, p. 130217</td>
</tr>
<tr>
<td>‘Mental balance’</td>
<td>Stenström et al., 1993, p. 237228</td>
</tr>
<tr>
<td><strong>Interpersonal</strong></td>
<td></td>
</tr>
<tr>
<td>‘Presentation of self’ focused</td>
<td></td>
</tr>
<tr>
<td>Avoid embarrassment</td>
<td>Bury, 1982, p. 1752142</td>
</tr>
<tr>
<td>‘Avoiding unpleasant reaction’</td>
<td>Locker, 1983, p. 101216</td>
</tr>
<tr>
<td>‘You don’t want people to see the pain you are suffering’</td>
<td>Williams and Barlow, 1998, p. 131227</td>
</tr>
<tr>
<td>‘To present themselves as morally competent actors in the workplace’</td>
<td>Pinder, 1995, p. 624221</td>
</tr>
<tr>
<td>Self-sufficiency</td>
<td>Stephens and Yoshida, 1999, p. 232229</td>
</tr>
<tr>
<td>‘Reciprocate for favours or help’</td>
<td>Stenström et al., 1993, p. 237228</td>
</tr>
<tr>
<td>‘Personal integrity’</td>
<td>Moss, 1997, p. 27224</td>
</tr>
<tr>
<td>‘Overcoming isolation and maintaining contact with family, friends and neighbours’</td>
<td>Stephens and Yoshida, 1999, p. 232229</td>
</tr>
<tr>
<td>‘Trying to retain as much autonomy as possible’</td>
<td></td>
</tr>
</tbody>
</table>

minimal, requires physical and psychological effort’ (p. 19). The ‘erstwhile, taken-for-granted world of everyday life’ becomes ‘fraught with danger’ and ‘everyday objects and events take on an alarming character and may give rise to feelings of insecurity’. One possible response to such dangers was to never leave the house, although such a response may leave people feeling that they were ‘locked-in’.

**Responses and coping strategies**

The studies showed that the main factors that affect people’s experiences of RA are the symptoms of the disease, the nature of the cultural backdrop, and expectations governing norms of social roles and behaviour. Certain consequences derive from these factors, which affect the ways in which people can or cannot cope with the disease. The psychologically oriented data also pointed towards a ‘career’ model of adaptation to RA, although an attempt to synthesise the studies along these lines was unsuccessful. However, the attempt yielded insights into both the nature of the studies themselves and the experience of adapting to life with RA. As with psychological coping strategies, it seemed as though the idea of an illness ‘career’ probably reflected more the professional and academic standpoint of the authors of the studies than it did the data.
gathered during the interviews. That is, concepts such as ‘mastery’\textsuperscript{177} or ‘taking charge’\textsuperscript{230} were more fundamentally associated with clinical imperatives surrounding treatment, cure and rehabilitation than they were patient perceptions. (Both reviewers felt that Shaul’s\textsuperscript{177} concept of ‘mastery’ of disease was not clearly evidenced by the findings presented in her paper.)

A further problem derived from the fact that many studies were cross-sectional in nature and were focused on people with more problematic or advanced disease, problems or disability. However, the failed attempt to synthesise the studies around the theme of illness career may say something about the nature of RA itself. Thus, it may be that the pervasive uncertainty and unpredictability of RA means that some people never manage to ‘master’ or ‘take charge’ in RA and, in effect, the disease ‘wins’. Alternatively, people may think that they will one day ‘master’ the disease, but repeated failed attempts lead to a sort of progressive disillusionment. So long as studies remain focused on captive clinical populations, we may never find answers to these questions, as the views of people who can cope, whose flare-ups are not so bad, or for whom the disease is ‘in remission’ were not included in the available research, which was more fundamentally concerned with describing the lives of chronically disabled people trying to cope with severe and debilitating disease.

<table>
<thead>
<tr>
<th>Type of Disruption</th>
<th>Studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Changed appearance</td>
<td>‘Visible changes … made them less attractive to their partners and potential partners’ (Williams and Barlow, 1998, p. 134)\textsuperscript{227}</td>
</tr>
<tr>
<td></td>
<td>‘People in their social worlds would see only the disease rather than the person within’ (Williams and Barlow, 1998, p. 131)\textsuperscript{237}</td>
</tr>
<tr>
<td></td>
<td>‘Difficulties finding a sexual partner (Bjørner and Hansen, 1993)\textsuperscript{219}’</td>
</tr>
<tr>
<td>Changed abilities</td>
<td>‘Reduction of personal resources’ (Wiener, 1975, p. 98)\textsuperscript{138} ‘Warrantability of a person’s changed behaviour (Bury, 1982, p. 180)\textsuperscript{142}’</td>
</tr>
<tr>
<td></td>
<td>‘Chronic illness undermines … autonomy and routine’ (Williams and Wood, 1988, pp. 130–1)\textsuperscript{119} ‘Their bodies appeared to have become detached, creating a hiatus between their wishes and their actions’ (Williams and Wood, 1988, p. 129)\textsuperscript{178}</td>
</tr>
<tr>
<td></td>
<td>‘Influences physical capacity’ (Stenström \textit{et al.}, 1993, p. 240)\textsuperscript{176} ‘Changes in the way work is tackled (Bjørner and Hansen, 1993)\textsuperscript{121}’</td>
</tr>
<tr>
<td></td>
<td>‘Inhibits sexual activity, makes sex painful (Lambert and Butin, 2000)\textsuperscript{232} Bjørner and Hansen, 1993\textsuperscript{219}’</td>
</tr>
<tr>
<td></td>
<td>‘Arthritis makes everyday life difficult (Lambert and Butin, 2000)\textsuperscript{178}’</td>
</tr>
<tr>
<td></td>
<td>‘Implications for current life definition and self-perception relative to life without disability’ (McPherson \textit{et al.}, 2001, p. 710)\textsuperscript{233}</td>
</tr>
<tr>
<td>Interpersonal relationships</td>
<td>Dread of dependency (burden on others) (Wiener, 1975, key concept; Donovan \textit{et al.}, 1989; Brown and Williams, 1995, p. 699)\textsuperscript{221}</td>
</tr>
<tr>
<td></td>
<td>‘Loss of friends through difficulties keeping plans and having to cancel at the last minute (Williams and Wood, 1988)\textsuperscript{18}’</td>
</tr>
<tr>
<td></td>
<td>‘Changes in roles/work-sharing (Bjørner and Hansen, 1993)\textsuperscript{219}’</td>
</tr>
<tr>
<td></td>
<td>‘The partner needs to be trained in how to do new things (Bjørner and Hansen, 1993)\textsuperscript{179}’ ‘Makes social relationships more complicated’ (Stenström \textit{et al.}, 1993, p. 240)\textsuperscript{176}</td>
</tr>
<tr>
<td>Cultural impediments</td>
<td>‘The illness is in many ways incompatible with the organisation of work’ (Locker, 1983, pp. 98–9)\textsuperscript{119} ‘Difficulties in maintaining the reciprocal nature of family relationships (Williams and Wood, 1988)\textsuperscript{178}’</td>
</tr>
<tr>
<td></td>
<td>‘The burden they felt their physical care caused for others’ (Brown and Williams, 1995, p. 699)\textsuperscript{222} ‘The way one is viewed by strangers (Moss, 1997, p. 30)\textsuperscript{122}’</td>
</tr>
<tr>
<td></td>
<td>‘Responsibility for domestic chores still resides with the woman (Bjørner and Hansen, 1993)\textsuperscript{219}’ ‘The wider social and cultural significance of illness in the workplace’ (Pinder, 1995, p. 605)\textsuperscript{233} ‘Society at large is not organised to help disabled people’ (Archenholtz \textit{et al.}, 1999)\textsuperscript{228}</td>
</tr>
<tr>
<td>Changing social status</td>
<td>‘Women frequently complained of their diminished roles as homemaker’ (Wiener, 1975, p. 102)\textsuperscript{138} ‘A deep sense of loss’ because ‘chronic illness caused them to fail in a valued role’ (Locker, 1983, p. 84)\textsuperscript{119} ‘Coming to terms with their own embarrassment about being “different”, as well as the discomfort it aroused in others’ (Williams and Wood, 1988, p. 131)\textsuperscript{178} ‘Feeling alienated from friends, family and co-workers’ (Shaul, 1995, p. 293)\textsuperscript{157}</td>
</tr>
<tr>
<td></td>
<td>‘Partners of women “taking on aspects of what had hitherto been women’s domains” (Brown and Williams, 1995, p. 699)\textsuperscript{220} ‘Illness carries … disturbing connotations relating to contagion, pollution and taboo’ (Pinder, 1995, p. 624)\textsuperscript{223} ‘Bodies which are flawed or conspicuously ill convey powerful symbolic messages concerning social order and disorder’ (Pinder, 1995, p. 624)\textsuperscript{234} ‘The presence of “visible disease” was invasive, threatening their private and public selves’ (Williams and Barlow, 1998, p. 131)\textsuperscript{227}</td>
</tr>
</tbody>
</table>
### TABLE 22a  General coping strategies in RA

<table>
<thead>
<tr>
<th>General coping strategies</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>‘Pacing/keeping up with normal activity’ (Wiener, 1975, p. 100)</td>
<td></td>
</tr>
<tr>
<td>‘Lowering expectations’ (Wiener, 1975, p. 101)</td>
<td></td>
</tr>
<tr>
<td>‘Developing a new set of norms for action’ (Wiener, 1975, p. 101)</td>
<td></td>
</tr>
<tr>
<td>‘Eliciting help’ (Wiener, 1975, p. 101; Donovan et al., 1989, p. 60; the last resort)</td>
<td>(decreases the potential of using other strategies)</td>
</tr>
<tr>
<td>‘Thinking it out’ (plan or avoid activities) (Locker, 1983, p. 71)</td>
<td></td>
</tr>
<tr>
<td>‘Individuals, families and others frequently test the limits of their relationships, particularly the amount of support and care they are able or willing to afford each other’ (Bury, 1988, p. 113)</td>
<td></td>
</tr>
<tr>
<td>‘Changing the way they did things, consciously or unconsciously’ (Donovan et al., 1989, p. 60; also ‘modify tasks’ in Stephens and Yoshida, 1999, p. 239; ‘Changing or adapting their jobs’ in Gilworth et al., 2001, p. 344)</td>
<td></td>
</tr>
<tr>
<td>Some women react to renewed illness by isolating themselves (Bjørner and Hansen, 1993; also ‘self-imposed isolation’ in Dildy, 1996, p. 180)</td>
<td></td>
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<tr>
<td>‘[They] became experts at recognizing the cues that signal the onset of a flare, of overwork or the need to change a pattern of activity through the acquisition of knowledge about the disease, its treatment, and experience of living with it’ (Shaul, 1995, p. 295)</td>
<td></td>
</tr>
<tr>
<td>‘Not use an assistive device; or be selective in terms of where help or an aid is used’ (Stephens and Yoshida, 1999, p. 239)</td>
<td></td>
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</table>

### TABLE 22b  Specific coping strategies in RA

<table>
<thead>
<tr>
<th>Intrinsic</th>
<th>Interpersonal</th>
<th>Public/societal</th>
<th>Environmental</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use alternative remedies (Donovan et al., 1989)</td>
<td>‘Concealing disability and/or pain’ (Wiener, 1975, p. 99)</td>
<td>Concealment, grooming behaviours (Williams and Barlow, 1998)</td>
<td>‘Modifying the dwelling structurally, using assistive devices, arranging physical space according to ability and adjusting the body to the environment’ (Moss, 1997, p. 27)</td>
</tr>
<tr>
<td>Take multiple medications (Rao et al., 1998, p. 255)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>‘For Marilyn, the mystification underlying the disease can be gotten around through shifts and changes in the physical and social aspects of her home environment’ (Moss, 1997, p. 28)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>‘Pay for help rather than receive it informally’ (Stephens and Yoshida, 1999, p. 239)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>‘[Florence] continues to neglect her health in the face of maintaining social interaction’ (Moss, 1997, p. 29)</td>
<td>‘Planning and routing not only reduce the physical difficulty of moving around the community, they do so in a manner which helps to preserve the person’s public identity’ (Locker, 1983, p. 87)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men felt through regular exercise ‘they were able to play an active role in controlling their disease’ (Williams and Barlow, 1998, p. 136)</td>
<td>Flexibility in the couple’s roles, so that the partner can step in and take responsibility for tasks when necessary (Bjørner and Hansen, 1993)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>‘Self-determination and agency’; ‘Struggle to do difficult tasks by themselves’ (Stephens and Yoshida, 1999, p. 239)</td>
<td>‘Refuse offers of assistance’; ‘Avoid people who give unwanted help’ (Stephens and Yoshida, 1999, p. 239)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Tables 22a and b outline the main coping strategies described in the studies. Those described in Table 22a seem to constitute ‘grand’ strategies related to activity in general. Those strategies presented in Table 22b are specific to different arenas: intrinsic strategies, interpersonal strategies, public strategies, and strategies relating to the environment.*

**Perceptions and experiences of treatment**

Surprisingly (given that the interview samples were derived largely from people in clinical settings), perceptions and experiences of care were not a major feature of these studies. An exception was Bury, who noted that as medical intervention appeared ‘both important and
limited’ (p. 173), medical knowledge in the context of RA was both a resource and a constraint. In a similar manner, Locker's finding that because all treatments involved risks in terms of side effects, ‘the disadvantages may outweigh the advantages to be gained’ (p. 62). Within a focus on individuals’ needs to ‘identify the causes of arthritis’ (p. 191), Williams noted, ‘the limitations of medical science in delivering a satisfactory explanation’ (p. 175) and also asserted within this context that ‘medicine can support political bureaucracy in preventing the establishment of social justice’ (p. 185).

Similarly, Brown and Williams reported that some of the seven women nursed by the interviewer expressed frustration at the perceived ‘medical impotence’ (p. 698). However, they were also relieved at being referred to a specialist and receiving ‘an acceptable name or diagnosis for their symptoms’ (p. 698). Similar themes were evident in Shaul’s: ‘In many cases, the first experiences of medical management were not helpful, as many were misdiagnosed or found that their complaints were not taken seriously’. Ryan reported complaints about ‘too much emphasis placed on the physical manifestations of the condition’ (p. 48), and not always being reviewed by the doctor, but these issues did not seem specific to patients with RA. Similarly, Archenholtz et al. mentioned ‘lack of continuity of care, not always the same doctor, and difficulties making appointments’.

The most detailed consideration of the issues was that provided by Rao et al.’s focus group study, which usefully compared RA patients with people with osteoarthritis. As in Lambert and Butin’s patient focus groups, ‘[some] believed that doctors could do very little to relieve pain or cure the underlying disease’ because ‘arthritis has no known cause or cure’. Interestingly, when compared with the osteoarthritis patients, RA patients more frequently mentioned ‘their trust in the physician’s ability to treat their arthritis’ (p. 259). Other findings were that medications were only effective for short periods of time (p. 255) and that ‘Several patients wished that a single medicine instead of multiple medications could be used to treat their arthritis’ (pp. 255–6).

Gender

Given that RA disproportionately affects women, it was perhaps unsurprising that gender roles were particularly emphasised in these studies. However, although some of the papers seemed to have chosen RA as a focus for study precisely because of the likely gender issues, the conceptual development in this field is weak and generally not related either to the existing studies in RA or to wider feminist or gender theory.

A generic weakness across the studies seemed to be the attempt to explain the experiences of women by analysing their accounts in isolation from those of men. A notable exception here was the study by Björner and Hansen. It was particularly disappointing that the conceptually rich account provided by Pinder seemingly excluded men at the analysis stage. We suspect that the reason for excluding male accounts was ostensibly to ‘strengthen’ the focus of the accounts on the experiences of women, but it could be argued that this approach actually had the opposite effect.

Once again, the main issues were covered in Wiener’s ground-breaking study and related to the fact that, within the traditional domestic division of labour, men and children have a stake in women remaining active. Where men take on what were once women’s roles, ‘role reversal may result in a permanent change in the household’s division of labour’ (p. 102). Ultimately, the relevant issues did not seem to point towards gender per se, but rather towards the more general loss of the ability to engage in activities once seen as mundane. Whether for worker or homemaker, it was precisely these taken-for-granted activities which constituted a large measure of self-identity. Some of the women interviewed in the studies complained that the support they were receiving from partners was ‘too much’, as such support was felt to be eroding.
Reading across the studies, however, it was evident that similar issues also applied to men:

I’m no longer the man I used to be. I am no longer the husband, the lover, you know, all of the aspects of my life, they’re no longer there.

The one attempt to seemingly systematically compare accounts from women with those from men was found in the study by Locker who, in contrast to the other studies already cited, noted that ‘The women experienced job loss as acutely as the men and few were content to settle into the role of housewife.’ Thus, we are left wondering whether or not the sorts of gender issues highlighted in the studies were particular to women, or whether or not they were made particular in the way that data were collected and analysed.

Interpersonal gender dynamics received more attention in the two Scandinavian studies, and the main findings seemed to further underline Bury’s assertion that much of the success in coping with RA seems to hinge upon the degree of flexibility possible in both formal institutions and informal relationships:

In the beginning Soren was going to help me with the cooking, but that was no good. I wanted everything done my way. Now we each do our own things, but duties are not fixed. Soren just does the things he knows I can’t do, and when I am feeling rough he just takes over. It isn’t something we talk about much.

**Discussion**

The main findings of the studies concerning coping strategies in RA, and the experience of living with the illness were probably best represented in the earlier ‘classic’ studies. Although the synthesis has not added much conceptually in relation to these issues, one of the main reasons for this was that many later studies were confirmatory rather than offering conceptual development or innovation. They did not refine or develop the concepts of earlier studies, in relation to either gender or the different ‘stages’ of the disease. Most of the studies were cross-sectional in nature. Although some authors (e.g. Shaul, Bury and especially Locker) did follow people over time, these respondents were seemingly already at the ‘chronic’ end of the spectrum to start with.

As we noted at the outset of this chapter, some of the earlier classic studies included in this synthesis were characterised by poor reporting of methods of data collection and/or analysis. Assuming the integrity of past classics, as we did, where transparency about the methods used is absent or poor is questionable. However, we feel that our decision was vindicated in this case because the later papers, for which the reporting of methods was much better, and in which there seemed to be no awareness of the earlier classic papers, nevertheless, produced very similar findings.

Our synthesis also raises all sorts of questions about the nature and value of qualitative research. Given the repetitive nature of the findings of the studies, and their large number, we would call into question the necessity of further work in this field, particularly work which fails to build on or develop what is already known. Although there was a conceptual coherence among these studies that rendered their synthesis easier, we found evidence that later studies had stopped referencing earlier work (Table 23). For example, Wiener was last referenced in Shaul’s paper and now seems to have been consigned to history, even though the synthesis has found that the findings of her study are still being rediscovered by more recent researchers. This could reflect a tendency in science and medicine to search only for more recent work, an implicit
assumption of this approach being that previous authors will have already read and incorporated the salient aspects of relevant work. The implication for qualitative researchers is that they need to search back further for relevant studies before repeating work that has already been done. The implication for synthesists of qualitative studies is that they will need to have an open-ended attitude (date wise) when searching for relevant studies.

We are likely to learn more about chronic illness through different forms of cross-cultural analysis, a point made by Wiener in 1975. She suggested that her theoretical framework could have been expanded by ‘sampling under different conditions, for example in another culture’ (p. 102). Unfortunately, later qualitative researchers chose to ignore her advice and instead reinvented the theoretical wheel of experience and adaptation in RA in clinically captive Western populations. This synthesis underlined the value of Wiener’s paper, which to this day seems the best at capturing the essence of lay experiences of RA. However, this point emerged only when her work was systematically compared and synthesised with subsequent studies.

This synthesis raises questions about the usefulness of attempting to synthesise a body of qualitative research within a very narrowly defined clinical area. With hindsight, it may have been more fruitful to synthesise studies within a broader field, say ‘chronic illness’, which seemed to be the topic that interested some of the study authors in the first place. Nevertheless, it may be that our findings could be presented to clinicians and patients in a way that would be helpful in order to highlight the patient’s perspective or provide information relevant for self-help or professional care.
Following our experiences in this and other synthesis projects, and discussions with others, we feel that there will be much to gain from future synthesists searching for the ‘absences’ and the ‘audible silences’\textsuperscript{15} in reports of qualitative studies. This synthesis is based on the substance of what was present in the findings of the included papers, but we could have learned more by focusing on what was missing. For example, it is noteworthy that patient experiences of health care did not feature much, even though most interviewees were derived from patient settings. Further, despite statements in some of the papers to the effect that the insights of qualitative studies can inform and improve patient care, specific implications or recommendations for care or treatment were also largely notable by their absence. Although some of the practitioners’ papers were indicative of studies undertaken by researchers at the beginning of their research careers, or primarily for professional development purposes, others\textsuperscript{179,233} constituted substantive studies. However, even these studies failed to draw out implications for treatment and care.

With more absences in mind, and notwithstanding our earlier comments about the usefulness of further research, our synthesis points to those areas in which further work might be done. We have already pointed out a need for cross-cultural studies and more detailed work around gender. Perhaps, more fundamentally, there is a need to capture the views of people with RA who inhabit the world outside the treatment system, or those who can cope, perhaps with milder disease. At present, our understanding of lay ‘coping strategies’ is largely skewed towards samples of people who, by definition, cannot cope. Essential concepts derived from the synthesis point towards a narrative or biographical approach, but this needs to somehow extend into the time horizon preceding disruption or reconstruction.
Chapter 8

Discussion

The value of syntheses

At the outset of this evaluation, we noted a number of objectives that might be attained through qualitative syntheses. In particular, it was suggested that they could bring together isolated qualitative studies and provide an accumulated understanding on a particular theme; that they could go beyond the detailed summarising of the traditional narrative review by achieving fresh insights, conceptual development and theory building; and that they could complement quantitative research by providing a more comprehensive understanding of a variety of phenomena. A primary purpose of this evaluation was to complete two full-scale qualitative research syntheses using meta-ethnography. We succeeded in this by producing a detailed synthesis of qualitative studies of medicine-taking, and a synthesis of qualitative studies of people’s experiences of living with RA. In doing so, we have demonstrated that these objectives are achievable.

The synthesis into medicine-taking has produced major new insights. It has shown that the reason why patients do not take medicines as prescribed is not because of failures in the doctor–patient consultation, as volumes of previous work have suggested, but because people have real concerns about the medicines themselves. These concerns had been voiced in qualitative studies, but it was only when brought together through the painstaking process of reciprocal translation that the significance of this evidence became apparent. This synthesis, however, went beyond simply drawing attention to these findings. It also produced an explanation for why people’s concerns about medicines were overlooked for so long and went on to propose resistance to medicine as a concept which best encapsulates lay responses to medicine. Another product of this synthesis was the model of medicine-taking which describes the various pathways which people might follow after a medicine has been prescribed for them.

On the other hand, it is possible that syntheses will not always reveal new insights. Although the RA synthesis shed new light on the range of ways in which the social relationships of those with RA were disrupted (see Table 21) and the general and specific coping strategies that those with RA employed (see Table 22), there was little to add to the conceptually rich earlier ‘classic’ studies. Thus, this synthesis did not produce significant new insights, but it did serve an important function in demonstrating what cumulative knowledge there is in this area.

Both of the meta-ethnographies conducted demonstrate the value of qualitative synthesis in establishing what is known and what remains unknown or hidden about a topic at a given point in time. They also illustrate how it is possible to trace conceptual development, or the lack of it, within a field. Within the field of medicine-taking, many of the earlier sociological studies focused on challenging the concept of compliance, rather than developing new theory, whereas the later, practice-based literature has been mainly descriptive. Similarly, in the earlier studies in the RA synthesis, there was evidence of considerable conceptual development around both patients’ experiences of RA specifically and of chronic illness more generally, but little by way of theoretical advance detectable in the later papers.
Additionally, our work illustrates that qualitative syntheses can highlight the various positions adopted by authors and, in the case of the medicine-taking synthesis, the involvement of sociologists with the medical agenda became clear only as the papers were synthesised. Perhaps most importantly, syntheses can provide a *weight of evidence*. In the medicine-taking synthesis, for example, only one of the studies \(^{173}\) dealt explicitly with the lay evaluation of medicines in any detail, yet when synthesised, the studies revealed many data on this topic, substantially more than any individual study produced alone. The synthesis could not complete the detail for some aspects of the lay evaluation of medicines but has revealed the practice to be widespread. This synthesis also produced a weight of evidence about people's concerns with the safety of their medicine that has not been found elsewhere. Single studies reporting worries about medicines are not taken to be sufficient evidence, yet when studies are brought together the weight of evidence is hard to ignore.

One of the great values of a qualitative synthesis, it seems to us, lies in the systematic and comprehensive approach that it brings to reviewing the literature. Both our syntheses revealed that only a minority of the studies referenced each other. This was the case in the medicine-taking synthesis even when papers were about the same medicines. We found a similar picture in our earlier pilot synthesis on patient experiences of diabetes.\(^{17}\) This suggests that qualitative health research is not an evolving process whereby new studies build on earlier ones and research is conducted only after the relevant literature has been reviewed and important questions identified. Rather, among the later studies in each of the syntheses, there appeared to be little regard for earlier relevant studies and, certainly within the medicines for treating HIV infection and experience of RA literature, a tendency for studies to replicate each other. Another important aspect of a qualitative synthesis may therefore be to indicate when it is time to draw a line and strongly discourage further research on a particular topic. In making this judgement, it may be helpful to apply the notion of saturation, rather as one would in primary qualitative research. In the RA synthesis, it was clear that in more recent studies there was no new conceptual or theoretical development taking place, suggesting that 'saturation point' had been reached, at least as far as studies of the experiences of clinical groups of people with RA were concerned. While wishing to prevent 'me too' qualitative studies in this way, it is important to recognise the possibility that further research could lead to new directions and insights, if it was, for example, informed by a new theoretical approach.

The fact that we have found, in meta-ethnographies that we have conducted on three different topics, that only a minority of studies reference earlier work is not just an argument for the value of qualitative synthesis, but also a source of concern about the research practice of primary qualitative researchers. This finding indicates that those of us providing training in qualitative research methods evidently need to do more to emphasise the importance of a thorough literature review as a prelude to any qualitative research.

In the same way that qualitative syntheses could be useful in revealing where there has been sufficient research, they may also help to indicate where further research effort needs to be focused. Our experience of conducting the RA synthesis in particular taught us that it was important to pay attention to absences or, as Noblit and Hare\(^{15}\) termed them, 'audible silences'. In the case of RA, it became clear that most of the qualitative studies had based their research on clinical groups of patients. Consequently, the strong focus in the research on the immense disruption that RA wrought on people's lives may have been a reflection of this. The voices of those with RA who were not receiving clinical care for their RA, and for whom RA may have represented less of a disruptive and disabling experience, were not present. Syntheses are also able to reveal aspects of a body of literature that are disguised; the medicine-taking synthesis unveiled people's real concerns about the safety of medicines which had in effect been rendered absent by their labelling as lay beliefs.
Meta-ethnography as a method of qualitative synthesis

Stages in a meta-ethnography

The studies in both syntheses were concerned with two specific topics but were conducted in several different countries, with disparate samples and methods and for varying readerships. In the case of the medicine-taking synthesis, they also involved a diverse group of medicines. Nevertheless, it was possible, using meta-ethnography, to systematically draw these studies together to present something new. Having completed three syntheses, it is our considered view that reciprocal translation is at the heart of why meta-ethnography is an effective method of qualitative synthesis. The requirement of systematically relating findings from different studies to each other makes it possible to establish new relationships between concepts, such that it is possible to conceptualise an issue in a coherent but fresh way. Following the reciprocal translation stage, the necessary reordering, relinking and reanalysis of material that a synthesis entails represents in itself new material or new findings and may in this case correspond to what Noblit and Hare\textsuperscript{15} called a ‘lines-of-argument’ synthesis. In the medicine-taking synthesis, the line of argument was both written and summarised and depicted diagrammatically in the model of medicine-taking (see Figure 16). However, further concepts may advance from this level, what Britten \textit{et al.}\textsuperscript{16} called ‘third-order’ concepts. In the medicine-taking synthesis, there were two such concepts: the notion that the medicines themselves needed to be brought back into the debate about medicine-taking and that the way in which many lay people respond to medicines is best characterised by the concept of resistance. In the RA synthesis, the lines-of-argument synthesis was presented in both a written and a tabular form.

Noblit and Hare\textsuperscript{15} suggested that there were three possible types of synthesis: reciprocal translation, refutational and lines-of-argument. As the preceding paragraph suggests, our experience is that there are three stages of qualitative synthesis in a meta-ethnography with reciprocal translation the first crucial stage. It requires the synthesiser to consider all the concepts and findings in each study in relation to all other studies. One of the concerns about qualitative synthesis is its potential to destroy the integrity of the initial studies. We consider that reciprocal translation protects against this because the method requires taking account of all the findings and concepts in each study and assiduously considering what each means. The next stage involves a process of reordering, relinking and reanalysis, finally leading to a representation of the synthesised material as line of argument. Beyond this there may be a further level of abstraction and conceptual development.

Identification of studies to be included

A key requirement of quantitative syntheses is to achieve an exhaustive search of the literature. Other research groups, using modified versions of meta-ethnography, have conducted extensive literature searches on broad topics. For example, Paterson\textsuperscript{80} followed the procedures of quantitative synthesis and retained all studies that satisfied the appraisal criteria, resulting in the meta-synthesis of 292 studies of chronic illness. Noblit and Hare\textsuperscript{15} regarded wide-ranging literature search as useful but not a requirement, depending on purpose of the meta-ethnography. They emphasised that the selection of studies should be made on the grounds of relevance to the topic and should include a range of studies likely to enable new insights. They warned that ‘…generalizing from all studies of a particular setting yields trite conclusions’ (p. 28).\textsuperscript{15} We searched only for papers within a 10-year time frame for both the RA and medicine-taking syntheses, although in the case of the former we subsequently included a purposive sample of earlier ‘classic’ papers. In addition, the medicine-taking synthesis included only English-language papers, whereas in the RA synthesis non-English-language papers were considered, and one such included paper required translation, thus increasing the resources and time required.
The reciprocal translation stage is very time-consuming. It and the subsequent stages of synthesis also require the synthesiser to be immersed in all the studies. This places practical limits on the number of papers that can be synthesised. In the medicine-taking synthesis, it was necessary to begin by grouping the papers according to the type of medication as a way of making the task more manageable. Even so, it was difficult to conceive of synthesising many more than the 38 papers included in that synthesis without there being a trade-off between the breadth of papers included and the depth of the scrutiny and thought invested in the various stages of synthesis.

The practicalities of handling large numbers of studies may result in using processes of data selection, management and computing that lead to context stripping, and a less detailed immersion of researchers in the studies to be synthesised. Specialist computer packages are being developed to assist the synthesis of qualitative research, and these are likely to encourage the synthesis of greater numbers of studies, but this approach will potentially change the nature of the meta-ethnographic synthesis towards a more procedural and less interpretive approach.

Noblit and Hare\textsuperscript{15} were mainly concerned with identifying the relevance of studies and their potential contribution to the question of interest. As we noted in \textit{Chapter 4}, most syntheses employing meta-ethnography, or methods of synthesis derived from meta-ethnography, have followed Noblit and Hare's\textsuperscript{15} approach and not undertaken a formal appraisal of quality. From our experience, an important benefit of the use of formal appraisal criteria was to encourage the close reading of studies as an aid to interpretation. We also found it helpful to be able to discuss papers within the research team. We did exclude papers on the grounds of relevance, or because they did not conform to our definition of qualitative research as involving both qualitative methods of data collection and data analysis. This latter criterion ensured that we excluded qualitative work that was fatally flawed. Yet, in spite of our very detailed appraisal of studies, we did not exclude any studies on quality grounds. Some studies made an important contribution to the synthesis because of their conceptual strength, although their methods, as formally assessed, were weaker (or inadequately described), whereas other studies largely fulfilled the quality checklist in terms of methods but made little conceptual contribution to the synthesis.

We regard the quality of qualitative research as a very important issue. However, we look on the appraisal of qualitative research as more of a concern for the funders and publishers of qualitative research than for the qualitative synthesiser. This is because we have found that weak studies either will not synthesise or will contribute only minimally to a synthesis, adding weight to the better studies. Although there is an argument that including weak studies gives them an unwarranted credibility, such studies do not unduly distort a qualitative synthesis in the way that a poor-quality, highly biased quantitative study could influence a meta-analysis. In a qualitative synthesis, it is the power of ideas that matters.

\textbf{Reproducibility and validity}

The commissioning brief for this project was for methodological work 'to enable users of qualitative work to both appraise and synthesise qualitative studies in a rigorous, replicable and formalised way'.\textsuperscript{10} Meta-ethnography, as described by Noblit and Hare,\textsuperscript{15} is a rigorous approach to synthesis but it is also an interpretive approach. Consequently, a high level of reproducibility would not be expected. Indeed, the authors acknowledged this and did not view it as problematic, rather the contrary. They advised that "Like all interpretations, a meta-ethnography is but a "reading" of what is studied. Other readings are possible and are to be encouraged. However, all interpretations must be grounded in the texts to be synthesized".\textsuperscript{15} Evidence from the reproducibility exercise included within this evaluation is equivocal. Without repeating an entire synthesis, it is difficult to know whether the similarities and differences between the two sets of initial syntheses produced would have become more or less pronounced. (Since the work for this report was completed, we have undertaken an exercise to explore this issue. PP, the key author of
the medicines synthesis, undertook a synthesis of all the papers in the RA synthesis. As expected her ‘reading’ of these papers was a little different, but the overall result was the same as the RA synthesis included in this report: it did not produce any substantial new insights.) Pilot syntheses and comparability/reproducibility exercises may have another valuable function, particularly for the first-time synthesiser, as a rehearsal for the full synthesis.

Noblit and Hare\textsuperscript{15} took a subtle relativist stance and accepted that an interpretation is but one possible interpretation, but also noted that the ethnographer reveals a limited relativism because the range of perceptions is influenced by context and socialisation. Noblit and Hare\textsuperscript{15} therefore adopted a position held by many qualitative researchers that worth is judged by the understandings and explanations derived rather than by assessments of the validity of processes. Noblit and Hare\textsuperscript{15} acknowledged that their approach may be regarded as overly relativist as they were essentially dealing with ‘interpretations of interpretation’. Nevertheless, they identified five criteria drawn from writings of qualitative researchers for what constitutes a good synthesis. These criteria related to the adequacy of metaphors (we have always understood metaphors to mean concepts in this context) in synthesising diverse studies which they suggested needed to be assessed according to the following criteria: economy – a metaphor is adequate when it is the simplest concept that accounts for the phenomena and has a superior ‘ease of representation’;\textsuperscript{234} cogency – a metaphor is adequate when it achieves the explanation without redundancy, ambiguity and contradiction;\textsuperscript{234} range – refers to the power of the metaphor in incorporating other symbolic domains;\textsuperscript{234} apparency – ability of language to (seemingly) ‘show’ us experience rather than merely refer to it;\textsuperscript{235} and credibility – adequate metaphors must be credible and understood by the audience.\textsuperscript{236} Noblit and Hare\textsuperscript{15} therefore, regarded the worth of a synthesis to be judged by the quality of its metaphors and whether or not the synthesis was regarded as useful and insightful by the intended audience.

Audiences

The issue of the audience for qualitative syntheses is an important one. There are a number of potential audiences for the two syntheses contained in this report. For the medicine-taking synthesis the relevant audiences are health policy-makers, clinicians, research funders, academic health researchers and those involved in the manufacture and control of medicines. In the case of the RA synthesis, the most relevant audiences are health practitioners and others caring for and working with people with RA (for example those providing social, educational, environmental, employment or housing services), people with RA and academic health researchers. These audiences are varied and the format for presenting the syntheses needs to vary accordingly. There is still work to be done on how to present and effectively disseminate the findings from qualitative syntheses. There are examples from within the quantitative review tradition that provide possible models, for example the ‘Guides for the public’ produced by the National Institute for Health and Clinical Excellence, which complement the formal guidance issued to health-care providers and any systematic reviews on which that guidance is based (www.nice.org.uk).

Place of meta-ethnography

Qualitative syntheses published prior to the present HTA project (and subsequently) have identified the value of this approach in informing policy and practice by contributing to conceptual and theoretical developments in the field. Qualitative syntheses may also advance the evidence base by achieving an accumulation of findings and enhancing individual studies by checking their comparability and replication. Moreover, qualitative synthesis has a potential empowering function, as it involves methods for combining multiple voices to seek new interpretations, rather than dismissing single case studies as locally bound.\textsuperscript{237}

Meta-ethnographies of qualitative health research could be stand-alone or could be a component part of a systematic review incorporating a variety of different types of evidence. This latter role
may be particularly useful in HTA in which meta-ethnographies of qualitative research could complement quantitative systematic reviews of clinical effectiveness and cost-effectiveness, thus ensuring that patients’ views and experiences are fully represented in the evidence base. The benefits of this can clearly be seen in the case of a review of the neuroprotective drug riluzole in the treatment of motor neuron disease.\textsuperscript{238} The review found that the cost of the drug to the NHS was greater than usually considered acceptable by the National Institute for Health and Clinical Excellence. Qualitative research found mixed views about the acceptability of the treatment for patients: some people with motor neuron disease were resistant to a drug that might marginally extend life without improving its quality; others felt that any extension of life was extremely important. On the basis of these findings, recommendations were made about the kind of information that should be given to patients about the possible benefits of the drug, so that they could make informed choices relating to their own requirements. It also highlighted to the policy-makers that there would be a limit to the number of people who would want the drug if it were made available.\textsuperscript{238}

**Different approaches to meta-ethnography**

The synthesis of qualitative research has opened up a very large agenda, reflecting the range of approaches and theoretical traditions that characterise qualitative research. This breadth and inclusiveness has led to the development of differing approaches to synthesis that reflect the varying assumptions and approaches to qualitative research. In addition, the fundamental distinctions between different schools of qualitative research have led to debates regarding appropriate methods for the conduct of each of the phases of study selection, appraisal and synthesis.

There is often a concern to identify a single ‘best’ approach to synthesis. As Barbour and Barbour\textsuperscript{239} observed, this may result in formalised, proceduralist approaches becoming dominant, and a corresponding neglect of the systematic and meticulous methods of qualitative research and its particular challenges for synthesis, thus leading to a kind of ‘mission drift’. We argue that it is important to acknowledge explicitly these differences and to distinguish between interpretive syntheses based on qualitative methods as developed by Noblit and Hare,\textsuperscript{15} and more structured proceduralist approaches that transpose checklists, standards and other terminology and procedures from quantitative research, with implications for appraisal, sample size, assessment of validity, etc. Whereas proceduralist approaches may be particularly suitable for conducting broadly focused syntheses involving large numbers of reports, or the synthesis of qualitative research for purposes of combining this with a quantitative synthesis, the strength of an interpretive approach based on the qualitative paradigm is in contributing to conceptual and theoretical development. As Hammersley\textsuperscript{14} stated, ‘I don’t think there is a single, all-purpose kind of research synthesis, whether we call it a systematic review or meta-ethnography. I think there are different kinds with different functions that make different demands on us.’ We endorse this view, and believe that what is therefore of paramount importance is to be clear about the aims and purpose of the synthesis and to select methods that correspond to these goals.

The value and general feasibility of qualitative synthesis has been demonstrated, although the development of methods for the synthesis of qualitative research is at present at an early stage and there is a need for further methodological work. Specific issues requiring clarification and/or further research include:

1. A need for an agreed terminology, as the rapid development of new approaches in the field has meant that key terms such as meta-synthesis and meta-ethnography are employed to refer to differing approaches and there is no single classification of methods of qualitative synthesis.
2. Qualitative syntheses have tended to focus on research adopting thematic and phenomenological approaches, and there is a need for further work that synthesises research based on other theoretical traditions and methods.

3. Further experience of undertaking meta-ethnography is required to provide guidance regarding specific aspects of this process. In terms of a qualitative interpretive approach, this includes questions about the advantages of defining phenomena of interest a priori compared with a broader approach to the field; ways of assessing the worth of qualitative synthesis; and the value of approaching and feeding back interpretations to the original researchers. Other questions that are particularly applicable to a more proceduralist approach relate to forms of purposive sampling, the number of studies that are manageable, and ways of reducing context stripping and the decontextualisation of studies.

4. A wider question that requires a comparative methodology is the investigation of how different methods of synthesis influence the outcome of the synthesis in terms of its focus and emphasis, which in turn will help to clarify the applications and contributions of different methods of qualitative synthesis.

5. Further developmental work is required on how to take account of relevant quantitative syntheses while retaining an interpretative approach to the synthesis of qualitative research.

Conclusion

We conclude that meta-ethnography is an effective method of interpretive qualitative synthesis. It allows a body of qualitative research to be drawn together in a systematic way, and can produce novel and important insights, even in fields which appear to have been thoroughly investigated. However, failure to produce new insights does not necessarily mean that the synthesis, or the synthesiser, has failed. It may mean that the field of enquiry has already yielded very rich material in the primary studies. In terms of HTA, meta-ethnography offers a method for including the views of patients in technology assessments in a way that complements the involvement of users/consumers on panels. In particular, it offers a method for including the views of larger numbers of patients in different contexts, and of identifying underlying concepts and explanations. However, meta-ethnography requires considerable qualitative expertise and should be considered an advanced qualitative research method. Meta-ethnography has considerable potential as a method of qualitative synthesis, but it is still evolving and cannot be regarded as a standardised approach to be applied in a routinised way.
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Contribution of authors

A team of people undertook this research project. In the report, where necessary, we have identified who did what. When it came to writing this report, specific people took responsibility for drafting particular chapters and their names are listed below. All authors then had the opportunity to contribute to the final drafts of all chapters.

- Chapter 1  Rona Campbell and Myfanwy Morgan.
- Chapter 2  Myfanwy Morgan, Gavin Daker-White and Rona Campbell.
- Chapter 3  Nicky Britten, Roisin Pill, Pandora Pound and Rona Campbell.
- Chapter 4  Rona Campbell, Lucy Yardley, Catherine Pope and Pandora Pound.
- Chapter 5  Pandora Pound, Gavin Daker-White and Jenny Donovan.
- Chapter 6  Pandora Pound, Nicky Britten, Myfanwy Morgan, Catherine Pope and Rona Campbell.
- Chapter 7  Gavin Daker-White, Jenny Donovan, Roisin Pill and Rona Campbell.
- Chapter 8  Rona Campbell, Myfanwy Morgan and Pandora Pound.
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# Appendix 1

## Appraisal form

### Paper ID

**1. Study information**

1a Study type (Rheumatoid arthritis = 1, medicine taking = 2)

**1b Date of paper’s entry into system (DD/MM/YYYY)**

### 2. Reference information

2a Reference type (1 = Journal article, 2 = Book Chapter, 3 = Book, 4 = Report, 5 = Serial, 6 = Thesis, 7 = Unpublished work, 8 = Conference proceeding, 99 = Other)

2c Title of article

2d Authors in the form Surname, Initials; Surname, Initials; etc.

2e Year of Publication

2f Web address

2g Notes

2h Keywords

2i If we have a copy of reprint, who is keeping it – 1 = Pandora, 2 = Gavin, 3 = Clerical Assistant
2j Journal name

2k Volume number

2l Issue no.

2m Start page

2n End page

2o Editors

2p City of publication

2q Publisher

2r Original language of publication (1 = English, 2 = Other)

2s Disciplinary background of majority of authors (1 = medicine, 2 = social sciences, 3 = nursing, 4 = multidisciplinary, 5 = unclear)

2t Geographical location of study (1 = UK, 2 = other Europe, 3 = North America, 4 = Australia and New Zealand, 5 = Other)

2u Corresponding or first author’s address

2v Was paper identified using an electronic database? (1 = Yes, 2 = No)

IF YES, please answer the following questions (2w–2bb):

2w Was it on MEDLINE (1 = Yes, 2 = No)
2x Was it on EMBASE (1 = Yes, 2 = No)

2y Was it on CINAHL (1 = Yes, 2 = No)

2z Was it on WoS (1 = Yes, 2 = No)

2aa Was it on PsychInfo (1 = Yes, 2 = No)

2bb Was it on Zetoc (1 = Yes, 2 = No)

2cc If paper not identified using electronic database, then how? (Give one option only)
1 = handsearch (hard copy or online), 2 = checking reference lists, 3 = serendipity/browsing, 4 = checking with experts, 5 = Concordance website, 6 = Reference manager

3. Appraiser information
3a Name of first appraiser (1 = Pandora, 2 = Gavin)

3b Date first appraisal done in the format DD/MM/YYYY

3c Name of second appraiser (1 = Rona, 2 = Jenny, 3 = Roisin, 4 = Nicky, 5 = Lucy, 6 = Myfanwy)

3d Date paper sent to 2nd appraiser in the format DD/MM/YYYY

3e Date paper returned from 2nd appraiser in the format DD/MM/YYYY

3f Date of 2nd appraisal in the format DD/MM/YYYY

3g Name of 3rd appraiser (1 = Rona, 2 = Jenny, 3 = Roisin, 4 = Nicky, 5 = Lucy, 6 = Myfanwy, 7 = Gavin, 8 = Pandora)

3h Date paper sent to 3rd appraiser in the format DD/MM/YYYY
3i Date paper returned from 3rd appraiser in the format DD/MM/YYYY

3j Date of 3rd appraisal in the format DD/MM/YYYY

APPRAISAL

AW = author’s words, i.e. respond in author’s words if appropriate (it’s OK to paraphrase)

RW = reviewer’s words, i.e. comment in your own words if appropriate

4. Initial screening questions

Please read whole paper through once before attempting to answer the questions in this section.

4a Does the paper report on findings from qualitative research and did that work involve both qualitative methods of data collection and data analysis?
1 = Yes, 2 = No

4b Comment using authors’ words if appropriate (AW comment)

4c Comment using reviewers’ words if appropriate (RW comment)

4d Is the research relevant to the synthesis topic?
1 = Yes, 2 = No

4e Comment

INSTRUCTION: If the answers to both the preceding questions were ‘YES’, continue with the appraisal. If one or both of the answers were ‘NO’, exclude the paper and elaborate on the reasons for this in the following questions:

4f Is the paper excluded?
1 = Yes, 2 = No

4g Reasons for exclusion

5. Aims

5a Is there a clear statement of the aims of the research?
1 = Yes, 2 = No

5b AW comment
6. Methodology
6a Is a qualitative methodology appropriate for authors’ stated aims?
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>6b AW comment</th>
<th>6c RW comment</th>
</tr>
</thead>
</table>

7. Theoretical perspective
7a Is a theoretical perspective identified?
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>7b AW comment</th>
<th>7c RW comment</th>
</tr>
</thead>
</table>

7d If yes, which theoretical perspective is identified by the authors?

<table>
<thead>
<tr>
<th>7e Comment</th>
</tr>
</thead>
</table>

7e How would you categorise the theoretical perspective?
(See notes at end to aid with categorizing into one of four following options: 1 = Phenomenology, 2 = Grounded theory, 3 = Ethnography, 4 = Action research, 5 = not classifiable according to grid)

<table>
<thead>
<tr>
<th>7f Comment</th>
</tr>
</thead>
</table>

8. Sampling
8a Is it clear which setting(s) the sample was selected from? (e.g. hospital/community)
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>8b AW comment</th>
</tr>
</thead>
</table>
8c RW comment

8d Is it clear why this setting was chosen?
1 = Yes, 2 = No

8e AW comment

8f RW comment

8g Is it clear and adequate information given on who was selected?
1 = Yes, 2 = No

8h AW comment

8i RW comment

8j Is it clear why these samples were selected?
1 = Yes, 2 = No

8k AW comment

8l RW comment

8m Is it clear how the sample was recruited?
1 = Yes, 2 = No

8n AW comment

8o RW comment
8p Is the sample size justified by the authors?
1 = Yes, 2 = No

8q AW comment

8r RW comment

8s Is it clear how many people accepted or refused to take part in the research?
1 = Yes, 2 = No

8t AW comment

8u RW comment

8v Is it clear why some participants chose not to take part?
1 = Yes, 2 = No

8w AW comment

8x RW comment

8y Overall, do you consider the sampling strategy appropriate to address the aims?
1 = Yes, 2 = No

8z AW comment

8aa RW comment

9. Data collection
9a Is it clear where the setting of the data collection was?
1 = Yes, 2 = No
9b AW comment

9c RW comment

9d Is it clear why that setting was chosen?
1 = Yes, 2 = No

9e AW comment

9f RW comment

9g Is it clear how the purpose of the research was explained and presented to the participants?
1 = Yes, 2 = No

9h AW comment

9i RW comment

9j Is it clear how the data were collected and why? (e.g. interviews/focus groups, etc.)
1 = Yes, 2 = No

9k AW comment

9l RW comment

9m Is it clear how the data were recorded? (e.g. audio/video/notes, etc.)
1 = Yes, 2 = No

9n AW comment

9o RW comment
9p Is there evidence of flexibility or an iterative process in the way the research was conducted?
1 = Yes, 2 = No

9q AW comment

9r RW comment

9s Is it clear who collected the data?
1 = Yes, 2 = No

9t AW comment

9u RW comment

9v Overall, do you consider that the data were collected in a way that addresses the research aims?
1 = Yes, 2 = No

9w AW comment

9x RW comment

10. Data analysis
10a Is it clear how the analysis was done?
1 = Yes, 2 = No

10b AW comment

10c RW comment

10d Is it clear how the categories/themes were derived from the data?
1 = Yes, 2 = No
10e AW comment

10f RW comment

10g Is there adequate description of the analysis?
   1 = Yes, 2 = No

10h AW comment

10i RW comment

10j Have attempts been made to feed results back to respondents?
   1 = Yes, 2 = No

10k AW comment

10l RW comment

10m Have different sources of data about the same issue been compared where appropriate (triangulation)?
   1 = Yes, 2 = No

10n AW comment

10o RW comment

10p Was the analysis repeated by more than one researcher to ensure reliability?
   1 = Yes, 2 = No

10q AW comment

10r RW comment
11. Research partnership relations
11a Is it clear whether the researchers critically examined their own role, potential bias and influence?
1 = Yes, 2 = No

11b AW comment

11c RW comment

11d Has the relationship between researchers and participants been adequately considered?
1 = Yes, 2 = No

11e AW comment

11f RW comment

12. Findings
12a Please outline the findings here in as much detail as possible

12b Were the findings explicit and easy to understand?
1 = Yes, 2 = No

12c AW comment

12d RW comment
12e What are the key concepts and interpretations? Please outline in as much detail as possible

13. Justification of data interpretation
13a Are sufficient data presented to support the descriptive findings?
   1 = Yes, 2 = No

13b AW comment

13c RW comment

13d Are quotes numbered/identified?
   1 = Yes, 2 = No

13e AW comment

13f RW comment

13g Do the researchers explain how the data presented in the paper were selected from the original sample?
   1 = Yes, 2 = No

13h AW comment

13i RW comment

13j Do the researchers indicate how they developed their conceptual interpretations of what the data contain?
   1 = Yes, 2 = No, 3 = not applicable (i.e. no conceptual development)

13k AW comment

13l RW comment
13m Are negative, unusual or contradictory cases presented?  
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>AW comment</th>
<th>RW comment</th>
</tr>
</thead>
</table>

13n AW comment

13o RW comment

13p Is there adequate discussion of the evidence both for and against the researchers’ interpretations?  
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>AW comment</th>
<th>RW comment</th>
</tr>
</thead>
</table>

13q AW comment

13r RW comment

| Overall, are you confident that all the data were taken into account?  
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>AW comment</th>
<th>RW comment</th>
</tr>
</thead>
</table>

13t AW comment

13u RW comment

14. Transferability

14a Is there descriptive, conceptual or theoretical congruence between this and other work?  
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>AW comment</th>
<th>RW comment</th>
</tr>
</thead>
</table>

14b AW comment

14c RW comment

| Are the findings of this study transferable to a wider population?  
1 = Yes, 2 = No

<table>
<thead>
<tr>
<th>AW comment</th>
<th>RW comment</th>
</tr>
</thead>
</table>

14d Are the findings of this study transferable to a wider population?
15. Relevance and Usefulness
15a Does the study add to knowledge or theory in the field?
1 = Yes, 2 = No

15b Comment

15c Comment

15d How important are these findings to practice?
1 = Very important, 2 = quite important, 3 = not very important, 4 = not at all important

15e Comment

15f Comment

16. Overall assessment of study
16a What is your overall view of this study?
1 = Excellent, 2 = very good, 3 = good, 4 = not very good, 5 = poor, 6 = very poor

16b Comment

16c Would you include this study in the synthesis?
1 = Yes, 2 = No

16d Comment

17. Time taken to conduct appraisal
17a How long did you take to conduct the appraisal of this paper? Please give time in minutes
### Theoretical perspectives

(To help with Q7e)

<table>
<thead>
<tr>
<th>Research tradition</th>
<th>Focus</th>
<th>Data</th>
<th>Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Phenomenology</td>
<td>Understanding social phenomena/ actions in terms of multiple perspectives</td>
<td>Primarily interviews, but increasing use of observation and focus groups</td>
<td>Constant comparison to describe/ develop explanations (discourse analysis and CA)</td>
</tr>
<tr>
<td>2. Grounded theory</td>
<td>Developing a substantive theory</td>
<td>Theoretical sampling. Multiple sources of data</td>
<td>Well defined procedures to develop theory</td>
</tr>
<tr>
<td>3. Ethnography</td>
<td>Study of a cultural system or group in terms of behaviour, customs and way of life OR Case study (in-depth study of bounded system, e.g. hospital ward, school)</td>
<td>Extensive fieldwork and time in field. Multiple sources of data As above</td>
<td>‘Thick’ narratives</td>
</tr>
<tr>
<td>4. Action research</td>
<td>Participation of those being researched as equals and negotiation of research design</td>
<td>Interviews and observation</td>
<td>Data collection and analysis iterative (responses to research are data). ‘Thick’ narrative</td>
</tr>
</tbody>
</table>
Appendix 2

Protocol

Systematic appraisal and synthesis of qualitative research: evaluating meta ethnography.

Dr Rona Campbell, University of Bristol.

Objectives

1. To conduct, using the meta ethnographic method, systematic appraisal and synthesis of quantitative research studies in 2 areas:
   i. patient experiences of living with rheumatoid arthritis and
   ii. lay beliefs about medicine taking in chronic disease.
2. To test our modified version of the CASP criteria for appraising qualitative research for their appropriateness, ease of use and inter reviewer agreement.
3. To evaluate the meta ethnographic method of synthesis including its reproducibility.
4. To complete our review of the methods available for appraising and synthesising qualitative research.
5. To document the effectiveness of the different elements of the search strategy in identifying relevant qualitative research studies.

Design

Systematic appraisal and synthesis of qualitative research.

Search Strategy

The strategy will include electronic searches of specialist databases and databases of grey literature, hand searching of key journals and collections of qualitative studies published in book form, contacting other qualitative health researchers in relevant areas, searching reference lists of studies retrieved.

Review strategy

The study will test our modified version of the CASP criteria for appraising qualitative research for their appropriateness, ease of use and inter reviewer agreement. The two syntheses will be undertaken using the meta ethnographic method which will be evaluated overall and for its reproducibility.

Expected output of research

1. Report containing:
   i. a review of methods available for synthesis,
   ii. results of the summative evaluation of our modified CASP criteria for appraising qualitative research,
   iii. two qualitative research syntheses,
   iv. an evaluation of the syntheses and a discussion of the contribution qualitative research synthesis could make to HTA.
2. Two papers on each of the syntheses and a methodology paper concerned with the processes of appraisal and synthesis of qualitative research to be published in peer reviewed journals.
3. Presentations of the synthesis findings to different audiences.
Expertise in team

Every member of the research team is a senior social scientist and an experienced qualitative researcher. The research team will be supported by a steering group whose multi-disciplinary membership is drawn from cognate subject areas including public health, epidemiology, anthropology, health economics and rheumatology.
**Health Technology Assessment programme**

**Director,**  
Professor Tom Walley, CBE,  
Director, NIHR HTA programme, Professor of Clinical Pharmacology,  
University of Liverpool

**Deputy Director,**  
Professor Hywel Williams,  
Professor of Dermato-Epidemiology,  
Centre of Evidence-Based Dermatology,  
University of Nottingham

**Prioritisation Group**

**Members**

<table>
<thead>
<tr>
<th>Chair</th>
<th>Professor Tom Walley, CBE, Director, NIHR HTA programme, Professor of Clinical Pharmacology, University of Liverpool</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dr Nick Hicks, Consultant Adviser – Diagnostic Technologies and Screening Panel, Consultant Advisor – Psychological and Community Therapies Panel</td>
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<tr>
<td>Ms Susan Hird, Consultant Advisor, External Devices and Physical Therapies Panel</td>
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</tr>
<tr>
<td>Professor Sallie Lamb, Director, Warwick Clinical Trials Unit, Warwick Medical School, University of Warwick</td>
<td></td>
</tr>
<tr>
<td>Chair – HTA Clinical Evaluation and Trials Board</td>
<td></td>
</tr>
<tr>
<td>Professor Jonathan Michaels, Professor of Vascular Surgery, Sheffield Vascular Institute, University of Sheffield</td>
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<tr>
<td>Chair – Interventional Procedures Panel</td>
<td></td>
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<tr>
<td>Professor Ruairidh Milne, Director – External Relations</td>
<td></td>
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<tr>
<td>Dr John Pounsford, Consultant Physician, Directorate of Medical Services, North Bristol NHS Trust</td>
<td></td>
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<tr>
<td>Chair – External Devices and Physical Therapies Panel</td>
<td></td>
</tr>
<tr>
<td>Dr Vaughan Thomas, Consultant Advisor – Pharmaceuticals Panel, Clinical Lead – Clinical Evaluation Trials Prioritisation Group</td>
<td></td>
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<tr>
<td>Professor Margaret Thorgood, Professor of Epidemiology, Health Sciences Research Institute, University of Warwick</td>
<td></td>
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<tr>
<td>Chair – Disease Prevention Panel</td>
<td></td>
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<tr>
<td>Dr Andrew Cook, Consultant Advisor – Interventional Procedures Panel</td>
<td></td>
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<tr>
<td>Dr Bob Coates, Consultant Advisor – Disease Prevention Panel</td>
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<tr>
<td>Dr Andrew Cook, Consultant Advisor – Interventional Procedures Panel</td>
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<tr>
<td>Dr Peter Davidson, Director of NETSCC, Health Technology Assessment</td>
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</table>

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| Dr Rafael Perera, Lecturer in Medical Statistics, Department of Primary Health Care, University of Oxford |

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Mrs Una Rennard, Public contributor

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Director, NIHR HTA programme, Professor of Clinical Pharmacology, University of Liverpool

Dr Morven Roberts,
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Mr Jim Reece,
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Clinical Senior Lecturer, Cardiff University

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Mr Jim Reece,
Public contributor

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Dr Pippa Tyrrell,
Senior Lecturer/Consultant, Salford Royal Foundation Hospitals’ Trust and University of Manchester

Dr Nefyn Williams,
Clinical Senior Lecturer, Cardiff University

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Dr Simon Padley,
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*We look forward to hearing from you.*