Summary of the roadmap document adopted on 4 February 2020 by the signatories of the Declaration

Towards access to at least 1 million sequenced genomes in the EU by 2022
Interpreting people’s health characteristics – including their genomes – is key to delivering effective health and care. Genome sequencing and data linking can help doctors predict, prevent, and diagnose diseases and treat people and patients better based on their individual genomic characteristics. Many more people could benefit from sharing data about their genomes and from doctors’ access to the wealth of linked genomic databases. People with four areas of disease – cancers, common complex diseases, infectious diseases, and rare diseases – may especially benefit. The potential advantages to health systems through enhanced prevention policies are enormous, thanks to available reference genomic data for screening and personalised medicine. Europe’s health systems are likely to gain from the savings that result from higher disease prediction rates, earlier diagnoses, and the specific tailoring of therapies to individuals. Benefits will come to the economy too: many data-driven innovations and the opening-up of new and novel markets to health and care-related services and products lie ahead.

Understanding the role of genomics and personalised medicine

Every individual is unique. So too is every genome. Combined with an understanding of people’s lifestyles and environments, personalised medicine can support improved disease diagnosis, treatment, and prevention based on each individual’s genomic characteristics. It can initiate a shift away from “one size fits all” treatments towards individually designed therapies and treatments.

Massive population-based analysis and significant advances in the early diagnosis and prevention of disease can only be achieved when a critical mass of collected data has been reached. This will be based on data and knowledge shared between organisations and across borders. Indeed, the larger the available sample collections and associated datasets, the more robust the analysis. Among future activities will be interpreting common yet complex disorders such as diabetes; understanding cancer in its many varieties; recognising individuals’ susceptibilities to infectious diseases; and diagnosing and finding leads for the treatment of patients with very rare diseases. In all of these domains, it will be important to generate large enough cohorts of people, with their genetic information, to investigate diseases and conditions bearing in mind the great diversity of Europe’s population.

Today, the key is to enable sharing of genomic and other data. Research and clinical data need to be used together by linking the information rather than bringing it all together physically in a central place. This needs to happen without breaching individuals’ privacy or compromising data security. Data-sharing is the future for Europe’s health and care. It will support and help in implementing a number of European Union initiatives and activities, such as Europe’s Beating Cancer Plan, Tackling cancers, common complex, infectious, and rare diseases

Tackling cancers, common complex, infectious, and rare diseases

Genome sequencing – the ability to analyse the unique sequence of an individual’s DNA – helps doctors and scientists understand diseases better (for example, how to predict, prevent, diagnose and treat diseases). Genomics can help determine the future risks of various debilitating diseases, such as cancer and many common complex diseases, as well as supporting the diagnosis of rare diseases.

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Collecting and analysing genomes will also help experts to understand and leverage pharmacogenomics – the branch of genetics concerned with determining an individual’s response to a particular medication – more effectively. As a result, doctors will be able to choose the right treatment for the right patient by predicting reliably the side-effects of medicines and identifying who would be a non-responder to a specific therapy.

Personalised medicine is expected to bring significant socio-economic benefits to Europe. It will result in faster and more accurate diagnosis of diseases, better identification of the likely responses of individuals to medication, and advances in preventive medicine. For people, the beneficial outcomes will include: better health, better quality of life, and increased life expectancy. For health systems, savings will result from the greater efficiencies and cost savings that are the consequence of personalised medicine. For the economy, these scientific and medical advances are likely to form the basis of data-driven innovation, and the development of new and novel markets for services, products, and medicines.
Louise has a long family history of breast cancer. One day, she heard on TV that mutations in the BRCA1 and BRCA2 genes increase the risk of breast cancer by up to 80%. Although it turned out that she doesn’t have mutations in these genes, her risk of developing the condition is still high due to her family history.

In order to detect early potential breast cancer, she started breast cancer screening at age 35, much earlier than the usual recommended routine screening. A few years later, Louise indeed developed breast cancer. To determine the most suitable type of treatment, she underwent a genomic test assessing the utility of chemotherapy versus other treatments in her case.

Test results showed that Louise would not benefit much from chemotherapy, so doctors opted for a more effective personalised treatment, which also prevented her from experiencing the unpleasant side effects of chemotherapy.

Justyna heard about the ongoing biobanking project in her country in the media, so she read more about the genetic risks of certain diseases. Her healthcare provider recommended a genomic analysis to estimate her Polygenic Risk Score (PRS), a test for most common diseases which was just introduced as a new clinical trial in the university hospital.

Now Justyna has to take statins. The genomic analysis also showed that one specific statin could increase her risk of muscle inflammation and should be avoided.

The 1+MG network will greatly increase the availability of this knowledge to oncologists, improving the choices for patients and the reducing over-treatment.

Developments in the sequencing of cancer genomes are rapidly improving insights and predictive power of genomic tests on risk, prognosis and treatment of cancer. The 1+MG network will greatly increase the availability of this knowledge to oncologists, improving the choices for patients and the reducing over-treatment.

Imagine we would have the EU federated genomic data network of 1+MG in place offering European clinicians the possibility to receive alerts when they encounter patients with similar characteristics. Patients’ increased susceptibility to life-threatening conditions caused by an infectious disease such as COVID-19 could be rapidly discovered and the necessary treatment adapted and personalised.

Pablo (27 years old) develops severe COVID-19 symptoms. Shortly after, his brother Pedro (30 years old) is also tested positive. Both need to be treated in intensive care. The two brothers do not belong to any of the known risk groups, but their close family relationship suggests a genetic risk factor.

Genetic profiling of the brothers is undertaken and compared to other disease cohorts and population biobank resources. It turns out that Pablo and Pedro are carriers of a rare genetic mutation in TLR7 (Toll-Like Receptor 7). This gene on the X chromosome regulates the interferon level as a defensive response to viruses. Based on this finding, Pablo and Pedro receive a specific treatment to restore the natural capacity of their immune system to fight the virus.
Cooperating Europe-wide on genomics

By 2018, several countries in Europe had established national initiatives that aim to generate genomic data and link it to phenotypic data (the observable health properties of a person). To help transform health and care in Europe, genomic and linked phenotypic data need to be accessed not only within but also across national boundaries.

Moving towards federated secure access to genomic data

Making collective use of Europe’s genomic data is a considerable challenge. Data security aspects play a critical role: trust needs to be ensured along several dimensions. This initiative pursues a federated approach which will bring together the strengths and qualities of national programmes, while drawing on their synergies and ensuring full compliance with data protection rules and standards. The initiative will not pool genomic data in a central repository. Instead, accredited users will be able to send approved algorithms to the many local databases involved.

Collaborating on the 1+ Million Genomes Initiative

The 1+ Million Genomes initiative, also referred to as ‘1+MG’, was launched on 10 April 2018, the European Union’s second Digital Day, when 13 European Union countries signed the declaration: “Towards access to at least 1 million sequenced genomes in the European Union by 2022”. The countries agreed to work collectively on a cross-border network of national genome cohorts associated with other relevant data. The aim is to advance data-driven research and healthcare solutions to benefit European citizens. By autumn 2020, more than 20 European countries have joined the initiative, a few others have observer status. The initiative is open for more Member States of the European Union, the European Economic Area (EEA) and European Free Trade Association (EFTA) to join it.

What is the 1+ Million Genomes initiative?

Pooling the knowledge encoded in genomic data is an area of great promise for Europe’s citizens, healthcare systems, and innovators. One country or one research institute working on its own cannot reach the scale of genomes needed to understand and improve the health of Europe’s people. Research, policy and care practice need to be brought together to focus on translating research outcomes into clinical applications and informing policy. This is why so many European countries are enrolled in the move forward on the 1+ Million Genomes initiative. Joining efforts means multiplying opportunities and will result in benefits for health and economies. Until 2023, Europe will be working on solutions to six key challenges that affect this pan-European initiative. Among them are governance, infrastructure, and interoperability challenges.

Among the six activities needed are: set up a mechanism for cross-border collaboration and data governance; examine the underlying ethical, legal, and social issues around expanding work on the collection and sharing of genomes; determine the minimal set of health data to be included in every dataset and develop good genomics practice; design the infrastructure needed and determine how to implement it; and apply the work to the main use cases of rare diseases, cancers, complex common diseases, and infectious diseases (e.g. COVID-19).

Today, Europe’s countries are at very different stages of development in the field of genomics. The aim is therefore to set up and achieve a programme to upgrade and connect already existing national and regional genomic initiatives or support their establishment where this is needed.

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23 countries have signed the 1+MG Declaration since 2018

The aim is to create a framework that will enable linking, access, and analysis of genomic and health data in Europe, both inside and across national boundaries.

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Which are the main areas of cooperation?

To create this framework with its focus on more than one million genomes, new ways of collaborating are needed. This is why the 1+ Million Genomes initiative was founded. The signatory countries are keen to move forward on accelerating developments in genomic science into enhanced healthcare. Various areas of activity are at the core of the initiative’s work: organisation; implementing personalised healthcare in health systems in cost-effective and efficient ways; ethical, legal, and social issues; health and phenotypic data; good genomic practice and standards; and infrastructure and interoperability.

Organisation of work

Strong coordination and collaboration of all the initiative’s efforts have been put in place. This is happening through cooperation among the signatory countries, supported by expert working groups. One expert group focuses on governance and ensures overall coordination. It covers several aspects of activity, including stakeholder engagement, industry engagement, a learning framework, and education and communication.

Ethical, legal, and social issues

A number of ethical, legal, and social issues need to be brought into focus and resolved. Examples include the values and ethics underpinning this pan-European framework; the underlying legal framework and challenges of finding ways to process genomic data across borders despite legal differences; the contractual arrangements related to data use and data inclusion; and the societal implications, transparency, policies, and processes to govern data access.

Genomics innovation in national healthcare systems

Embedding genomics expertise and research outcomes into personalised healthcare will lead to greater efficiencies and, thus, cost savings. Accelerating the transfer of research knowledge into improved and cost-effective healthcare is key (including improving disease prediction rates, diagnosis, and the tailoring of therapies and treatments to individuals). This requires innovation in healthcare systems across Europe: the 1+MG initiative will help the countries involved to learn from each other in this respect. Analysing the potential costs and benefits of the adoption of genomics in healthcare will be part of these steps. The 1+MG framework will thus assist in the analysis of economic viability, and in the harmonisation of cost assessment and economic evaluation models. This will help towards generating comparable evidence across European healthcare systems and facilitate countries to make informed policy choices about the implementation of genomics-based personalised medicine.

Clinical and phenotypic data

Agreeing on a minimal set of phenotypic and clinical data to be collected in each genomics study is crucial. This agreement will enable comparative studies across data collections that may have different backgrounds. The initiative will build consensus to implement this minimal dataset across European laboratories and clinics.

Good genomic practice and standards

When sequencing genomic data, it is important to draw on good practice. As a result, guidelines and appropriate standards and forms of accreditation will need to be drawn up. Genome sequencing standards, benchmarking and metrics, as well as good quality control of data, are needed.

Infrastructure and interoperability

The initiative’s proposed infrastructure uses a federated approach. Each country in Europe is invited to contribute to the endeavour, by bringing to the enterprise the strengths and qualities of its own national programme. The initiative is not centralised and will not pool genomic data in one place. Rather, through interoperability, it will bring data warehouses up to agreed standards and link them with a secure infrastructure. Users, like accredited research institutions, healthcare professionals, and health policy agencies, will be able to access this secure network to execute approved algorithms and analyse available data remotely.
What is the timeline and next steps?

The 1+ Million Genomes initiative has set up an intensive agenda of activities to reach its goals by 2022. Its signatory countries and expert groups are working together on many stepwise, concrete actions. By aligning the investments and advances expected in 2020-2022, the aim of creating a cohort of at least 1 million sequenced genomes can be achieved.

### Laying out the 1+MG Roadmap 2020-2022

Between 2020-2022, three phases of activity will concentrate on engaging, translating, and driving the 1+MG initiative forward.

**2020 – Engage** – Engaging countries and other stakeholders in governance and collaboration

- Assemble national, regional, and local public authorities as well as a broad range of international stakeholders in the initiative. They will link together different genomic medicine resources in line with the initiative’s declaration.
- Define a governance model of coordination and cooperation. The cooperation model will cover the terms and conditions needed for secure access to genomic data and associated health data across borders, and the use of national and regional data collections for research and improving healthcare.

**2021 – Translate** – Translating the mission into concrete infrastructure, guidelines, and pilots

- Pilot the technical infrastructure for distributed, authorised and secure access to national and regional genetic data, and facilitate the interoperability of registries and databases.
- Support the development of an ethical and legal framework for cross-border access to genomic datasets to advance research across Europe, while taking appropriate measures to protect the privacy of individuals.
- Test-run the secure infrastructure and legal framework to enable cross-border sharing and analysis of genetic data and associated health datasets, anonymised as appropriate. The datasets will come from multiple European countries, build on existing initiatives in genomics and personalised medicine, and focus on four use cases.
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- Test-run the secure infrastructure and legal framework to enable cross-border sharing and analysis of genetic data and associated health datasets, anonymised as appropriate. The datasets will come from multiple European countries, build on existing initiatives in genomics and personalised medicine, and focus on four use cases.

**2022 – Drive** – Moving forward on sharing, scaling, and sustaining the initiative

- Embed the coordinated data governance framework and infrastructure for Europe-wide large-scale processing of genome and health data in national and regional science and healthcare systems.
- Scale activities to enable distributed and secure access to research cohorts of at least 1 million sequenced genomes.

The underpinning aim of this three-year initiative will be to support a European data strategy that reinforces the creation of data spaces in key strategic sectors and domains of public interest. In the health sector, this will translate into a European health data space, essential for making advances in preventing, detecting, and curing diseases and for making informed decisions that will improve the accessibility, effectiveness, and sustainability of healthcare systems.

A Horizon 2020 project “Beyond 1 Million Genomes” (B1MG) will provide support and coordination to the implementation of the roadmap.

For more general information, the 1+ Million Genomes initiative encourages people to visit its website:
1+Million Genomes
Roadmap 2020-2022

More information
European Commission
DG CONNECT
Unit H3 - eHealth, Well-Being and Ageing
EUF0 02/161
L-2557 Luxembourg
CNECT-H3@ec.europa.eu

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www.digitalhealtheurope.eu

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Gesellschaft für Kommunikations- und Technologieforschung mbH
Oxfordstr. 2 · 53111 Bonn, Germany
https://www.empirica.com