The European health report 2012:
charting the way to well-being
Executive summary
This executive summary contains an overview of *The European Health Report 2012: charting the way to well-being*, the WHO Regional Office for Europe’s flagship publication issued every three years. Details are outlined in four sections, addressing health status in Europe, European targets for health and well-being, the case for measuring well-being and an agenda to address measurement challenges.

**Keywords**

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Overview

As part of its mandate to monitor and report on the health of nearly 900 million people in the WHO European Region, the WHO Regional Office for Europe publishes the European health report, its flagship publication, every three years. The report has three purposes. First, its timing offers an excellent opportunity to provide policy-makers and public health professionals with the epidemiological evidence base underpinning the strategic objectives, targets and priorities of the new European policy framework for health, Health 2020. Second, the report analyses social, economic and environmental determinants of health and puts well-being on the agenda as a marker of social progress for the European Region. Finally, it identifies key challenges for health measurement for the coming decade and sets out a collaborative agenda: to collect, analyse, and make use of health data Region-wide.

The report has four sections. The first introduces the report by describing health status and trends in Europe at present, covering 53 countries and highlighting individual countries and subsets of countries. Topics covered include demographic trends, life expectancy, mortality, causes of death, the burden of disease, risk factors, and social determinants of and inequalities in health, including health systems. Key messages from this analysis show that people across Europe are living longer, but with changing patterns of disease burden and increasing inequalities in health and its determinants. Life expectancy has increased to more than 76 years for men and women combined, mainly as a result of decreases in certain causes of death and improvements in the prevalence of risk factors and socioeconomic and living conditions. Yet these improvements and the conditions that foster them have not been shared equally within and between countries; substantial differences persist and are increasing in many instances.

The second section presents the baseline for the overarching targets selected to monitor progress for the new European health policy, Health 2020. Specifically, it documents the intense process through which the WHO Regional Office for Europe consulted with representatives of Member States and the work of several expert groups, leading to the identification of six overarching targets to be achieved by 2020. The targets are set at the regional level and are quantifiable, with indicators to mark progress towards 2020. The use of targets builds on previous European efforts such as those of the Health for All and HEALTH21 strategies. With almost 30 years of experience in target setting, the Regional Office and the Region have learned valuable lessons that can benefit Member States in their efforts to set national-level targets.

The third section recognizes that a core aspect of Health 2020 is improving population well-being, specifically in the context of health. This offers a platform to develop a common agenda across sectors and stakeholders, including a whole-of-government approach. This is based on the WHO Constitution’s definition of health: a “state of complete physical, mental and social well-being and not merely the absence of disease or infirmity”. Nevertheless, WHO has neither measured nor reported on well-being. To incorporate well-being in the work of WHO and to develop a European target and relevant indicators by 2013, the section
Executive summary

of the report lays out issues and processes needed to develop a common concept and measurement approach to well-being that allows valid and comparative data collection and yields information useful to policy-makers and in programme evaluation.

The fourth section concludes the report by identifying important challenges in measuring progress towards Health 2020 and by outlining an agenda to overcome them, with WHO working with partners and Member States. This also reflects the Regional Office’s support to countries in enhancing their collection, analysis and reporting of health information, as well as its work with the European Commission (EC) and Organisation for Economic Co-operation and Development (OECD) to set up a common, integrated health information system for the European Region, covering all 53 Member States.

1. Where we are: health status in Europe – the case for Health 2020

Demographic trends

The WHO European Region is undergoing important demographic and epidemiological changes that are shaping the needs for health promotion, disease prevention and care in the future. Such transitions are occurring at varying speed and intensity for different country groups and populations, however, creating new challenges and a mosaic of health situations that requires specific approaches. This highlights some of the major challenges for health in Europe that the Health 2020 policy addresses. The information provided in this section identifies key elements on which the policy is based, sets the agenda for action and provides a baseline for monitoring progress towards defined priorities and areas for policy action.

The population of the 53 countries in the European Region reached nearly 900 million in 2010. Decreasing fertility rates across the Region mean that population growth will soon plateau. Moreover, the population is ageing rapidly, with projections estimating that more than 25% of the total population in the Region will be aged over 65 years by 2050.

An estimated 73 million migrants (with women representing 52%) live in the European Region, accounting for nearly 8% of the total population. Overall, this population inflow, mostly towards EU countries, comprises an increase of 5 million in the migrant population since 2005 and accounts for nearly 70% of population growth during this period.

The proportion of the population of the European Region living in urban areas reached nearly 70% in 2010 and is expected to exceed 80% by 2045. As a consequence, people are being exposed to different risks and determinants of health.

Life expectancy

Life expectancy at birth increased by 5 years after 1980, reaching 76 years in 2010. Average life expectancy at birth differs across countries, with the highest level at 82.2 years, and the lowest at 68.7 years in 2010, a gap of 13.5 years. In 2010, women reached an average of 80 years, while men lived 72.5 years on average, a gap of 7.5 years. Inequalities in life
expectancy across countries were larger for males than females, particularly after 1993. Projections suggest that life expectancy in the European Region will increase to nearly 81 years by 2050, a similar pace to that in 1980–2010. Data from individual countries show large inequalities in life expectancy and years gained, especially when analysed by sex or by subregions.

More people in the Europe Region are living over the age of 65 years: an estimated 15% in 2010. Their remaining life expectancy after that age is 15.5 years on average. By 2050, this age group is projected to represent more than 25% of the Region’s total population.

**Mortality**

Overall mortality from all causes of death continues to decline in the European Region, reaching an age-standardized death rate (SDR) of 813 deaths per 100 000 population in 2010. Mortality trends show wide variations between country groups across the Region. Differences in mortality patterns in the Region tend to be low or very low from the early years of life up to young adulthood and then rapidly increase in older age groups. Child mortality indicators in the European Region are the lowest in the world and have continued to decline since 1990. The average reported infant mortality rate in 2010 was 7.3 per 1000 live births, following a 53% reduction between 1990 and 2010.

The European Region’s maternal mortality ratio for 2010 was 13.3 maternal deaths per 100 000 live births, showing a 50% reduction since 1990 with much variation across countries. Women’s level of education and socioeconomic status affect maternal mortality. This highlights the importance of policies and interventions addressing gender and other social determinants of health, including health system functioning.

As the European population ages, mortality trends help to anticipate some of the challenges that health systems will face and some of the adjustments needed to address them. All-cause mortality among people aged over 65 years decreased in the European Region as a whole, reaching its lowest rate, 4549 per 100 000, in 2010: a 25% reduction since 1980.

**Causes of death**

Noncommunicable diseases produce the largest share of mortality, accounting for about 80% of deaths in the European Region in 2009. Cardiovascular diseases remain the leading cause of mortality, accounting for nearly 50% of all deaths, followed by cancer, accounting for nearly 20%. External causes of injuries and poisoning are responsible for 9% of all deaths. Distribution patterns by subregion have changed over time.

The main cardiovascular diseases are ischaemic heart and cerebrovascular diseases, which together account for 35% of all deaths in Europe. Nevertheless, they vary widely between countries by age, sex and the distribution of their determinants. Risk factors are described below.

According to the European Health for All database, overall cancer incidence and mortality rates in the European Region are 379 cases and 168 deaths per 100 000, respectively. Incidence has increased by 32% and mortality decreased by 10% since the mid-1980s.
Reductions in mortality rate are observed with cancers for which diagnosis, prevention and health care have improved. Among men, lung, colon, stomach and prostate cancer account for nearly 50% of cancer deaths, while breast, lung, stomach, colon, cervical and ovarian cancer account for 60% of deaths among women.

Monitoring of mortality trends for the main external causes provides additional information for future projections, and insight into environmental safety conditions and some harmful behaviours in the population. For example, although levels differ, suicide rates have decreased by 25–40% in all the subregions of Europe, following an increase in the mid-1990s. Recent trends indicate, however, a slower decrease coinciding with the economic downturn since 2008. Mortality from transport accidents provides information on road safety, including infrastructure conditions, protective measures and regulation. The overall trends for the European Region show a 50% decrease since 1990, to a rate of 10 per 100 000 in 2010. This situation may be related to a decrease in road traffic accidents, particularly those involving alcohol.

Communicable diseases are less common in the European Region than in other parts of the world. Concern focuses mainly on tuberculosis (TB), HIV/AIDS, other sexually transmitted diseases and hepatitis viral infections. Nevertheless, recent poliomyelitis, rubella and measles epidemics in Europe have re-emphasized the need to sustain or improve public health activities, such as disease surveillance and preventive activities, including health promotion and immunization. Mortality rates for TB have decreased in the European Region after increases in the 1990s, reaching a rate of 6 per 100 000 in 2010: a 30% reduction during this period. Co-infection with HIV presents further challenges to reducing mortality from tuberculosis, particularly in areas of eastern Europe where both infections coexist and where multidrug (including antimicrobial) resistance is present. Patterns of HIV incidence differ among groups of countries. AIDS incidence is decreasing in all subregions, indicating the importance of effective treatment.

**Premature mortality**

More than 70% of mortality occurs at ages over 65 years, when disease processes have been underway for several years. Monitoring of premature mortality (deaths before the age of 65) provides useful information for developing public health policy, programmes and interventions to delay disease and the onset of disability. The gap between men and women shows a 50% excess risk for men reaching 60 years, and the risk for men is twice that for women by the age of 65. Nevertheless, rapid population ageing necessitates the monitoring of mortality, and particularly morbidity and disability, in the groups aged over 65.

Cancer has replaced cardiovascular diseases as the foremost cause of premature death in 28 of the 53 countries in the European Region, as a result of the sustained decrease in cardiovascular mortality and an associated increase in life expectancy, and the long latency period in developing cancer. At the country level, the highest rates of premature mortality from lung cancer are found in central Europe (particularly Hungary, Serbia and Poland): over 25 per 100 000. This pattern is associated with the prevalence of tobacco smoking, which in turn is determined by the affordability of tobacco products, marketing strategies and
permissive public health policies. The premature mortality rates for breast cancer in the Region have decreased over the past decade to 14 per 100 000: a reduction of 21% since the peak in the mid-1990s.

Rates of premature mortality from respiratory diseases in the European Region have steadily decreased by 40% since the mid-1990s, to 16 per 100 000 in 2010. Mortality from this cause takes its toll on two distinct populations: children and older people. The main specific causes of death are chronic obstructive pulmonary disease (COPD), asthma, pneumonia and influenza, all closely associated with conditions and exposures in the outdoor and indoor environments. Interventions to prevent them are known, but some require intersectoral action.

Premature mortality from diseases of the digestive system in the European Region showed an increasing trend from 1990 to 2010, when the rate reached 25 per 100 000: a 30% increase. In this group, chronic liver disease and cirrhosis, and ulcers of the stomach and duodenum are the main causes of death and are mainly associated with harmful intake of alcohol and processed foods. Chronic liver disease and cirrhosis have also been associated with a range of viral causes, such as hepatitis B and C infections, and toxins and drug misuse. Alcohol abuse, however, particularly when it is heavy and sustained, probably makes the largest contribution.

Diabetes is a major public health problem in Europe because of its direct and indirect effects – ranging from renal, neurological and ophthalmological microvascular damage to vascular damage of the limbs, brain and heart – with various severe consequences. Premature mortality from diabetes in the European Region showed a 25% reduction from 1995 to 2010, when it reached a rate of 4 per 100 000. Identifying diabetes as a cause of death poses some difficulties, so mortality from this cause may be underestimated.

### Burden of disease

The distribution of the European Region’s total burden of disease (combining mortality, morbidity and disability) for 2004, the latest year with available data by mid-2012, shows a range of 10–28% in estimated disability-adjusted life years (DALYs) lost per country population. This represents an almost threefold gap between the most and least favourable country situations. Total DALYs have been attributed to different leading risk factors across the European Region. This information enables the identification of the most important areas for interventions – such as nutrition, physical activity and reduction of addictive substance consumption – mainly to reduce obesity, high cholesterol and blood pressure, and alcohol and tobacco use. Such interventions require intersectoral participation and the use of various cost-effective strategies.

### Risk factors

Understanding the underlying causes of disease, including risk factors and socioeconomic and health system determinants, is important for disease prevention and control and health promotion. Among the major groups of diseases causing high mortality, morbidity and disability – such as cardiovascular diseases, cancer, external causes of death, respiratory system diseases and digestive diseases, as noted above – the two main risk factors to tackle
are tobacco smoking and harmful alcohol consumption. Prevalence and consumption levels of these two factors among the European population remain high across the Region, in spite of the availability of knowledge and technology to control them.

Tobacco smoking prevalence in the European Region reached 27% of the population aged 15 years and older around 2008, but has gradually decreased towards 25%, particularly among men. Alcohol consumption is another factor that determines the frequency of health problems, and according to WHO estimates it accounts for nearly 6.5% of all deaths in Europe. In addition to volume, the type of alcohol consumed and patterns of binge drinking are also relevant, due to their potential effects on health. Unfortunately, several countries in the Region have not reported recent data.

Differential access or exposure to diverse environmental factors over the course of a lifetime are known to determine the occurrence of major health problems, including cardiovascular, respiratory and digestive diseases, cancer and external causes of death. Access to clean water and hygienic sanitation services, housing conditions, road safety, air quality, the work environment and exposure to extreme climate conditions directly or indirectly contribute to shaping the health profile of the European population.

Social determinants and health inequalities, including health systems

The WHO Commission on Social Determinants of Health defines health equity as “the absence of unfair and avoidable or remediable differences in health among population groups”. Global evidence suggests that at least 25% of health inequalities (differences found within a country’s population) are associated with a lack of access to effective health services, and this percentage increases when adding differential access to basic public health interventions such as access to safe water. The social determinants of health contribute to another 50% of health inequalities, covering political, socioeconomic and environmental factors. These differences are unfair, so health inequities make up the bulk of health inequalities. These unfair inequalities are also called “the causes of the causes” of health inequities, reflecting their fundamental influence on disease causation and the systematic social patterning of health outcomes, including life expectancy. Health inequities are a major concern in Europe, given the widening gap and social gradient in life expectancy between and within many countries. As one of the most important social determinants, gender norms and relations continue to shape the way health systems are organized and services delivered, often to the detriment of girls and women or people who do not fit within accepted gender roles.

Prerequisites to tackle health inequities include:

a. commitment to ensuring that all people have equal opportunities to improve or maintain their health;

b. assessment of health policies and programmes in relation to their specific effects on inequities in health, from inputs to outcomes, using disaggregated data on different subpopulations;
c. understanding of the pathways from social determinants to differential exposures, vulnerabilities, interaction with the health system, and differential health outcomes; and
d. identification of entry points and actions for change involving participation of the community and other stakeholders, drawing on the evidence base on what can be done to reduce health inequities.

Action within the health sector or health system can include strategies to reshape existing programmes to enhance equitable access to care and address processes that exclude people who are disadvantaged or vulnerable. Action on the social determinants of health often requires that multiple sectors align their objectives, work together and achieve multiple social goals. For example, early child development programmes are associated with better health, education and nutritional outcomes for children and across the life-course, as well as improved incomes and contribution to greater social cohesion.

Distributions of wealth, education and occupation status – within or across countries – reflect how stratified a population is, and these distributions are also important social determinants of health within and across countries. The European Region is characterized overall as one of the wealthiest in the world, where the annual average income per capita was about US$ 24 000 in 2009. Although increasing since 1990, per capita income levels within the Region are still highly inequitable, ranging from about US$ 715 to US$ 105 000, with nine countries in the Region with levels below US$ 5000. The recent economic downturn affecting the European Region increased the average unemployment level to 8.7% of the economically active population in 2009, a reversal of the more optimistic trend over the previous decade. At the country level, the gap between low and high unemployment shows a thirty-five-fold difference.

The environment represents another important health determinant. Recent assessments of the contribution of environmental factors to health have estimated that they may be responsible for as much as 13–20% of the burden of disease in Europe, depending on their mortality pattern classification. WHO recently assessed the impact of environmental factors on health inequalities in the European Region.

An effective health system is a prerequisite for responding to the changing epidemiological situation and the health needs of the population through governance, health financing, human resources and institutions that provide services covering prevention, treatment and palliative care. Progressive financing of health systems and ensuring social protection for households from catastrophic expenditure are important components of universal coverage without financial risk. Direct or out-of-pocket (OOP) payments by households, as a percentage of total health expenditure, provide a good indicator of financial risk and the level of equity in financing. Data on this indicator are available from all 53 European Member States. Average OOP expenditure across the European Region stands at about 23%, but significant variation exists across countries, from a high of almost 80% to a low of just under 6%: a fourteen-fold gap. In 10 countries, OOP expenditure remains the main source of payments (about 50% or more). Global evidence shows that, when reliance on direct payments falls to less than 15–20% of total health expenditure, the incidence of financial catastrophe routinely falls to
negligible levels (with financial catastrophe defined as when households annually spend more than 40% of their income after food on paying for health services).

2. What we are aiming for: European targets for health and well-being

The strategic goals of Health 2020 are taking shape in collaboration with Member States. At its sixty-first session in September 2011, held in Baku, Azerbaijan, the WHO Regional Committee for Europe endorsed the proposals that Health 2020 will:

a. set out an action framework to accelerate attainment of better health and well-being for all;

b. be adaptable to the different realities that make up the Region; and

c. formulate regional targets for achievement by 2020.

The process of target setting has been informed by previous efforts and by participatory detailed discussion, written consultation and approval by representatives of governing bodies at each stage.

Previous experiences with target setting and monitoring

European Region targets were suggested as part of the first common health policy in the Region, the European strategy for attaining Health for All. In the early 1980s, the then 32 Member States in the WHO European Region debated European targets aligned to the new policy. The agreement on targets was a major undertaking, with more than 250 experts from across the Region working with the WHO Secretariat. More than 20 drafts and an in-depth consultative process with Member States resulted in the presentation of 82 targets for consideration by the governing bodies of the Region. A reduced set of 38 targets was unanimously adopted in 1984 at the thirty-fourth session of the Regional Committee for Europe in Copenhagen. The first European health policy was published, and included these 38 targets, along with 65 regional indicators to monitor and assess progress.

These European targets and indicators were reported on within the Health for All database. By 2012, the Health for All database had evolved to cover 53 countries and it is widely available and used across the Region. The database includes a selection of core health statistics covering basic demographics, health status, health determinants and risk factors, and health care resources, utilization and expenditure; these data are compiled from various sources, inform WHO atlases of health in the European Region, and are updated twice a year.

To reflect the changes that have taken place in the Region since the mid-1980s, the 38 targets were revised in 1991. The WHO Regional Office for Europe supported the implementation of the targets by responding to Member States’ requests and aligning the Organization’s budgets and programmatic activities to the target areas.

During the 1990s, major political, economic and social changes in the Region changed the European landscape. One result was a dramatic increase in the number of European Member
The European health report 2012

States, which now stands at 53. The Regional Office revisited its European health policy and regional targets. In 1998, a new set of 21 targets was identified within *HEALTH21: Health for All in the 21st century*. Adopted at the forty-eighth session of the Regional Committee in 1998, these targets were published in 1999. The focus of this second iteration remained on the construction of targets at the country and local levels, with no regional reporting.

The WHO European Region has approximately 30 years of experience in target setting as part of regional health policies and strategies, in the context of a Europe that has changed dramatically. Lessons learned include the following:

a. a broad consensus needs to be developed among stakeholders;

b. targets need to be limited to a manageable number;

c. any plan should be based on evidence of effectiveness;

d. targets need to be linked to resources; and

e. technical challenges remain that require collaboration with partners and technical support to countries that ask for it.

Experience in the Region has shown that setting targets and monitoring indicators can be a hugely motivating factor as countries collect and incorporate in their routine information systems the necessary data to inform health policy, even where in the past such data did not exist.

**Technical issues in selecting targets and indicators**

A well-organized mechanism is important to formulate specific, measurable, achievable, relevant and timely (SMART) targets. SMART targets are more likely to be accomplished than general goals. To arrive at measureable targets, concrete criteria for measuring progress must be established. For targets to be achievable, they must be realistic, while set against a defined time scale – in this case up to 2020 – with interim progress monitoring. Targets are considered relevant when they represent objectives towards which the policy is able to work. Every target, if achieved, should represent real progress that can be quantified.

Once a target area and potential indicators are identified, several approaches exist to select a target level. The counterfactual method identifies a biologically achievable or theoretical minimum or maximum value, which is compared to the existing situation based on available information. Trend analyses show where trends in rates can be used to arrive at a target in the future. Intervention study pooling and comparative risk assessments also provide methods of selecting target levels based on evidence of what works in different contexts and subpopulations. A monitoring framework and structured reporting are also needed, as well as a guide to interpret the indicators and targets. With these lessons in mind, extensive consultations with Member States and technical experts have taken place to develop regional targets in line with the development of Health 2020. The full report includes a template for the monitoring framework.
Consultation with representatives of European Member States

Three meetings of the European Health Policy Forum for High-level Government Officials took place that provided detailed discussions and input to the process to identify a limited number of European targets including technical criteria, alignment with Health 2020, and agreement on a framework and shortlist. Delegations from across the European Region met in March 2011 in Andorra, in November 2011 in Israel and in April 2012 in Belgium. A shortlist of six overarching European target areas was identified as a mechanism for accountability and solidarity across the Region, noting that Member States should be encouraged to develop their own national targets for health.

Member States also contributed to the technical deliberations, working closely with the WHO Secretariat, reflecting the proposal of the Standing Committee of the Regional Committee (SCRC) in May 2011. Representatives of the following Member States were nominated for this working group: Andorra (previous chair of SCRC), Poland, Sweden (current chair of SCRC), Turkey, Ukraine, the United Kingdom and the former Yugoslav Republic of Macedonia (former SCRC Chair). The group was chaired by a representative of a Member State (Sweden) with extensive experience in this area, and co-chaired by the WHO Regional Director for Europe. Each meeting of the group resulted in clear recommendations to narrow down a list of potential targets and indicators in line with the three broad areas identified as part of Health 2020:

a. burden of disease and risk factors;

b. healthy people, well-being and determinants; and

c. processes including governance and health systems, for wider consultation.

The WHO Secretariat collated inputs and recommendations on the process to set targets, as well as potential targets for inclusion, for the various consultations with Member States described above. All technical divisions at the WHO Regional Office for Europe submitted potential targets and monitoring indicators. From an initial list of 51 proposed targets, the SCRC working group recommended that 21 proposed targets should be retained in January 2012. This shortlist was subject to extensive written and face-to-face country consultation during February and March 2012, resulting in an initial framework of 16 potential targets and associated indicators, largely drawn from existing data reporting by countries; these were further reduced to 6 overarching or “headline” targets. The May 2012 meeting of the SCRC, held in Geneva, provided full support to the target work, further endorsed the six overarching targets, agreed that they will feature in all Health 2020 documents and confirmed that indicators will monitor progress and achievement by 2020.

Overarching targets

The sixty-second session of the Regional Committee, in September 2012 in Malta, endorsed the six overarching targets were, along with the need for their quantification, the identification of indicators and further development as appropriate. The rationale for choosing these targets – listed below – is that either they are in line with contemporary global target setting efforts, such as in the area of noncommunicable diseases, or they extend and
update previous European target setting strategies and approaches already acknowledged or agreed upon by European Member States. Here are the targets.

1. Reduce premature mortality in Europe.
2. Increase life expectancy in Europe.
3. Reduce inequities in health in Europe.
4. Enhance the well-being of the European population.
5. Provide universal coverage in Europe.
6. Establish national targets set by Member States.

With the assistance of an international expert group, the WHO Secretariat is completing the identification of indicators during 2013, in collaboration with Member States, for subsequent submission to the Regional Committee in September 2013. Different experiences from Member States across the European Region also highlight existing efforts to develop national targets.

3. How we are getting there and what we value: the case for measuring well-being

The WHO definition describes health “not merely as the absence of disease or infirmity” but as “physical, mental and social well-being”. Nevertheless, for more than 60 years WHO has neither measured nor reported on well-being. Instead, it has focused its reporting on death, disease and disability. While this monitoring function is clearly a core mandate for the Organization, WHO is partnering with other institutions to describe the well-being of populations and to measure progress on the enhancement of well-being in Europe in the context of Health 2020.

What does well-being mean? What makes up a “good life” is one of the basic moral discussions common across all philosophical traditions. Across countries, people usually agree on the “big picture” or the minimum ingredients of well-being, even if the identification of important areas or components remains a normative exercise. What matters to people’s lives is also surprisingly constant, indicating that what we value does not change easily. It is a multidimensional concept. Well-being and health are interactive concepts, with some common determinants, such as the health system.

Direct relevance to Health 2020

Discussions with representatives from European Member States and technical experts during the extensive consultation on Health 2020 provided qualitative evidence that, across the European Region, people value health and want to minimize disease. In terms of important broader determinants of health and well-being, they value social cohesion and inclusion, so that all people have a fair chance for health. People also value security and safety, which are related to health in the context of well-being. Common values across Europe increase the possibility of having a regional target for health and well-being.
Why is this important for health? Policy-makers, public health practitioners and people living in communities across Europe agree that well-being includes health, and that health is an essential part of – if not a prerequisite for – well-being. Health matters for well-being, and specifically several aspects of health, including the physical, mental and social. Moreover, research shows that there are two-way relationships between different areas of well-being: it is clear that health influences overall well-being, but well-being is also an indicator of future health or illness. Further, reviews of studies to date suggest that the effect of well-being on health is substantial (though variable), and has a comparable effect to other risk factors that are more traditionally targeted by public health interventions, such as a healthy diet.

Why is this important to governments and societies across Europe? The past few years have witnessed a number of national and international initiatives promoting the policy use of well-being indicators that reach beyond measuring economic performance and, within the health sector, that can supplement standard metrics of mortality, disability or disease. These initiatives are very diverse in their scope, methods, targets and key audiences. Another goal shared by some of these initiatives is to involve citizens in the definition of measures of well-being and progress.

Improving – or at least maintaining – well-being is part of the social contract between governments and the people they represent. Ensuring a good life is not owned by any particular sector or service, as it is a multidimensional concept with multiple determinants. Improving population well-being can be a platform to develop a common agenda, including a whole-of-government approach, across sectors and stakeholders. In addition to governments, major actors interested in well-being include civil society groups, patient groups, wellness and health promotion practitioners and the media.

**What we can build on**

Efforts that have considered measuring well-being at the population level are in practice more relevant than others that focus on specific clinical subpopulations in order to develop targets and indicators to monitor and report on health and well-being at the level of the European Region. The WHO Regional Office for Europe made a systematic literature review of validated tools for the measurement of well-being, to increase understanding in this area. The complete report also highlights and reviews efforts led by those working with the Regional Office as partners on this common challenge, including national governments in Europe, other international organizations, WHO at the international level, private firms and efforts commissioned by the United Nations. All feature health as an important component of well-being, or a factor directly affecting it; a few draw on the same data sets collected through international surveys; and many measure the same components, or domains, of what makes up a good life, although they sometimes use different words for a particular domain, such as quality of life, subjective well-being or happiness.

**What challenges we face**

Despite general agreement on what makes a good life, as well as multiple tools and approaches to measure health and well-being, researchers agree that the field of measuring
well-being would benefit from additional clarity and more rigorous assessment methods. Some of the challenges are a narrow conceptualization of health and well-being; limited data sources, yet a vast number of tools and indices; greater reliance on mortality or illness measures than those that assess positive health; and a lack of meaningful approaches to communicating and interpreting multidimensional concepts.

With strong support for an overarching target addressing health and well-being as part of the new European health strategy, Health 2020, the WHO Regional Office for Europe is working with technical partners to provide operational clarity on how health is measured in the context of well-being. In parallel, the Regional Office has entered a process of intense consultation with Member States. With the approval of governing bodies, the Regional Office expects to have the following results by 2013, elaborated on in the full report:

- a framework and definition of well-being that is conceptually sound (as far as possible the operational approach to measurement should draw on existing models that have been used at the population level);
- identification of the range of domains and subsequent indicators – for example, linked to the International Classification of Functioning, Disability and Health (ICF), WHO’s framework for measuring health and disability at both individual and population levels, which complements WHO’s International Classification of Diseases (ICD);
- indicators and an approach to their measurement, identified to measure each aspect of the health domain, that are tied to an agreed target identified for monitoring progress towards the Health 2020, to improve population health in the context of well-being;
- clarity on the way policy-makers, health professionals and other interested stakeholders across the European Region can use this information as input to policy-making and interventions, joined up with different sectors (for use within Health 2020, both the information content of well-being measures and the entry points need to be considered carefully, along with potential limitations to using well-being indicators); and
- a recognition that a large number of countries in the Region does not currently have national efforts (whether within the health ministry, other ministries or national statistical agencies) to conceptualize, collect or use information on health and well-being (any effort to improve well-being at the regional level should consider options to support a broad range of countries, with different data and measurement starting points).

Improving health and well-being is recognized as an essential component of Health 2020. A wide range of ongoing activities measuring well-being at the international level in Europe, as well as many national initiatives, provides a strong basis for the WHO Regional Office for Europe to build on work in this field: in particular, measuring health in the context of well-being and setting out a research agenda that improves both the methods for assessing and communicating measures and the policy processes to enhance the use of information that improves health and well-being.

The Regional Office will also support the policy use of health and well-being measures:

- drawing on strategies reflecting the WHO European Region’s comparative advantage in several areas, including approaches to disseminating policy relevant information, working in collaboration with partner European institutions and Member States;
Executive summary

- advising on how well-being indicators should be interpreted and used in connection with standard measures of mortality, morbidity and health system performance indicators; and
- undertaking a more innovative role in providing evidence on the mechanisms and tools for the health sector to enhance well-being in other sectors.

4. Countdown to 2020: marking progress

Although the previous sections document a plethora of health information available in the WHO European Region, the measurement challenges remain considerable. Relevant data to measure progress are not available for all countries; definitions vary between countries and disease classifications are not homogeneously applied; vital-event registration varies in population coverage and is not yet compulsory in all countries in the Region.

One of the major challenges for the implementation and impact of Health 2020 is data availability to monitor progress at the country level. The overarching or “headline” targets and their indicators have been selected for their importance in achieving Health 2020 but also for their availability. Most proposed indicators for the targets listed are supplied by information that is either routinely or regularly collected in most European countries. Nevertheless, differences in definitions, coverage of the population and data quality make comparisons difficult. WHO is committed to assisting Member States in enhancing their health information reporting and in monitoring progress with the implementation of Health 2020.

Key requirements for addressing the challenges that the WHO Regional Office for Europe faces are listed below. These are also the areas where the Regional Office could add value with its partners:

- addressing the range of data sources and norms/standards – from vital statistics to household surveys – so that these contain common ways to disaggregate population data, by social strata or other types of strata, to enable and increase accountability of monitoring of health inequalities within and across countries;
- establishing how to allocate all 53 countries within meaningful and policy relevant sub-European aggregations and report on subregional trends. This is important as 14 Member States are not included in the three subregional groups (covering the EU and the Commonwealth of Independent States) currently used;
- increasing the validity and reliability of data reported to the Health for All database from across all 53 countries, in light of collaborative work with Member States and other regional entities;
- improving the measurement and reporting of the overall burden of disease, including morbidity and other conditions particularly addressing mental health, thus providing a more comprehensive picture of the regional burden of disease;
- collecting, linking and making accessible data across health and other sectors, to support health in all approaches and intersectoral analysis, policy-making, monitoring and evaluation;
• making progress on surveillance and outbreak monitoring, as well as compliance with the International Health Regulations (IHR) across the European Region, and via connections with other regions (through governance for health and tackling communicable diseases); and
• developing a roadmap to address these challenges and achieve a solutions, identifying collaborators, resources and processes, and a research agenda to support joined-up action.

To respond to these challenges, WHO will support the development, updating or redesign of tools and instruments to enhance health information collection, analysis and reporting at the country level. WHO is working with the EC and OECD to establish a single integrated health information system for Europe, covering all 53 Member States. This work is well under way, along with the development of a health information strategy for Europe.