European Research on Migration and Health

International Organization for Migration (IOM)

Background Paper

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“Assisting Migrants and Communities (AMAC): Analysis of Social Determinants of Health and Health Inequalities” project
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Executive summary

This paper reviews the different kinds of research that are required in order to identify, analyse and remedy problems in the field of migrant health. It is divided into five sections (A-E).

Section A explains why research in this field is so important. International migration has increased steadily and health is an important factor affecting the well-being and successful integration of migrants. Conversely, migrants’ health suffers when the transition to a new society is problematic or unsuccessful. A sound knowledge base is crucial to shaping the development of enlightened policies and practices in this area.

Section B describes the different topics that need to be investigated, explaining the specific importance of each of them. These are as follows:

1. Background information on migrants and their situation: demographic characteristics and the social, historical, political and legal situation of (different groups of) migrants.
2. Migrants’ state of health: particular areas of vulnerability and increased risk, as well as strengths.
3. The entitlement of different categories of migrants to health care: gaps in coverage, discrepancies between policies and their implementation.
4. Accessibility of health services: linguistic and cultural barriers, methods for reaching migrant populations more effectively.
5. Quality of health services (prevention, health promotion and care): identifying problems and evaluating solutions.
6. Measures for achieving change: monitoring interventions of different types and at different levels. What are the levers for change?

In Section C, these topics are considered in turn to examine which agencies carry out research on each of them, what methods they use and what difficulties they encounter.

1. The collection of background information on migrants and their social situation is the task of national, local and municipal authorities, as well as researchers in the field of ‘migration studies’. In itself it is not health research, but it is an essential foundation for health research. Researchers may encounter both practical and political objections and obstacles to the collection of such data.
2. Migrants’ state of health can be monitored and analysed using data from clinical or population-based samples. Here too, however, there are barriers to data collection.
3. Migrants’ entitlement to health care can be surveyed on the basis of official policies, but implementation is sometimes at variance with these. Migrants may not know their rights; some health workers may ignore these rights, while others may turn a blind eye to restrictions on care.
4. Accessibility can be analysed using quantitative data on health service utilisation, but qualitative studies are indispensable for the interpretation of these data. Such research investigates migrants’ experiences of health care, the cultural and linguistic barriers they may encounter, as well as conflicting expectations and ideas about health and health care.
5. Investigating the quality and effectiveness of services involves similar issues and methods. There is a serious shortage of studies examining the effectiveness of different treatment methodologies for migrant patients: so much so that it is doubtful to what extent ‘evidence-based’ health care for migrants exists at present. ‘Good practices’ developed for use with migrants are far too seldom evaluated.

6. The study of policy development and levers to change is in its infancy.

Section D argues that international collaboration is essential, particularly within Europe, to make progress on all these fronts. Since 2000, collaborative efforts have multiplied, and the EU, CoE, IOM and WHO have all promoted the topic of migrant health.

Finally, in Section D some important gaps in existing knowledge are identified and the current lack of coordination of research efforts is criticised. It is argued that the establishment of a single European agency to oversee activities in this area – a ‘European Migrant Health Observatory’ – would go a long way to furthering a more effective research effort in this field.
Why is research on migrant health important?

In many parts of the world, particularly in Europe, an increased rate of immigration is confronting host societies with challenges and opportunities – the most familiar being in the labour and housing markets, inter-ethnic relations and the educational system. Until recently, much less attention has been paid to the consequences of migration for the health system, but this omission is hard to justify. Health is an extremely important factor in the lives of migrants and their families. To begin with, it is closely linked with integration:

Migrants who are burdened or handicapped by health problems are hampered in the task of integration. [...]. Illness exacerbates marginalisation and marginalisation exacerbates illness, creating a downward spiral.

At the same time, integration is a prerequisite for effective health care delivery, which is often impeded by inadequate access. Access to effective health care should be seen as no less important than housing and education for the well-being, and thus the integration, of migrants.

(Ingleby et al., 2005, p. 1)

Access to good quality health care is an important aspect of the social inclusion or exclusion of migrants. Existing service provisions are the outcome of a long process of adaptation to the needs of the majority native population: it is only to be expected that they often fail to meet the needs of other groups (Watters, 2002). Most concern with migrant health therefore focuses on the topic of care provision.

However, health is not only determined by the quality of health care – far from it, in fact; most experts regard the environmental factors that influence health as even more important. As the slogan “health in all policies” implies (Stahl et al., 2006), almost all aspects of social life can have an impact on the health of citizens. Poverty and marginalisation are factors which often affect migrants to a disproportionate extent, and in the recently published WHO report on the Social Determinants of Health (CSDH, 2008), particular attention is devoted to the health risks of migrants. Not only poverty, but also bad housing, discrimination and work-related health risks can have a serious negative impact on the health of migrants.

At the present time we can observe an increase in the level of activities aimed at improving the health of migrants and ensuring that they have good access to appropriate care. The amount of activity in a given country or region is related to the proportion of migrants in the local population, as well as to prevailing attitudes to migration. Where public opinion is hostile to migrants and a policy of assimilation prevails, only religious or charitable organisations are likely to show concern about their health. In such countries, the dominant attitude is that migrants must learn to stand on their own two feet and the host society has no obligation to adapt to their presence. At the other end of the political spectrum, more ‘migrant-friendly’ attitudes are accompanied by greater willingness to accept the host society’s responsibility for the social conditions of migrants and to tackle the threats to their health.

Such activities may be justified on humanitarian or human-rights grounds (“health as a fundamental human right”, cf. Pace, 2002), or on the more utilitarian grounds associated with public health policies since their inception in the nineteenth century (“health as a service of general (economic) interest”, cf. Huber, Maucher et al., 2008). In a few countries, migrants and ethnic minorities have enough political influence to be able to lobby directly for healthier conditions and better care.

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1 In this paper, the term “migrant” will be used in the broadest sense, to refer not only to those who change their country of residence voluntarily but also to asylum seekers, refugees and victims of trafficking. Since the consequences of migration may also extend to the second generation or even further, we also use the term to cover the descendents of migrants, rather than referring each time to “migrants and ethnic minorities”.
The importance of a sound knowledge base

The topic of migrant health encompasses two main issues: on the one hand the state of health of migrants, on the other the quality and accessibility of the care available to them. Obviously, any practical initiatives in these two areas need to start from a sound knowledge base. We need to know how healthy or unhealthy migrants are, and what special risks they are exposed to. In what respects do health services fail migrants, and what can be done to remedy these shortcomings? We also need to know which strategies are effective for getting things changed. All this points to the fundamental importance of good research on migrant health.

The knowledge and insights generated by such research can be used at different levels:

- At the level of policy-making (by governments, health care authorities, or individual health care providers), research is needed to inform decision-making. Politicians and managers must know where the problems are, how they can be remedied and how urgent they are.
- At the level of service delivery (by doctors, nurses, clinics, hospitals, public health agencies etc.), research is needed to inform health care workers about the special needs of migrants, so that they can provide more effective care.

Modern health care is increasingly dominated by the principle that all interventions must be ‘evidence-based’. It is not enough to have wisdom and professional experience: facts and figures are required, based on reliable and valid research data. However, these demands are hard to meet in the field of migrant health, where research is to a large extent still in its infancy. The lack of a sound knowledge base is one of the chief obstacles to progress in this area. Fortunately, at the present time many initiatives are under way to improve this situation. In some countries a substantial knowledge base has already been built up, and there has been a dramatic increase in the number of research projects at European level concerned with migrant health.

In this paper we will review European research on migration and health, focussing on the situation in host countries. Of course, issues related to health also arise in the countries of origin: for example, the effect of remittances on household health expenditures, the entitlements of returning migrants, contagious diseases and the effect of separation from the breadwinner on the family left behind. In research up to now these topics have been largely neglected, but with increasing internal migration in the EU more attention needs to be paid to them.
What kinds of research are needed?

In order to reduce a topic to manageable proportions, it is first necessary to break it down into its constituent parts. Ingleby et al. (2005) identified six main areas in which research on migrant health is needed. We will examine each of these in turn.

1. Background information

This topic does not concern health data as such, but other information which is nevertheless indispensable for informed decision-making. Such information includes the number and origin of migrants and their migration history; social and demographic characteristics of migrant populations; the legal position regarding immigration, integration, nationality and citizenship; public opinion and representations of migrants in the media.

Health workers and researchers need such information for two main reasons. Firstly, it is necessary to know the size and structure of migrant and ethnic minority populations in order to reach conclusions about their state of health. For example, general practitioners may have the impression that they see a lot of Turkish men with lower back pain. But how large is the proportion of Turkish men in the area they serve? What is the age structure of this group? In epidemiology, the denominator (the size of the relevant background population) is just as important as the numerator (the clinical data). All too often, information about the background population is simply not adequate for reaching reliable conclusions about migrants’ state of health. As we shall see in the next section, this has proved a major obstacle to investigating migrants’ state of health.

An additional problem here is that there is no international consensus about how individuals should be categorised in terms of their migrant status or their ethnic, religious or cultural background – or, indeed, whether they should be categorised in such ways at all. In several countries, such as France or Hungary, there are severe legal restrictions on the collection and storage of such data. These restrictions are usually there for political reasons, e.g. the republican ideology (“all citizens are equal”) or a desire to make sure that nothing comparable to the ethnic cleansing and genocide practised by the Nazi’s can ever happen again. Such considerations are by no means redundant today, in the light of the ‘moral panics’ that arise from time to time in some countries over such groups as Muslims or Roma. A further objection to laying down hard and fast categories of ethnicity or migration status is this: phenomena that are in essence shifting social constructions should not be treated as if they were timeless facts of nature. Fifty years ago, ‘ethnicity’ was regarded as an objective biological or demographic characteristic: nowadays, even in census data, it is treated as a personal choice.

A second purpose for which background information is important is to enable health workers to interpret migrants’ complaints and to respond in an appropriate way. The importance one attached to this issue will depend on one’s approach to medicine. A purely ‘biomedical’ approach, in which the patient is reduced to a bundle of symptoms and causal mechanisms, will not encourage interest in the patient’s living situation and cultural background: such an approach treats only the disease – not the person who has it. A ‘biopsychosocial’ approach, by contrast, will insist on viewing the patient in their social context, in order to understand what their symptoms might be linked to and what treatment is appropriate. Another word for this approach is ‘holistic’: the fact that this term is increasingly associated with the alternative circuit indicates how strongly modern medicine is dominated by the biomedical approach.

Spectacular recent advances in ‘high-tech’ medicine have helped to strengthen the grip of the biomedical approach. Medical anthropologists, however, point out that there is a crucial distinction between ‘disease’ and ‘illness’: ‘disease’ refers to the physiological processes that may underlie the patients’ complaints, while ‘illness’ refers to the experience of these complaints and their impact on the patient’s daily life. It is
the ‘illness’ for which the patient seeks help; awareness of their social context is crucial to understanding and alleviating it.

The contextual information that we need in order to better understand the illnesses of migrants and ethnic minorities relates to the demographic, socio-economic, legal, political and historical factors influencing the conditions in which they live. What kinds of work do they have, if any? What education and qualifications do they have and how useful are these in the host country? What hardships and deprivations do they experience? Do they have a valid residence permit? Have they claimed asylum, and if so, what will happen to them if their claim is rejected? What is the climate of public opinion concerning their group – are they subject to discrimination? None of these issues are in themselves ‘medical’, but all of them may be relevant in trying to understand migrant health.

2. State of health

Information about the state of health of migrant and minority groups is needed for two main purposes:

a) to identify problems that may call for special efforts in the area of health promotion and prevention;
b) to give health workers insight into the kind of health problems that migrant and minority patients may be particularly prone to, as well as the factors which may underly these problems.

Before going further, however, it is necessary to dispel two widespread misunderstandings about migrants’ state of health. One is that it is necessarily worse than other people’s. A contrary notion, the ‘healthy migrant’ concept introduced by Raymond-Duchosal in 1929 and often confirmed in later studies, suggests that migrants may start off with a health advantage compared with the host population, which they tend to lose over time. However, some of this effect may simply be due to the fact that migrants tend to be younger than the host population. Another possible explanation is that immigration policies sometimes restrict the migration of people with health problems. Even if there is, among some groups, a ‘healthy migrant effect’ at the point of arrival, most migrants will occupy an inferior socio-economic position in the host country, and this in itself will undermine their chances of staying in good health.

Nevertheless, there are certain respects in which migrants may enjoy a health advantage. In the UK, some minority ethnic communities have higher levels of education and employment, and lower levels of disability, than White communities (Aspinall & Jackson, 2004). According to McCormack et al (2008), “breast cancer incidence rates vary sixfold between industrialized and less-developed countries, and migrants from low-risk countries to high-risk countries have an intermediate risk”. Cultural and religious practices may offer health advantages over a modern Western life-style: for instance, Muslims who adhere strictly to the ban on alcohol are less at risk for alcohol-related diseases. These examples are sufficient to demonstrate that it is foolish to attempt any generalisations about the general level of health of all migrants. The answer will depend on the condition one is studying and the particular migrant group concerned. Sometimes, indeed, the findings will also differ between men and women, or between first-generation migrants and their children.

A second misunderstanding concerns the importance of health disparities for action on migrant health. Health disparities can indicate areas of special need where extra attention should be paid to research, prevention, health promotion and treatment. However, it is not necessary to show that migrants have particularly severe or unusual health problems in order to justify the provision of appropriate and accessible health care services for them. Indeed, the assumption that migrants always have unusual health problems is part of an older discourse which has its roots in colonialism, in which the migrant is portrayed as threatening, alien or exotic. More often than not, the health problems for which migrants seek help are ‘common-or-garden’ complaints that anybody may suffer from.
As we will explain in the next section, research on the state of health of migrant and minority groups uses two main methods: **clinical studies** and **population-based (epidemiological) surveys**. The former generally take specific diagnoses as their starting-point, while the latter are usually confined to information that can be provided by informants themselves (self-report). Often, respondents are asked to make a subjective assessment of their general level of health. Such assessments often reveal striking differences, but it is difficult to know to what extent the self-ratings correspond to more ‘objective’ measures of health.

Investigations of migrants’ state of health sometimes go further and attempt to explore the determinants of particular health problems. Stronks et al. (1999) put forward a model in which four basic contextual factors can underlie ethnic differences in health:

1. the process of migration
2. cultural factors
3. socio-economic position
4. social context

Specific determinants of health or illness which may arise in these contexts are:

1. genetic factors
2. lifestyle
3. physical environment
4. social environment
5. psychosocial stress
6. health care utilisation

### 3. Entitlement to health care

The next topic on which good research is vital concerns the conditions under which migrants are entitled to receive health care. This question is usually subsumed under the topic of ‘access’, but we consider it sufficiently important and distinctive to deserve separate consideration.

Entitlement to care can be broken down into three components: ‘coverage’, ‘health basket’ and ‘cost-sharing’ (Huber, Stanciole et al., 2008). ‘Coverage’ refers to whether a person’s health expenses are paid for by a State or private insurance scheme. ‘Health basket’ refers to the range of services that are covered, while ‘cost-sharing’ refers to the out-of-pocket financial contribution which is required from the service user.

The entitlement of migrants to health care varies from country to country and according to the category of migrant concerned (employed, unemployed, asylum seeker, undocumented, etc.)

The rules governing entitlement are often complex, fast-changing and poorly understood by migrants – and even by the people supposed to be applying the rules. Research is needed to clarify the situation in each country and its consequences for health.

### 4. Accessibility of care

We use this category to refer to obstacles to obtaining care other than problems of entitlement. In order to access care, a sick person (or someone in their environment) must first of all realise that they need it. This will depend on their level of ‘health literacy’, in particular their knowledge about the treatments available and the signs that treatment is necessary.
It is sometimes said that health literacy tends to be low among migrants, but it would be more accurate to say that knowledge about health is of different kinds, and the knowledge that migrants have is often at variance with that of the majority population. Particularly when a person has only recently migrated or has not had much contact with the host society, their knowledge about illnesses and health care is likely to reflect mainly the culture and health system of their country of origin.

Moreover, different cultures may have widely different ways of categorising, describing, assessing and responding to illnesses. Kleinman (1978) introduced the concept of ‘explanatory models’ to describe these culture-bound belief systems. Health care systems also differ enormously across the globe, both in their formal structures and in the unwritten rules governing interactions between health care workers and patients. Because of this, it can easily happen that migrants are perceived by health workers as unaware of basic ‘facts’ about health, having ‘irrelevant’ expectations of the health system, behaving in ‘inappropriate’ ways, and so on, when all they are doing is drawing on the stock of health knowledge they have acquired in their country of origin.

Although one can argue about the relative merits of different medical concepts and systems, it is necessary for migrants in European countries who wish to use mainstream health services to learn how people in the host society think about health, how the health care system is structured, and what the formal and informal ‘rules of the game’ are for both staff and patients. Indeed, everybody has to acquire this knowledge and keep it up to date: it is an important ingredient of socialisation and ‘health citizenship’.

A basic task for researchers in this area is to find out whether, and if so in which areas, health care is less accessible for migrants. A crude indication of this is the level of ‘care consumption’: if fewer people are using a service than would be expected on the basis of the (assumed or measured) incidence of illness, this may be a sign that they need help in order to find their way to the service.

Another explanation of a reduced take-up of services, however, may be that people know perfectly well that the service is there, but have different views on what is wrong with them and what should be done about it. This is especially likely to be the case when Western medical views differ markedly from those prevailing in source countries.

In the West, the medical domain has expanded enormously in the last 50 years. This has been accompanied by a substantial increase in expenditure on health services – most notably in the USA, where total per-capita health spending, adjusted for inflation, has increased eightfold during this period². In the larger West European countries, expenditure is currently around 50% of the US level – but in many of the countries that migrants come from the corresponding percentage is far lower (e.g. Afghanistan, 0.4%; Pakistan, 0.7%; India, 1.5%; Morocco, 4%; China, 5%; Turkey, 9%; most African countries, below 1%)³. These figures show that in terms of the amount spent on health care, the contrast between sending and receiving countries can be very extreme (see also CSDH, 2008).

A higher amount spent on health care in a given country will go hand in hand with increased readiness to seek medical help and a broadening of the criteria for ‘illness’. In the West, conditions which were previously regarded as natural or inevitable, or not medical conditions at all, now qualify for diagnosis and treatment. The field of mental health is a prime example of this, with a massive increase in the number and variety of problems regarded as signs of ‘mental disturbance’. In 1952, the Diagnostic and Statistical Manual of the American Psychiatric Association listed only 60 categories of abnormal behaviour; in 1994 this number had increased to 410. It is therefore understandable that migrants coming from countries where psychiatric provisions are few and primitive, catering only for the most extreme cases, may resist being told that they have a ‘mental health’ problem. They are likely to associate the term with extreme forms of insanity and will therefore go out of their way to avoid having it applied to them or their family members.

² This figure is based on data from Reinhardt (2002) and other sources.
³ The figures are for 2005 and were obtained from the WHO Statistical Information System (WHOSYS).
The medicalisation of ageing, pregnancy and childbirth provides a further illustration of this Western trend. People brought up to regard these as natural processes which should be allowed to run their course – a belief not confined to migrants, but shared by many Europeans – resist the idea that a pregnant or elderly person should live their lives under constant medical supervision. Poor take-up of antenatal care and low compliance with medical advice may thus be more a result of differing health beliefs and values than of ignorance about what is available. In this connection it is important not to assume axiomatically that modern Western practices are preferable to all others and that the only thing that needs to be done is to educate migrants in ‘good’ health care attitudes. The massive investment in mental health care services in the West has not been accompanied by a commensurate improvement in the happiness of its citizens; this calls for appropriate modesty about the superiority of Western approaches. Though the USA spends more of its GNP on health care (currently about 17%) than any other country in the world, many other countries enjoy better levels of health.

Besides divergent health beliefs and values, there may be practical barriers to using a particular health service (e.g. location, transport, opening hours), or social barriers in the form of stigma and fear of gossip (particularly, as we have seen, in the case of mental health problems). Another factor that can discourage service use is be a lack of consideration for cultural practices and customs (for example, concerning hospital food or contact between female patients and male doctors – although it is worth remembering that many European-born women sometimes prefer a female doctor.) Such barriers to access fall into the category of ‘institutional discrimination’, a term which gained currency in the context of the American Civil Rights Movement: a recent definition is as follows:

Institutional discrimination occurs when the culture, policies, systems and procedures in an organisation inherently discriminate against a group or groups of people. This happens because the systems and processes were designed without taking into account the diverse needs of groups within the community in relation to e.g. their race, disability or gender (ESCC, 2009).

Institutional discrimination is largely unconscious, but access to health services by migrants can also be undermined by conscious, direct, individual discrimination at any level from receptionist to consultant. Since 2008, the European Union Agency for Fundamental Rights (FRA) has included health care among the areas in which it produces annual reports on (direct) discrimination. In most countries, however, this form of discrimination seems to be much less of a problem than the institutional kind.

A highly specific barrier to accessing health care is the threat to undocumented migrants that they will be denounced to the authorities if they do so. Some countries require health care staff by law to report undocumented migrants who make use of their services; even where there is no legislation requiring this, such migrants may perceive the risk of denunciation to be greater than the risk of leaving health problems untreated.

As we have seen, ‘accessibility’ is a broad concept covering many diverse phenomena, and many kinds of research are necessary to investigate the accessibility of health care for migrants. For example, we need to know whether migrants are adequately informed about illnesses and the health care system: health authorities need to know what information is needed and in which languages, and how it can be most effectively presented. We also need to know how migrants’ beliefs and values concerning health may be at odds with those presupposed by health care workers. Research is also needed to uncover direct and indirect discrimination.

And it is not only the problems that need to be investigated: so, too, do the solutions which are proposed for them. Research is needed to evaluate the effectiveness of the ‘good practices’ that are put forward to make services more accessible. In this respect, health education and health promotion are important but often-neglected areas. People’s behaviour is notoriously intransigent when it comes to avoiding health risks, and it is a great challenge to devise methods that are actually capable of changing behaviour. In particular, research is needed to identify the most effective methods of persuading members of migrant and ethnic minority communities to take good care of their health.
5. Quality of care

While ‘accessibility’ refers to obstacles on the path to care, ‘quality’ refers to what happens in the caregiving situation. Of course, many migration-related factors, such as language barriers, divergent health beliefs and discrimination, undermine both the accessibility and the quality of care. A wide range of interventions and methods have been put forward as ‘good practices’ for improving the quality of health care for migrants. The task of the researcher is (a) to find out when something is going wrong in the treatment situation, (b) to identify what it is, and (c) to evaluate the different solutions proposed for dealing with it.

To assess the quality or appropriateness of care, a number of measures can be used. Subjective measures set out to measure the degree of satisfaction of clients and their caregivers. This can be asked for directly, or estimated from levels of drop-out or compliance. Objective measures investigate the effectiveness of particular practices. Do the standard procedures lead to worse outcomes for migrant patients? Are particular ‘good practices’ capable of producing better outcomes? In order to evaluate a procedure or treatment method, researchers need ideally to carry out a ‘randomised clinical trial’ or RCT, in which patients are allocated at random to different groups (preferably without them or the researcher knowing which group they are in, i.e. ‘double-blind’). However, the practical and ethical obstacles to such research, which is usually very expensive, mean that RCT’s are hardly ever carried out to establish the effectiveness of treatments or ‘good practices’ for patients from migrant or minority ethnic groups.

6. Achieving change

Under this heading researchers examine all the activities that are undertaken to encourage the development of adequate health care for migrants. At the outset we can make a distinction between ‘structural’ and ‘incidental’ changes.

- ‘Structural’ improvements to health care services as those that are embedded in policy. Policy may be laid down at national, local or municipal levels, as well as by service providers or professional bodies.
- ‘Incidental’ improvements arise more spontaneously, as a result of the activities of individuals or NGO’s responding as they see fit to perceived needs.

However, the boundary between the two may shift overnight: an intervention may start out as a spontaneous individual initiative, yet because of its success receive the blessing of an institution or a ministry and become incorporated in official policy. The boundary between ‘structural’ and ‘non-structural’ provisions may also be blurred: in countries where there is little recognition by government of the health needs of migrants, services run by NGO’s such as Médecins du Monde, or religious organisations such as the Jesuit Refugee Service, may become virtually part of the landscape and even receive funding from central government. Nevertheless, there is increasing realisation that the structural embedding of measures in policy is essential for sustained progress (see the section “Why ‘good practices’ are not enough” in the report on Good Practices for the Portuguese EU Presidency Conference (Portugal et al., 2007, p. 17)).

Under the heading of ‘achieving change’, researchers examine the wide range of activities that may be undertaken to lobby for migrant health, to bring together those interested in it, to form pressure groups, centres of expertise, research networks and ‘think tanks’. One of the most important of all these activities is teaching — educating students, professionals, politicians and the public about the importance of migrant health and showing how research findings can be applied.
Who carries out this research, what methods are involved, and what are the difficulties?

1. Background information

Much of this information is collected by national, regional or municipal authorities. As mentioned in the previous section, the amount of official information concerning such variables as (parental) place of birth, nationality, migrant status, ethnicity or religion can range from non-existent in some countries, to fairly detailed in others. Where official statistics are scarce, researchers have to try and remedy this hiatus as best as they can themselves.

In spite of the various legal and practical barriers to data collection, it is increasingly common for population data to be collected on people’s country of birth (and that of their parents), or their nationality, religion, or ethnicity. Unfortunately, the precise data available vary from country to country, making trans-national studies of migrants’ state of health very difficult. As far as demographic data on migration are concerned, Europe resembles a patchwork quilt.

Apart from data which is routinely collected on all inhabitants, many governments (national or local) conduct or commission surveys to collect specific types of information, including variables relating to migration or ethnicity, on a particular age cohort or sample of the population. Sometimes these surveys are longitudinal and enable conclusions to be drawn about the causal factors affecting people over the life course. The UK and Scandinavian countries have perhaps the most extensive collections of such data on their inhabitants. Thanks to this, and to the possibilities for linking different data sets with each other, these countries have been able to undertake some of the most large-scale surveys on the health of migrants of ethnic minorities. A recent review article (Ingleby, 2008) describes Scandinavian and British studies published in recent years on the incidence of schizophrenia among migrants, which have revolutionised views on this topic with the help of databases covering millions of people.

Academic research in the area of ‘migration studies’ provides a copious source of data on topics such as education, housing, legislation, discrimination, public opinion, policy making and (media) representations. To make this data accessible it is necessary for researchers in the field of health to collaborate closely with their colleagues in the social sciences. An example is the work of Adam & Devillard (2008), which reports a study carried out by the IOM reviewing immigration laws in the 27 EU Member States.

2. State of health

Two main types of research can be carried out to reach conclusions about the state of health of particular social groups such as migrants and ethnic minorities: clinical studies and population-based studies.

Clinical studies

These studies start from data generated in contacts with health service providers (family doctors, hospitals, well-baby clinics, etc.) For example, the number of patients with a particular diagnosis or receiving a particular type of treatment may be recorded. This can either be done routinely, or at the special request of a researcher. Where such data are collected routinely, existing clinical records will provide a wealth of information for the researcher.

However, the prevalence of a given condition among migrants or ethnic minorities can only be estimated from such data if (1) information about migration or ethnicity has been collected at the same time, and (2)
data are available on the size of the underlying populations. Data on the underlying populations falls in our category of ‘background information’, and it will be obvious that the best clinical data in the world is useless for estimating prevalence rates if good statistics are not available about the underlying population.

We can illustrate this point with an example. Consider – once again – general practitioners recording the number of male patients of Turkish origin reporting lower back pain. Suppose we have data from all the GP’s whose catchment area falls in the city of Rotterdam. We will only be able to estimate incidence rates from these data if, at the same time, data are available on the ethnic origin and gender of the inhabitants of Rotterdam. It will also be very helpful to have data about the age structure of different groups; if the Turkish inhabitants are older than average, this could be the reason for the higher incidence, because lower back pain is associated with ageing.

But there is yet another problem concerning the use of clinical data. Incidence rates in the clinic are only equal to incidence rates in the general population if access is perfect. In this example, it could be that access by Turkish men to health care is limited, for all the reasons we have discussed above. Figures obtained from clinical practice shed no light at all on what is going on with the people who do not show up at the clinic. As Goldberg & Huxley (1980) point out, a number of ‘filters’ intervene between illness and treatment.

In general, we can say that the numbers of people who receive treatment will reflect two things: the proportion who are ill, and the proportion of these who succeed in getting into treatment. If (and only if) one of these variables is known, it is possible to estimate the other. But if we know nothing about accessibility we cannot say anything on the basis of clinical data about incidence; and if we know nothing about incidence, we can say nothing about accessibility. For this reason, figures concerning the amount of ‘care consumption’ by different groups need to be interpreted with great caution.

Many other problems surround the use of clinical data (for example, the reliability and validity of diagnoses), but it will be clear by now that the use of such data to shed light on the state of health of the migrant and ethnic minority population is fraught with difficulties. In many health care settings, information relating to migration or ethnicity is simply not recorded. Even when it is, the relevant catchment area may not be known, and there may be no information concerning the size of the migrant or ethnic minority populations in that area – let alone concerning gender and age. Moreover, access may vary between groups. Fortunately, however, there are other types of studies which can be used.

Population-based studies

For epidemiological purposes there are enormous advantages in collecting data on health from the general population, rather from the highly selected sub-sample which is found in clinical settings. By doing this, of course, one immediately loses the advantages of the clinical setting: there, a diagnosis has to be made anyway, often with the help of highly sophisticated procedures. Outside the clinic, the information that can be collected is more basic in nature, and generally depends on self-report data from the respondent. Only when funds are available for medical screening of a group can population-based studies begin to match the sophistication of the diagnoses that clinical data is based on.

Population-based studies may concern very large populations, or they may be quite small-scale; they may be focussed on a single health problem, or cover a whole range of conditions. Increasingly, governments carry out large-scale surveys that include questions about health in order to monitor the health needs of their citizens. For our purposes, however, such surveys are only useful if questions are also included about migration status or ethnicity. It may, however, also be possible to couple population data with health data obtained from special surveys.

The following indicators may be used to assess the general health of a group:
The mortality of the group (i.e. the death rate). Such figures have to be interpreted with great caution. In the first wave of any migration, the people who migrate tend to be fit and healthy, ready to face the hazards of journeying far from home. Some migrant groups keep close contact with their home country and return when they become sick or old. Their deaths are not recorded in the host country, so that official statistics on mortality rates may not give an accurate impression of the health of that group.

Inaccuracies may arise in several other ways. For example, if members of some ethnic groups are inaccurately classified in a national census, but their origins are accurately recorded when they die, then the death rate of these groups will be overestimated. Conversely, if ethnicity is not correctly classified when a person’s death is officially registered, then the death rate will be underestimated.

A widely used indicator of general health is life expectancy, but this can also be difficult to interpret in the case of migrants. For instance, a Dutch study showed that life expectancy for Moroccan-born men is 3.5 years longer than for Dutch natives, while for Turkish and Surinamese-born men it is 1.5 years shorter. The explanation for these differences is very unclear (RIVM 2002). It may have something to do with the tendency of migrants to return home when they become old or sick.

Another indicator that is often used in survey research is subjective health. When asked about their experienced state of health, 79% of the native Dutch population describe this as ‘good’ or ‘very good’, compared to 71% for first-generation Western immigrants and only 63% for first-generation non-Western immigrants (CBS 2004). Similar findings have been reported in the United Kingdom, e.g. by Dunnell (2008), and in Switzerland (Garbadinho et al. 2007). However, the cross-cultural validity of such self-report measures is unknown (Chandola & Jenkinson, 2000; Bruijnzeels, ed. 2004: 89; Menec et al., 2007).

Epidemiological research on migrants and ethnic minorities is in its infancy and most of the information currently available on their health is based on clinical studies – with all their attendant disadvantages. Bhopal (2007) gives an authoritative introduction to this area of research. Such studies belong to the domain of Public Health: they may be carried out by (large) health care providers, municipal authorities, national health research agencies, independent contractors, NGO’s and university-based researchers. However, the resources required for epidemiological research are usually substantial, so that only major organisations are able to fund them.

3. Entitlement

Research on the entitlement of migrants to health care is in its infancy, although Appendix A lists a number of comparative studies – mostly carried out on behalf of international agencies or NGO’s. In theory, one can learn everything one needs to know about migrants’ entitlement to health care by studying documents setting out the policies of state health care schemes or health insurers. In these documents the answers to questions about who is covered, for what kinds of treatment, and how much money they have to contribute out of their own pockets, are set down in black and white. Nevertheless, we cannot assume that theory and practice overlap perfectly when it comes to entitlement. Rules have to be interpreted (for example, regarding the definition of what constitutes ‘emergency care’) and different service providers may operate different definitions. Moreover, knowledge of the rules may be incomplete or out of date: for example, in response to the current confusion about the health care entitlement of asylum seekers and undocumented migrants in Britain, new guidelines and summaries follow on each other’s heels with bewildering rapidity.

‘Implementation gaps’ between policy and practice seem, in fact, to be inherent to modern organisations (Hogwood & Gunn, 1984). Sometimes service providers will be more permissive than the rules allow; health workers will turn a blind eye to policy restrictions, out of a sense of professional responsibility or concern for human rights. Sometimes, however, they will be more restrictive, and will withhold information about clients’ rights because they disagree with the policy, or because of prejudice against the
person they are dealing with. In this respect a receptionist at the hospital’s front desk can wield more power than the rest of the medical hierarchy.

4. Accessibility

Research on accessibility often takes quantitative data as its starting-point and proceeds to investigate it qualitatively. Where there are signs that a given service is being ‘under-used’, research may be undertaken to find out what is holding people back from using it: generally, the best way to find this out is to ask them. In theory, quantitative methods such as postal questionnaires could be used, but there are doubts about the validity of data gathered in this way. If a person is mistrustful of a service, they will probably also be wary of someone carrying out research on behalf of the service.

Researchers often regard migrants and ethnic minorities as difficult groups to investigate. Because relations between them and the majority population are often strained, there may not be much willingness to cooperate if the researcher is perceived as representing ‘the authorities’, or simply looks white and middle-class. Often, people are tired of being approached at regular intervals by researchers; they may also have misgivings about the use to which the data will be put. The researcher’s conclusions will then be based on those who do cooperate, whereas the opinions of those who don’t may be much more relevant. One way of overcoming some of these problems is to employ members of migrant or ethnic minority groups to carry out the research. A striking example of this is the book edited by Clarke (2005), reporting research on migrants in Finland carried out entirely by migrants.

Studies on the accessibility of health care for migrants and minorities are a fairly recent phenomenon and are typically small-scale in nature. Whereas epidemiological research is mostly large-scale and expensive, studies on accessibility are often commissioned by individual service providers. Many of these studies are classified as ‘grey literature’: because they are not generalisable, they do not make their way into mainstream scientific literature but instead enjoy only limited circulation. Hundreds of such studies have been undertaken by students for their final-year dissertations, and even these can provide highly useful data. In recent years, internet has provided possibilities for disseminating ‘grey literature’ cheaply and on a wide scale: local authorities and NGO’s are making increasing use of this medium to publicise their findings.

5. Quality

There are two types of evidence that are relevant to the assessment of the quality of care. First, evidence about cultural incompetence – the ineffectiveness of standard approaches which do not take any account of diversity; second, evidence about cultural competence – the effectiveness of measures for dealing with diversity. Who carries out this research and what methods are used?

Regarding the first type of evidence, we have already mentioned the fact that in this era of ‘evidence-based medicine’, proof of the effectiveness of new procedures and treatment methods is required before they are introduced. However, such studies usually pay no attention to the possibility that effectiveness may vary between different ethnic groups. Worse still, members of minority populations may actually be excluded from clinical trials. Graham (1992) showed that 96 per cent of the studies published in the 1970s and 1980s in four leading journals of the American Psychological Association excluded African American subjects.

There is, in fact, little hard-and-fast data available on differences in the effectiveness of treatments for different ethnic groups. However, some drug companies have begun to highlight this issue, in the hope of identifying products which can be marketed as specially appropriate for particular groups.
Concerning the evaluation of ‘good practices’ introduced to increase cultural competence, a major report was compiled in the USA by Fortier & Bishop (2003) in which many hundreds of studies were reviewed. Also in the USA, Griner & Smith (2006) demonstrated considerable effect sizes of measures to improve cultural competence in a meta-analysis of 76 studies concerned with mental health programmes. However, Bhui et al. (2007), who examined 109 articles describing programmes for improving cultural competence among health care professionals in the UK, found that only nine of these had included an evaluation of the model described.

On the topic of health promotion, even less work has been reported. The research project ‘Healthy Inclusion’ is currently being carried out in seven countries to improve the effectiveness of health promotion activities for migrants.4

As noted above, most research in this area focuses on satisfaction or ‘procedural evaluation’ and much of it is qualitative in nature. Often, such studies are undertaken by the originators of the method and are thus far from impartial. However, the state of the art in this area is continually improving and much current research is published in high-quality, peer reviewed journals.

6. Achieving change

To find out what activities are being undertaken in order to improve health care for migrants and minorities, studies have usually been undertaken by national research institutions or by researchers in universities. Systematic surveys involve contacting service providers and asking them a number of questions about the measures that have been taken to improve the accessibility and quality of services for these groups. Alternatively, a qualitative overview of initiatives taken may be made on the basis of literature reviews, internet searches and ‘snowball’ sampling methods, in which one informant will suggest other informants.

Concrete examples of research in the above areas

A good impression of the ‘state of the art’ in the six areas we have discussed can be obtained from the extensive review undertaken by Philipa Mladovksy (2007) of the Health and Living Conditions Network of the European Observatory on the Social Situation and Demography, European Commission. During 2009, detailed information from 17 countries will become available from the project MIGHEALTHNET (http://mighealth.net), which will be summarised as one of the outputs of the project.

Research at the European (or global) level

Thanks to the activities of bodies such as the WHO, the IOM, the European Commission, the Council of Europe and various international NGO’s, opportunities have arisen for carrying out international collaborative research on migrant health. In this section we will discuss the advantages of this kind of research and discuss some examples.

As far as Europe is concerned, perhaps it would be more appropriate to start by describing the disadvantages of confining research to the national level. Imagine what the state of research would be in the USA if each of the 50 states spoke a different language and there were no nation-wide arrangements for organising research, sharing knowledge and regulating professional organisations. Imagine also that few researchers moved from state to state. Yet this is the baseline situation in Europe. While not wishing to underestimate the richness of Europe’s enormous cultural and social diversity, the fact that health research is mainly carried out on a national basis is a great obstacle to progress. It reduces the scale of efforts and limits the possibilities for sharing data, methods and insights. For this reason, the emergence of health research at a European level is greatly to be welcomed – especially in relation to migrants and ethnic minorities.

Particularly important is the possibility of joining forces to increase the ‘critical mass’ of the research community. In each country, migrants and ethnic minorities form a small segment of the population whose interests easily get overlooked. As a result, research on migrant health often occupies a highly marginal position. On a European or international level, such research relates to far more people and can therefore claim more resources.

Apart from these advantages, cross-national comparisons of patterns of migrant health or approaches to health care can yield new insights into underlying factors and the nature of ‘good practice’. Are the higher rates of cardiovascular disorders among some Asian communities in the UK also found in the rest of Europe? What does the distribution of these disorders tell us about the causes? What are the relationships between national health-care philosophies and provisions for migrants? Under what conditions are particular methods effective? The use of data from several different countries introduces greater variability in the factors studied, making it easier to disentangle the influence of different factors.

In Appendix A a number of international collaborative research studies are described, together with a short description of their scope and the results of completed projects. The matrix of DG SANCO projects constructed by the IOM in the context of the AMAC project represents an important step towards obtaining an overview of current efforts.
Conclusions

Though research on migrant health has shown remarkable growth during the last decade, there still remain many ‘blind spots’ where research is urgently needed. To start with, we make here three suggestions – many more are possible:

1. An inventory needs to be made of legal, ideological and practical barriers to the collection of data on migrant status or ethnicity in each country. The MEHO project (Migrant and Ethnic Health Observatory), co-financed by DG SANCO, has a special project on “Data sources and indicators”, but this does not investigate legal restrictions. A study is urgently needed which would clarify the situation and explore ways of overcoming the obstacles.

2. As noted above, ‘good practices’ to improve the accessibility and/or quality of health care delivery are seldom evaluated. More attention to this question is required. The methods used to do not have to be RCT’s – in fact in many cases it would be impossible or unethical to satisfy the conditions for an RCT. As Fortier and Bishop (2003) show, many other research strategies (e.g. ‘process evaluations’) can be used to assess these value of these approaches. The participation of migrants themselves in the design, implementation and evaluation of interventions is of course crucial.

3. Finally, as we noted at the beginning of this paper, hardly any attention has been paid to health issues connected with migration in the countries of origin. Thanks to successive enlargements, the EU contains many such ‘sender’ countries and research in this area needs to be expanded.

We have seen that international collaborative projects are of great value in this area, and it is encouraging to note the substantial increase in funding for this type of work. However, as will be obvious from a study of Appendix A, these efforts suffer from a lack of cohesion and a systematic research policy. A rigidly centralistic, top-down policy would have perhaps even greater disadvantages: it would stifle creativity, innovation and diversity. Yet the philosophy of “let a thousand flowers bloom” can lead to wasting of scarce resources. At present, new projects arise in a fairly haphazard way: different agencies do not coordinate their efforts with each other and there is little synergy between different projects, with the result that efforts are not spread in a rational way. Moreover, knowledge does not seem to develop in a cumulative fashion. Considering that the first studies were undertaken as long ago as 1983, there is a great deal of repetition of the same findings and recommendations (‘reinventing the wheel’).

One illustration of the lack of a cohesive policy is the fact that it is often very unclear why certain countries get included in an international project or network while others are not. Inclusion probably has most to do with the international contacts that the proposer of the project happens to have in his or her address book.

Coordination of efforts is required not only within funding agencies, but also between them. For example, the European Commission’s DG SANCO, DG Research and DG Employment, Social Affairs and Equal Opportunities have all commissioned pioneering studies in this area; yet there is no sign that efforts have been coordinated between these agencies. The same remarks apply to the lack of harmonisation between EC-funded projects and those initiated by the WHO, the IOM and various large private foundations.
In conclusion, we may observe that while diversity in research is in itself a good thing, more attention should be paid to the need to develop shared approaches and to avoid unnecessary duplication of research effort. Projects such as the COST Action HOME (Health and Social Care for Migrants and Ethnic Minorities in Europe) can help to reduce the fragmentation of research efforts, but a more coordinated (‘joined-up’) approach by the EC and other international bodies is also urgently required. The establishment of a single European agency to oversee research activities in this area – a ‘European Migrant Health Observatory’ – would go a long way to furthering a more effective research effort.
Note about the author

David Ingleby graduated in Psychology from Cambridge University. After working as researcher for the Medical Research Council he returned to teach at Cambridge. In 1984 he was appointed Professor of Developmental Psychology at Utrecht University. Since 1991 he has focused on health care, cultural differences and the situation of migrants and minority groups. He is Scientific Coordinator of the project MIGHEALTH.NET and Chair of the COST Action HOME. In 2007 he held the Willy Brandt Memorial Chair at Malmö University: at present he is Consultant to the Council of Europe Committee of Experts on Mobility, Migration and Access to Health Care.

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References


Appendix

Published reports and collaborative projects on migrant and ethnic minority health in Europe

This appendix lists reports and projects in which issues concerning migrant and ethnic minority health are examined in a number of European countries simultaneously.

1. Published reports on migrant and ethnic minority health in Europe

Fig 1. Increase in number of published reports, 1985 – 2008

![Bar chart showing the increase in number of published reports from 1985 to 2008.](chart)

Note: the total for 2005-2008 has been adjusted to take account of the fact that it is a four-year period, not a five-year one.
Which countries have been surveyed?

For some reports – not all – it has been possible to list the countries which have been surveyed. In this table you can see which countries are mentioned in which reports.

### Report (see table on next page)

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Table 1. Countries covered in each report
**Summary table**

Please note that it is not certain whether this list of reports is exhaustive. However, we have tried to make it as complete as possible.

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* For some reports it is not possible to determine the number of countries studied
** Second phase of project 7

**Abbreviations, number of projects sponsored**

DG Directorate General (EC)
EC European Commission
ERF European Refugee Fund (DG Justice) (1)
ESAEO Employment, Social Affairs and Equal Opportunities (5)
ICMH International Centre for Migration and Health (2)
IOM International Organisation for Migration (2)
JUSTICE Justice and Home Affairs (1)
MdM Médicins du Monde (1)
MRC MRC (UK government) (1)
RESEARCH Sixth Framework programme (1)
SANCO Health and Consumers (2)
WHO World Health Organisation (2)
List of reports


Summary: This book addresses the research and policy issues that emerge from the interface of different cultures as a consequence of migration. It includes articles on the following issues: (1) the contribution of the social sciences to an understanding of migrant health needs; (2) health care across cultural boundaries; (3) health care for labor immigrants; (4) philosophical considerations of health care policy and the position of ethnic minorities; (5) health care research and evaluation in a host country: The Netherlands; (6) health care and Moroccan and Turkish immigrants; (7) problems of health and health care research with particular reference to ethnic minorities; (8) health care research and evaluation from the country of origin: Turkey and Morocco; (9) social and health problems of migrant workers; (10) social and health care of Moroccan workers in Europe; (11) aspects of health care intervention in host countries; (12) advocating for migrants' health; (13) migrants' special needs in sexuality and family planning in Belgium and Germany; (14) health care and education aids for foreigners in the Netherlands; (15) mental health of migrants; and (16) psycho-social problems of migrants. The book concludes with recommendations for researchers and practitioners.


Bollini (1992) who studied the policy regarding Migration and Health in seven industrialized countries (France, United Kingdom, Switzerland, Italy, Sweden, United States and Canada) already indicated that these countries can be divided into two groups: those which have a passive attitude, that is, which expect immigrants to adapt to the health system designed for the native population (Italy, France, Switzerland and the United States); and those which have acknowledged the health problems posed by immigrant groups and who have actively tried to provide alternative solutions, for instance by providing interpreter services during medical encounters (United Kingdom, Sweden and Canada).


Abstract--The paper reviews the available evidence on access to health care and two health outcomes, perinatal mortality and accident/disability, for migrant and ethnic minorities in selected receiving industrialized countries. The health of these communities is analyzed using the entitlement approach, which considers health as the product of both the individual's private endowments and the social environment he or she faces. Migrants, especially first and second generations, and ethnic minorities often have reduced entitlements in receiving societies. Not only are they exposed to poor working and living conditions, which are per se determinants of poor health, but they also have reduced access to health care for a number of political, administrative and cultural reasons which are not necessarily present for the native population.

The paper argues that the higher rates of perinatal mortality and accidents/disability observed in many migrant groups compared to the native population are linked to their lower entitlements in the receiving societies. Policies aimed at reducing such health gaps need to be accompanied by a more general effort to reduce inequalities and to promote full participation of these groups in the mainstream of society.
http://ceris.metropolis.net/Virtual%20Library/other/reitz1/reitz4.html

From the Summary: This review of nearly 400 publications from Canada, the US, Britain and Australia has identified a large number of studies supporting the conclusion that very often, recent immigrant groups experience low rates of utilization of many important social and health services, despite evidence of significant need. The barriers most often identified include those related to language, lack of information about services, cultural patterns of help-seeking, lack of cultural sensitivity by service providers, financial barriers, and lack of service availability.


Contains country reports on Belgium, France, Germany, The Netherlands, Spain, Sweden, and the UK as well as other chapters.


Summary of a review requested by the European Commission in 1997.

Abstract - The paper gives a brief overview of a wide spectrum of health issues and problems, ranging from communicable disease to mental health and family formation, which affect migrants and host countries.


First report of the project ‘Salute per Tutti’ – ‘Health for all, all in health’ – European experiences and strategies against social exclusion of immigrant people by health care services’.

Countries covered: Italy, Spain, Sweden, Belgium and the Netherlands


Abstract: This paper offers an examination of mental health services for migrant groups in a number of European countries. It draws on a range of recent studies to highlight some of the key and emerging issues in relation to the provision of mental health services within an increasingly multi-ethnic and multicultural Europe. The results of a preliminary mapping exercise of mental health services for migrant groups are presented and their broader implications are considered. The aim of the mapping exercise was to collect and examine information on mental health services for migrant groups against a backdrop of broad policy developments in the mental health field and
the emergence of multicultural approaches in public policy. The results of a questionnaire survey of service providers in 16 European countries are summarised, with special attention to three of the participating countries, Sweden, the Netherlands and Spain. The information from it is placed in a context of current research in the field of race, culture and mental health. In examining the results of the preliminary mapping exercise, key areas for policy development and service provision are identified and an agenda for future research in this area is suggested.

Available online at www.ercomer.eu/ingleby
Contains country reports on the UK, Netherlands, Spain and Portugal, also some material on Canada and Australia.
The results of this project are also summarised in:

(Report not confined to Europe)
From the introduction:
This report is a comprehensive review of primary literature on internal and international migration and health. It is the result of searches using five on-line databases, a list of health and migration related keywords, and strict inclusion and exclusion criteria (see section 2).
These searches produced 362 papers, of which 136 papers met the criteria and were included in the report (see section 5). These papers were summarised and separated into internal migration, and four subgroups within international migration: ‘all cause and cardiovascular mortality’, ‘cancer mortality’, ‘mental health’, and ‘morbidity, risk factors and anthropometry’.
The review investigates the morbidity or mortality rates of many immigrant groups from around the world moving between different countries or within a country. Although patterns and profiles vary for different immigrant groups in different countries, general themes were apparent.

www.salutepertutti.org
Research report of “Partners for Health, Phase II”: Combating discrimination in health care.
Extract: A qualitative study on experiences of discrimination and exclusion has been done in the countries of the five participants of the project: Italy, Spain, Sweden, Belgium and the Netherlands. The history of migration, the origin of migrants, the focus of research on migrants and health care in general and development of health care for migrants differ to a large extent between the five countries. Therefore, it was decided to focus on migrant groups that belong to the field of expertise of the participants in the project, or that were determined as groups, which have
clear problems of accessibility of health care. In Italy the research population was migrant women, with a focus on reproductive health. In Spain the research focused on migrant users of primary health care and hospitals. In Sweden the researchers focused on women from the Middle East and Somalia. In the Netherlands research was done in mental health care, in particular focused on Cape Verdean migrants. In Belgium, the researchers focused on mental health care for Turkish women with somatic complaints.


This publication presents an overview of evidence gathered from 52 good practices that reveal how the health field can foster social inclusion. The largest numbers of good practices focus on ethnic minorities residing in the country for a significant period of time, or illegal or newly arrived migrants.


Contains an overview of theoretical issues as well as country reports on The Netherlands, Switzerland, Greece and Portugal.


General overview of health problems and policy issues associated with migration. Not confined to Europe. Unfortunately, the paper does not include a list of references.


Very extensive report, containing a range of information on almost all European countries. Some are covered extensively, others only in passing.


Contains reports on Austria, Belgium, France, Germany, Hungary, Italy, Netherlands, Portugal, Spain, Sweden, United Kingdom


Three countries’ migrant health policies were described, revealing considerable differences. The issue of undocumented migrants and their access to health care systems in those three countries was also addressed.


This report was prepared in the framework of the Conference “Health and Migrations in the EU” (Lisbon, 27-28 September 2007) and aimed at presenting a collection of existing Good Practices in the field of health and migration, including practices in the public sector as well as many provided by non-governmental organizations. The first part of the report analyses common aspects of Good Practices, such as the methodology, the models and aims, and provides a critical overview of the limitations of Good Practices, highlighting the importance of structural change. In conclusion, some national cases of health policies are analysed to illustrate the diversity within the EU, and some recommendations are presented.

The Annexes present the collection of Good Practices divided into chapters: Transnational cases, country cases and Portugal.


The report compares data from Finland, Germany, Greece, Netherlands, Poland, Spain, Romania, and United Kingdom. Access and quality of care is examined for the following vulnerable groups:
- Migrants, asylum seekers and illegal immigrants
- Older people with functional limitations
- People with mental health problems.
The report begins with an in-depth analysis of the factors which may undermine access and service quality, and shows how these factors may affect each group in turn, using data from the 8 countries surveyed.

More detailed country reports are in preparation (see http://www.eurocentre.org/detail.php?xml_id=866 )


Starting with 2008, the FRA includes health care as one of the areas covered in its annual report on discrimination in the 27 EU Member States. Data on discrimination and barriers to access in health care is collected by the FRA’s 27 RAXEN National Focal Points (NFPs). Attention is also paid to the efforts that have been made in each Member State improve the situation for migrants and ethnic minorities. In this report, Roma and undocumented migrants are identified as the groups most likely to experience barriers to access to health care.
2. Current projects

A. Projects co-funded by DG SANCO (12)

MIGHEALTHNET Information network on good practices in health care for migrants and minorities in Europe.

www.mighealth.net

The MIGHEALTHNET project aims to stimulate the exchange of knowledge on migrant and minority health through the development of interactive data bases in each of the participating countries. These 'wikis' will contain the following sorts of data:

1. Background information concerning migrant and minority populations
2. The state of health of migrants and minorities
3. The health care system and the entitlement of migrants and minorities to health care
4. Accessibility of health care
5. Quality of care: ‘good practices’ developed to improve the matching of service provisions to the needs of migrants and minorities
6. Centres of expertise, general reports and policy documents, journals, training programmes, E-mail groups etc.

By facilitating the transfer of knowledge and expertise and stimulating network formation within and between European countries, the project hopes to further the development of good practices concerned with the health of migrants and minorities.

Countries participating: Belgium, Bulgaria, Czech Republic, Denmark, Germany, Greece, Hungary, Lithuania, Netherlands, Norway, Poland, Portugal, Romania, Sweden, Switzerland, Turkey and the United Kingdom.

Healthcare in "NowHereLand": Improving services for undocumented migrants in the EU.

http://www.ausl.re.it/HPH/FrontEnd/Home/Default.aspx?channel_id=58

This project aims at improving the level of health protection for the people of Europe by addressing migrants’ and immigrants’ access, quality and appropriateness of health and social services as important wider determinants for health, focusing on healthcare services for undocumented migrants (UDMs) as an especially vulnerable group, an increasing public health risk and a group providing difficulties for healthcare providers and health policy.

MEHO: Migrant and Ethnic Health Observatory.

http://www.meho.eu.com/

The main objective of MEHO is to develop indicators to monitor the health status of immigrant/ethnic minority groups in Europe. Therefore existing health related databases and surveys will be used and an European network of epidemiological observatories on migrants’ health will be established. This network will generate an European overview of comparable and exchangeable data on sociodemographic and health profile of immigrant/ethnic minority groups for selected health problems. Specific attention will be paid to the conceptual, methodological, ethical and practical issue of identifying immigrants and ethnic minorities in health databases and to the assessment of valid comparisons between these groups within countries and between countries.

We do not only include immigrant/ethnic minority groups in Western Europe but also the Roma population in Central and Eastern Europe.

We focus on five critical health areas for which we know already that ethnic specific health data is available: mortality, cardiovascular diseases and diabetes, infectious diseases, cancer, self-perceived health and health care use.
EUGATE: Best practice in access, quality and appropriateness of health services for immigrants in Europe.
http://mighealth.net/eu/index.php/EUGATE_contact_details_and_associated_partners

EUGATE (in 16 countries) aims to provide a European-wide definition of what is meant by ‘migrant’, explore legislation, policies and funding arrangements relating to migrants and minorities, investigate organisation and utilisation of services, describe evaluation and monitoring methods, and identify and compare models of best practice. In addition they are going to develop a 'tool kit' for improving access, quality and appropriateness of health and social services for migrants and ethnic minorities, as well as creating a multi-lingual, searchable website on best practice.

Health and migrations in the European Union
http://mighealth.net/eu/index.php/Health_and_migrations_in_the_EU_-_contact_details

The project aims to:
- improve the knowledge about the health status of immigrants coming from third countries, and its health determinants considering: demographic dynamics of migratory processes and their impacts, the specific political and legal frameworks at the national and international levels;
- identify best practices about immigrants’ access to healthcare (including health promotion, prevention and healthcare services);
- contribute to the definition of health policies and strategies that could be implemented both at the EU and Member State level, aiming at better migrants’ integration.

AMAC: Assisting migrants and communities: analysis of social determinants of health and health inequalities.
http://mighealth.net/eu/index.php/AMAC_contact_details

The project aims to consolidate the outputs of the vigorous 2007 migration health agenda in Europe and promote multi-stakeholder dialogue and engagement to fight health inequalities linked to migration. In parallel to this policy dialogue, the project will review, through seven background papers developed around three workshops, priority health issues affecting migrants across Europe such as mental health, maternal and child care, care for the elderly, as well as legislation and research in the field, bioethics and training for health professionals. In the “Health in All Policies” spirit, the AMAC initiative will also expand the dialogue beyond health professionals to include stakeholders in linked fields such as education, social affairs and interior. The project will conclude with a final EU-level consultation in May-June 2009 and a final report including the consultation’s conclusions and recommendations.

AVERROES Network: Improving access to health care for asylum seekers and undocumented migrants in the EU.
http://mighealth.net/eu/index.php/AVERROES_contact_details_and_associated_partners

The project contributes to enhancing the EU general population’s health, by improving asylum seekers’ and undocumented migrants’ access to healthcare. The project is also strategically relevant since Portugal has decided to
focus on “Migration and Health” issues, during its EU presidency. Actually, the Portuguese National Institute of Health has even recently selected the MdM European Network observatory and advocacy project as a best practise for its “Health and Migration” programme, funded by the Community Action Field of Public Health. To achieve its objective the project proposes to create an NGO network covering 19 member states, that will carry out research, field surveys, awareness raising and advocacy activities at national and EU levels. By doing this, the network intends to sensitize the main stakeholders to the project’s advocacy message as well as to convince and encourage policy-makers to favour the elaboration of binding community regulations to improve asylum seekers’ and undocumented migrants’ access to healthcare in the EU.

**Health and the Roma community: analysis of the situation in Europe.**
http://mighealth.net/eu/index.php/Health_and_the_Roma_community_-_contact_details_and_associated_partners

The project will:
- contribute to the reduction of health inequalities affecting the Roma community in Europe; obtain reliable and objective data about the social/health situation of the Roma population and the use made of healthcare resources available for the mainstream society;
- identify factors considered vital in improving the Roma situation and promote equity;
- promote synergies between public/private sphere (health centres, hospitals, social organisations, public administrations, etc.).

**TAMPEP - European Network for HIV/STI prevention and Health Promotion among Migrant Sex Workers**
http://www.tampep.com/

A Network of community-based service providers, public health and social services cooperating in 26 European countries. Main objective: To reduce HIV vulnerability of migrant and mobile sex workers across Europe.

**Aids & Mobility Europe (A&M) 2007-2010**
http://www.aidsmobility.org/index.cfm

A network for the support of European organisations that provide HIV/AIDS prevention and care to mobile and migrant populations. A&M pays special attention to young migrant people.

**PHBLM**

**Increasing Public Health Safety Alongside the New Eastern European Border Line**

Funded under the European Commission’s 2006 Public Health Programme (PHP), this IOM project aims to increase the public health safety of all EU member states, build the capacity for border management, and support appropriate health provision to border guards and healthcare to migrants entering the European Union border area as a fundamental human right. The project is being implemented with the University of Pécs in Hungary, in partnership with the governments of targeted countries, and is collaborating with the European Centre for Disease Prevention and Control (ECDC), Frontex, and the World Health Organization (WHO).
Running from June 2007 to December 2009, the PHBLM project includes four main components: situational analysis of the current public health conditions and border management procedures; development and regional testing of a training module; development of minimum public health standards and a proposal for structural changes; and dissemination of the project results. The project focuses on the eastern external border of the enlarged European Union where the Schengen criteria are currently being implemented (Hungary, Poland and Slovakia). Romania will also be included in the preliminary situation analysis as well as in the regional testing of the training module.

Further information: contact Roumyana Petrova-Benedict at rpetrovabenedict@iom.int

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**Healthy Inclusion**

http://www.wrk.at/cn/?xml_file=frkeng.xml&new_start=/cn/was_wir_tun/frk/startseite.php

(See "Research focus points / Health promotion and prevention").

The project is concerned with improving the access of migrants to health promotion interventions. The project will:

- Provide information about migrants’ perceived barriers for participating in health promotion interventions as well as about facilitating factors
- Provide examples of good practice and suggested means of enhancing migrants' participation in health promotion interventions
- Develop specific recommendations on how health promotion interventions at the community level can be adapted to better meet the needs of migrants.

The results will be disseminated to the health promotion community and to policy makers in each partner country.

Further information: contact Barbara Kuss at barbara.kuss@w.roteskreuz.at
B. Projects (co-)funded by other bodies (9)

HOME – Health and Social Care for Migrants and Ethnic Minorities in Europe
COST Action IS0603
http://www.cost.esf.org/index.php?id=233&action_number=IS0603

The recent increase in the numbers of migrants in Europe has generated a growing volume of research on their state of health and the need to adapt care services to their needs. Scientific progress in this field, however, is held up by a lack of interdisciplinary and international collaboration. Moreover, the addition of a cross-national perspective can yield new insights into the causes of ill-health and can further the exchange of good practices. In Southern, Central and Eastern European Countries, work on migration and health is in particular need of strengthening and encouragement. This Action will bring together an international group of experts to consolidate and review work carried out so far, identify blind spots and persistent problems, and recommend ways forward. Its three Working Groups will be concerned with social and policy factors, migrants’ state of health, and improvements in service delivery. The Action will produce ‘state of the art’ reports on the most urgent themes and will organise workshops, conferences, joint publications and training activities to discuss and disseminate its findings.

Sixth Framework Network of Excellence CINEFOGO
http://www.cinefogo.org/

Within this project, the University of Trento is investigating migrant health. They have a project "Giving new subjects a voice. Cultural diversity in the health-care system", financed by the Volkswagen Foundation and directed by Prof. Patrizia Nanz.
http://www4.soc.unitn.it:8080/dart/content/e1366/index_eng.html

BOMEME (Birth Outcomes of Ethnic Minorities in Europe).
Coordinated by Anders Hjern, Adjunct Professor of Paediatric Epidemiology, National Board of Health and Welfare and Karolinska Institutet, Stockholm, Sweden (anders.hjern@socialstyrelsen.se).

IMISCOE (International migration, Integration and Social Cohesion in Europe).
http://www.imiscoe.org/

Cluster B5 has a group working on Migrant Health. Contact David Ingleby (j.d.ingleby@uu.nl).
WHO-HPH task force on migrant-friendly and culturally competent healthcare (MFCCH).
http://www.ausl.re.it/HPH/FrontEnd/Home/Default.aspx?channel_id=38

The Task Force on Migrant Friendly and Culturally Competent Health Care is established within the international HPH network with a specific mandate for coordination assigned to the HPH regional network of Emilia-Romagna (Italy) by the General Assembly and the Governance Board of the international HPH network. The provider is the Health Authority of Reggio Emilia, which is the coordinating institution of the regional HPH network of Emilia Romagna.

EUROMED European-Mediterranean Network on Migration and Mental Health.

Key areas of work:
1. Improvement of mental health services through research and training
3. Participation in policy development and future planning in issues related to migration and mental health.

PROMO

This project aims to review policies, services and best practice in the promotion of mental health and prevention of mental ill health of socially marginalised groups in Europe. The project is coordinated by Queen Mary University, London. (Prof. Stefan Priebe)

PROMO assesses best care for ‘socially marginalised groups’ with mental health disorders across Europe. The marginalised groups that are specifically studied are a) the long-term unemployed, b) homeless people, c) prostitutes, d) ‘travellers’, e) illegal immigrants, and f) asylum seekers and refugees. The study will identify relevant legislation and policies, assess all available health care and social services in representative areas, and evaluate the overall quality of care for each group. The ambitious aim is to develop recommendations for the European Commission for how care for socially marginalised groups is best organised and delivered.

The project has partners in 14 European countries, is funded by the European Commission and will last three years. It is coordinated in the Unit for Social & Community Psychiatry in Newham. The Principal Investigator is Prof. Stefan Priebe, and the project manager is Donna Wright (D.J.Wright@qmul.ac.uk).

EU Partnerships to Reduce Migrants' Vulnerability to HIV Infection

Project funded by the Government of Portugal in 2007 and managed by IOM HQ and Brussels.


Active Ageing of Migrant Elders across Europe
http://www.aamee.eu/

Project promoted by the Ministry of Intergenerational Affairs, Family, Women and Integration of the State of North Rhine-Westphalia, Germany, and the Directorate General for Employment, Social Affairs and Equal Opportunities, with a grant from the EU.

The AAMEE project focuses on the promotion of active ageing and social, cultural and economic integration of migrant and minority ethnic elders, emphasising volunteer activities and the emergence of new culturally sensitive products and services in the fields of, for instance, housing, care, education, leisure, culture and marketing. This will be done on the basis of a mixture of practical and scientific activities and a variety of approaches.

3. Projects on Roma health (5)

Two projects listed under (1) or (2) are listed again here

MEHO: Migrant and Ethnic Health Observatory (DG SANCO)

(The main project is described under category 1)

- Objectives:
  - to define the most important criteria identifying Roma health
  - to set an existing databases with available information on Roam health issues in CEE
  - to collect and analyze data on health of Roma comparison with the indigenous majority population

- Description of work:
  - to summarize outputs from all accessible health data characterizing Roma health group outputs, we suppose the data are very scarce and limited and needs to be summarized
  - to establish a database of organization and scientists dealing with Roma health issues in CEE, considering governmental, academic research, private and NGO sectors
  - to develop ethical issues particularly in reproductive health data
  - to support all kinds of structural; and targeted research dealing with Roma health

  - to develop guidelines how to receive or extract data about Roma from routine health statistics
  - focus on at least 3 countries
  - focus at least on 3 critical health areas cardiovascular, infectious disease, reproductive health

Health and the Roma community: analysis of the situation in Europe. (DG SANCO)

Listed above under category 1.
Open Society Institute (OSI) - Roma Health Project
http://www.soros.org/initiatives/health/focus/roma

The OSI Roma Health Project (RHP) works to promote equal access of Roma to appropriate and quality health care services. The project focuses on the protection of the rights of the Roma population in the health care setting, by promoting involvement of Roma communities in advocating for access to health services, addressing discrimination against Roma in the health sector and raising visibility around the obstacles impeding access health care. RHP is working to affect the development of sound public health policies targeting Roma, and combating the perpetuation of myths and stereotypes about Roma communities and health.

Building Healthy Roma Communities Project (IOM)

Funded by the Government of Belgium, this IOM project was completed in February 2008 and targeted Hungary, Poland, Romania and Slovakia. The twelve-month, second phase aimed for the best possible utilisation of the research and training from phase one, through capacity-building of professionals and stakeholders working with Roma communities, specifically in the areas of health and social affairs.

The project implemented a country-level training course in each of the target countries during 2007. These courses were led by the trainers certified during the project’s first phase and supported by the project’s Expert Team. A pilot community centre (Equality Support Unit) was also established in Romania, whose goal is to provide assistance for the Roma community while also encouraging self help, health promotion and awareness, and ensuring dissemination of information.

Powerpoint presentation on the whole project: http://mighealth.net/eu/images/3/3a/Iom22.pdf

Euroma - European Network on Social Inclusion and Roma under the Structural Funds
http://www.euromanet.eu/

In January 2008 twelve countries of the EU together with a wide representation of the European Commission launched in Seville (Spain) a transnational Network aiming at the promotion of the use of Structural Funds (SF) to enhance the effectiveness of policies targeting the Roma and their social inclusion.